# **MPRAflow Documentation**

Release 2.1

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## Installation Getting Started

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This pipeline processes sequencing data from Massively Parallel Reporter Assays (MPRA) to create count tables for candidate sequences tested in the experiment.

- Installation & Getting Started Instructions for the Installation of the program and some examples to get you started.
- MPRAflow Workflows An overview of how MPRAflow works and documentation for the MPRAflow sub workflows.
- MPRAflow Examples Muliple examples from the literature are listed for every sub workflow in MPRAflow.
- **Project Information** More information on the project, including the changelog, list of contributing authors, and contribution instructions.

## CHAPTER 1

Quick Example

## CHAPTER 2

## Features

- Association This utility takes in library association sequencing data (FASTQ) and a design file (FASTA) to assign barcodes to the corresponding elements tested. Functionality includes filtering for quality and coverage of barcodes. This utility must be run before the COUNT utility.
- **Count** This utility processes sequence data (FASTQ) of barcodes from the DNA and RNA fractions of the MPRA experiment and outputs count tables labeled with the element tested and a label provided in the design file. This utility can process multiple replicates and conditions in a parallelized manner. Based on a user specified flag, the pipeline will either output normalized activity for each tested sequence, or will combine the results into a single count matrix compatible with MPRAnalyze.
- Association Saturation mutagenesis This workflow is about assocation variant calls with barcodes. Variants are introduced by an error-prone PCR. The workflow takes the sequencing of the region, with barcodes in index read and the reference sequence and maps the reads to the reference, calls variants and associates them with the corresponding barcode.
- **Saturation mutagenesis** This workflow is about getting single variant effect from a target with multiple mutations generated by error-prone PCR. The workflow takes counts (e.g. from the count workflow), combines them with an association file (variants to barcodes, e.g. from saturation mutagenesis assocation) and uses a generalized linear model to to detect single variant effects.

## CHAPTER 3

## Feedback

The best place to leave feedback, ask questions, and report bugs is the MPRAflow Issue Tracker.

## 3.1 Indices and tables

- genindex
- search

## 3.1.1 Quickstart

1. Create an experiment.csv in the format below, including the header. *DNA\_R1* or *RNA\_R1* is name of the gzipped fastq of the forward read of the DNA or RNA from the defined condition and replicate. *DNA\_R2* or *RNA\_R2* is the corresponding index read with UMIs (excluding sample barcodes) and *DNA\_R3* or *RNA\_R3* of the reverse read. If you do not have UMIs remove the columns *DNA\_R2* and *RNA\_R2* or leave them empty.

```
Condition, Replicate, DNA_R1, DNA_R2, DNA_R3, RNA_R1, RNA_R2, RNA_R3
condition1, 1, cond1_rep1_DNA_FWD_reads.fastq.gz, cond1_rep1_DNA_IDX_reads.

→fastq.gz, cond1_rep1_DNA_REV_reads.fastq.gz, cond1_rep1_RNA_FWD_reads.fastq.gz
condition1, 2, cond1_rep2_DNA_FWD_reads.fastq.gz, cond1_rep2_DNA_IDX_reads.

→fastq.gz, cond1_rep2_DNA_FWD_reads.fastq.gz, cond1_rep2_RNA_FWD_reads.fastq.

→gz, cond1_rep2_RNA_IDX_reads.fastq.gz, cond1_rep2_RNA_FWD_reads.fastq.gz
condition2, 1, cond2_rep1_DNA_FWD_reads.fastq.gz, cond2_rep1_DNA_IDX_reads.

→fastq.gz, cond2_rep1_DNA_REV_reads.fastq.gz, cond2_rep1_RNA_FWD_reads.fastq.gz
condition2, 2, cond2_rep2_DNA_FWD_reads.fastq.gz, cond2_rep1_RNA_FWD_reads.fastq.gz
condition2, 2, cond2_rep2_DNA_FWD_reads.fastq.gz, cond2_rep2_RNA_FWD_reads.fastq.gz
condition2, 2, cond2_rep2_DNA_FWD_reads.fastq.gz, cond2_rep2_RNA_FWD_reads.fastq.gz
condition2, 2, cond2_rep2_DNA_REV_reads.fastq.gz, cond2_rep2_RNA_FWD_reads.fastq.gz
condition2, 2, cond2_rep2_DNA_REV_reads.fastq.gz, cond2_rep2_RNA_FWD_reads.fastq.gz
condition2, 2, cond2_rep2_DNA_REV_reads.fastq.gz, cond2_rep2_RNA_FWD_reads.fastq.gz
cond2_rep2_RNA_IDX_reads.fastq.gz, cond2_rep2_RNA_FWD_reads.fastq.gz
```

2. If you would like each insert to be colored based on different user-specified categories, such as "positive control", "negative control", "shuffled control", and "putative enhancer", to assess the overall quality the user can create a 'label' tsv in the format below that maps the name to category:

insert1\_name insert1\_label
insert2\_name insert2\_label

The insert names must exactly match the names in the design FASTA file.

3. Run Association if using a design with randomly paired candidate sequences and barcodes

**Note:** This will run in local mode, please submit this command to your cluster's queue if you would like to run a parallelized version.

4. Run Count

Be sure that the experiment.csv is correct. All fastq files must be in the same folder given by the --dir option. If you do not have UMIs please use the option --no-umi. Please specify your barcode length and umi-length with --bc-length and --umi-length.

5. Run saturation mutagenesis

**Note:** The experiment file is different from the count workflow. It should contain the condition, replicate and filename of the counts, like:

```
Condition, Replicate, COUNTS
condition1, 1, cond1_1_counts.tsv.gz
condition1, 2, cond1_2_counts.tsv.gz
condition1, 3, cond1_3_counts.tsv.gz
condition2, 1, cond2_1_counts.tsv.gz
condition2, 2, cond2_2_counts.tsv.gz
condition2, 3, cond2_3_counts.tsv.gz
```

The count files can be generated by the count workflow, are named: <condition>\_<replicate>\_counts.tsv.gz and can be found in the outs/<condition>/ <replicate> folder. They have to be copied or linked into the --dir folder.

## 3.1.2 Installation

Installation should take less than 10 minutes

#### System Requirements

CentOS Linux 7 or above

#### **Required packages**

conda 4.6 or above

Download here: https://docs.conda.io/en/latest/miniconda.html

#### **Clone repository**

git clone https://github.com/shendurelab/MPRAflow.git

#### Set up conda environment

This pipeline uses python2.7 and python3.6 with additional R scripts. Three *.yml* files are provided to create the appropriate environments and is completely handled by nextflow. The whole pipeline is set up to run on a Linux system. The general environment with nextflow located in the home directory called *environment.yml*.

Install the the conda environment. The general conda environment is called MPRAflow.

```
cd MPRAflow
conda env create -n MPRAflow -f environment.yml
```

If you do not have access to the internet, you have to run the previous command on a node with internet. Afterwards you need to start nextflow too (see Steps to run the pipeline). After creation of the second conda environment by nextflow you can cancel it and start it on your internal node. Be aware that folders must have access on all nodes.

Nextflow has problems using conda 4.7 and highet, because the source activate command is replaced by conda activate. If you get error messages after running you can make a symbolik link of the activate command from you bin folder of the conda or miniconda folder to your MPRAflow environment bin folder. E.g. like:

ln -s ~/miniconda3/bin/activate ~/miniconda3/envs/MPRAflow/bin/activate

#### **Quick test**

```
conda activate MPRAflow
nextflow run count.nf --help
nextflow run association.nf --help
nextflow run saturationMutagenesis.nf --help
```

#### 3.1.3 Running MPRAflow on HPC Cluster

Nextflow gives us the opportunity to run MPRAflow in a cluster environment. Right now we split up processes into two main groups: *longtime* and *shorttime*. We can define different job setting for both groups. As you can imagine from the names *longtime* defines processes that takes a while when running. Sometimes several days. *shortime* defines processes that are quicker and are usually done in several minutes.

To enable the submission to your cluster you have to edit the *conf/cluster.config*, allowing each process to be run as a separate *qsub*, *sbatch* or similar command. The config contains example code for SGE, LSF, and SLURM

architectures. Please remove the \ for the architecture you would like to use and place \ in front of any architectures not currently in use. A \ in front of all of them runs the pipeline on your local machine (default). If you run MPRAflow on a cluster system make sure be that you export all environment variables. E.g. this can be done with the -V option by SGE.

**Note:** Please consult your cluster's wiki page for cluster specific commands and change clusterOptions = to reflect these specifications. Additionally, for large libraries, more memory can be specified in this location.

## 3.1.4 Overview

This pipeline processes sequencing data from Massively Parallel Reporter Assays (MPRA) to create count tables for candidate sequences tested in the experiment.

This package contains three utilities:

## ASSOCIATION (association.nf)

This utility takes in library association sequencing data (FASTQ) and a design file (FASTA) to assign barcodes to the corresponding elements tested. Functionality includes filtering for quality and coverage of barcodes. This utility must be run before the COUNT utility. See *Association* for more details.

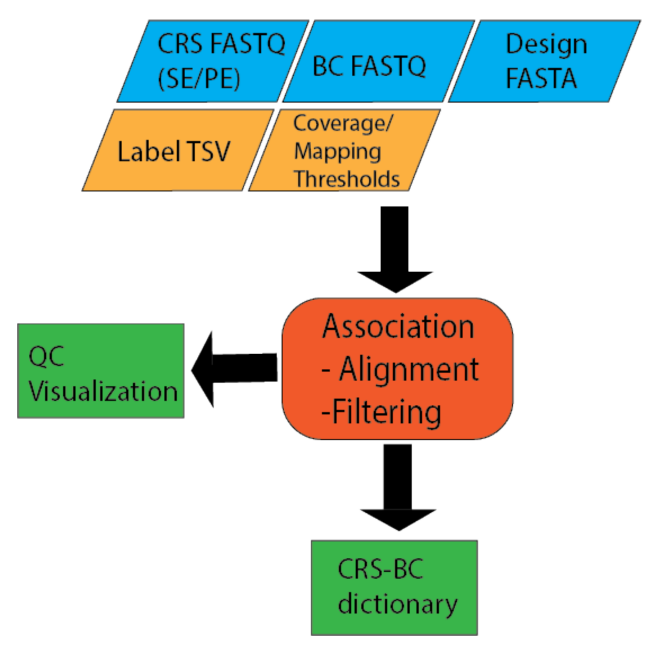
## COUNT (count.nf)

This utility processes sequence data (FASTQ) of barcodes from the DNA and RNA fractions of the MPRA experiment and outputs count tables labeled with the element tested and a label provided in the design file. This utility can process multiple replicates and conditions in a parallelized manner. Based on a user specified flag, the pipeline will either output normalized activity for each tested sequence, or will combine the results into a single count matrix compatible with MPRAnalyze. See *Count* for more details.

## Saturation mutagenesis (saturationMutagenesis.nf)

This workflow is about getting single variant effect from a target with multiple mutations generated by error-prone PCR. The workflow takes counts (e.g. from the count workflow), combines them with an association file (variants to barcodes) and uses a generalized linear model to to detect single variant effects. See *Saturation mutagenesis* for more details.

## 3.1.5 Association



## Input files

## **Fastq Files**

2-3 Fastq files from library association sequencing –Candidate regulatory sequence (CRS) sequencing, 1 forawrd read and an optional reverse read if paired end sequencing was used –Barcode sequence, 1 read covering the barcode

## **Design File**

Fasta file of of CRS sequences with unique headers describing each tested sequence

#### Example file:

>CRS1	
GACGGGAACGTTTGAGCGAGATCGAGGATAGGAGGAGCGGA	
>CRS2	
GGGCTCTCTTATATTAAGGGGGTGTGTGAACGCTCGCGATT	
>CRS3	
GGCGCGCTTTTTCGAAGAAACCCGCCGGAGAATATAAGGGA	
>CRS4	
TTAGACCGCCCTTTACCCCGAGAAAACTCAGCTACACACTC	

## Label File (Optional)

Tab separated file (TSV) of desired labels for each tested sequence

Example file:

```
CRS1 Positive_Control
CRS2 Negative_Control
CRS3 Test
CRS4 Positive_Control
```

**Note:** If you provide a label file, the first column of the label file must exactly match the FASTA file or the files will not merge properly in the pipeline.

#### association.nf

#### **Options**

**Optional:** 

With --help or --h you can see the help message.

#### Mandatory arguments:

fastq-insert	Full path to library association fastq for insert (must be surrounded with quotes)
fastq-bc	Full path to library association fastq for bc (must be surrounded with quotes)
design	Full path to fasta of ordered oligo sequences (must be surrounded with quotes)
name	Name of the association. Files will be named after this.
fastq-insertPE	Full path to library association fastq for read2 if the library is paired end
	(must be surrounded with quotes)
min-cov	(must be surrounded with quotes) minimum coverage of bc to count it (default 3)
min-cov min-frac	
	minimum coverage of bc to count it (default 3)

cigar	require exact match ex: 200M (default none)
outdir	The output directory where the results will be saved and what will be used as a prefix (default outs)
split	Number read entries per fastq chunk for faster processing (default: 2000000)
labels	tsv with the oligo pool fasta and a group label (ex: positive_control) if no labels desired a file will be automatically generated

#### Processes

Processes run by nextflow in the Association Utility. Some Processes will be run only if certain options used and are marked below.

**count\_bc or count\_bc\_nolab (if no label file is provided)** Removes any illegal characters (defined by Piccard) in the label file and design file. Counts the number of reads in the fastq file.

create\_BWA\_ref Creates a BWA reference based on the design file

- PE\_merge (if paired end fastq files provided) Merges the forward and reverse reads covering the CRS using fastq-join
- align\_BWA\_PE or align\_BWA\_S (if single end mode) Uses BWA to align the CRS fastq files to the reference created from the Design File. This will be done for each fastq file chunk based on the split option.

collect\_chunks merges all bamfiles from each separate alignment

map\_element\_barcodes Assign barcodes to CRS and filters barcodes by user defined parameters for coverage and mapping percentage

filter\_barcodes Visualize results

#### Output

The output can be found in the folder defined by the option --outdir. It is structured in folders of the condition as

#### **Files**

count\_fastq.txt number of barcode reads

count\_merged.txt number of aligned CRS reads

design\_rmIllegalChars.fa Design file with illegal characters removed

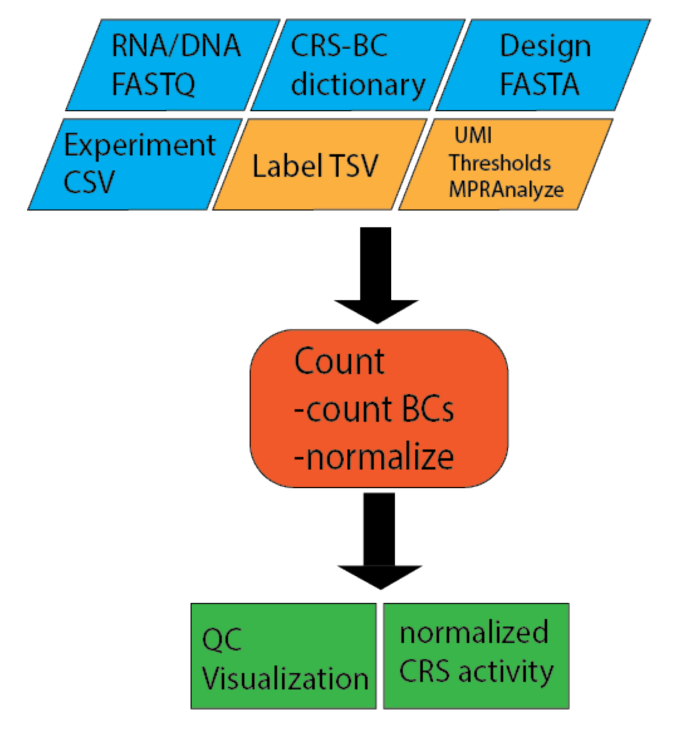
label\_rmIllegalChars.txt Label file with illegal characters removed

s\_merged.bam sorted bamfile for CRS alignment

**\${name}\_coords\_to\_barcodes.pickle** pickle file containing a python dictionary of CRS/barcode mappings

\*.png Visualization of number of barcodes mapping to enhancers

## 3.1.6 Count



## Input files

## **Experiment File**

Comma separated file (CSV) that assigns all fastq files present in a directory to a condiditon and replicate. Each line represents an experiment, which will all be processed in parallel

Condition, Replicate, DNA BC F, DNA UMI, DNA BC R, RNA BC F, RNA UMI, RNA BC R Condidtion1,1,C1R1\_DNA\_barcode\_F.fastq.gz,C1R1\_DNA\_barcode\_UMI.fastq.gz,C1R1\_DNA\_ →barcode\_R.fastq.gz,C1R1\_RNA\_barcode\_F.fastq.gz,C1R1\_RNA\_barcode\_UMI.fastq.gz,C1R1\_ ⇔RNA\_barcode\_R.fastq.gz Condidtion1,2,C1R2\_DNA\_barcode\_F.fastq.gz,C1R2\_DNA\_barcode\_UMI.fastq.gz,C1R2\_DNA\_ →barcode\_R.fastq.gz,C1R2\_RNA\_barcode\_F.fastq.gz,C1R2\_RNA\_barcode\_UMI.fastq.gz,C1R2\_ →RNA\_barcode\_R.fastq.gz Condidtion1, 3, C1R3\_DNA\_barcode\_F.fastq.gz, C1R3\_DNA\_barcode\_UMI.fastq.gz, C1R3\_DNA\_ →barcode\_R.fastq.gz,C1R3\_RNA\_barcode\_F.fastq.gz,C1R3\_RNA\_barcode\_UMI.fastq.gz,C1R3\_ →RNA\_barcode\_R.fastq.gz Condidtion2,1,C2R1\_DNA\_barcode\_F.fastq.gz,C2R1\_DNA\_barcode\_UMI.fastq.gz,C2R1\_DNA\_ →barcode\_R.fastq.gz,C2R1\_RNA\_barcode\_F.fastq.gz,C2R1\_RNA\_barcode\_UMI.fastq.gz,C2R1\_ →RNA\_barcode\_R.fastq.gz Condidtion2,2,C2R2\_DNA\_barcode\_F.fastq.gz,C2R2\_DNA\_barcode\_UMI.fastq.gz,C2R2\_DNA\_ →barcode\_R.fastq.qz,C2R2\_RNA\_barcode\_F.fastq.qz,C2R2\_RNA\_barcode\_UMI.fastq.qz,C2R2\_ ⇔RNA\_barcode\_R.fastq.gz Condidtion2, 3, C2R3\_DNA\_barcode\_F.fastq.gz, C2R3\_DNA\_barcode\_UMI.fastq.gz, C2R3\_DNA\_ -->barcode\_R.fastq.qz,C2R3\_RNA\_barcode\_F.fastq.qz,C2R3\_RNA\_barcode\_UMI.fastq.qz,C2R3\_ →RNA\_barcode\_R.fastq.gz

## **Design File**

Fasta file of of CRS sequences with unique headers describing each tested sequence

#### Example file:

>CRS1
GACGGGAACGTTTGAGCGAGATCGAGGATAGGAGGAGCGGA
>CRS2
GGGCTCTCTTATATTAAGGGGGTGTGTGAACGCTCGCGATT
>CRS3
GGCGCGCTTTTTCGAAGAAACCCGCCGGAGAATATAAGGGA
>CRS4
TTAGACCGCCCTTTACCCCGAGAAAACTCAGCTACACACTC

#### **Association Pickle**

Python dictionary of CRS to Barcodes

#### Label File (Optional)

Tab separated file (TSV) of desired labels for each tested sequence

Example file:

```
CRS1 Positive_Control
CRS2 Negative_Control
CRS3 Test
CRS4 Positive_Control
```

#### Count.nf

#### **Options**

#### Mandatory arguments:

dir	Fasta directory (must be surrounded with quotes)			
e,experime	nt-file Experiment csv file			
design	Fasta of ordered insert sequences.			
association	Pickle dictionary from library association process.			
Optional:				
labels	tsv with the oligo pool fasta and a group label (ex: positive_control), a single label will be applied if a file is not specified			
outdir	The output directory where the results will be saved (default outs)			
bc-length	Barcode length (default 15)			
umi-length	UMI length (default 10)			
no-umi	Use this flag if no UMI is present in the experiment (default with UMI)			
merge_interse	ect Only retain barcodes in RNA and DNA fraction (TRUE/FALSE, default: FALSE)			
mpranalyze	Only generate MPRAnalyze outputs			
thresh	minimum number of observed barcodes to retain insert (default 10)			

#### **Processes**

Processes run by nextflow in the Association Utility. Some Processes will be run only if certain options used and are marked below.

create\_BAM or create\_BAM\_noUMI (if no UMI sequence) creates a bamfile of barcode and UMI sequences

raw\_counts creates a table of counts for each barcode (where UMIs, if present, are deduplicated)

filter\_counts Remove barcodes that are not the appropriate length

final\_counts Record overrepresended UMIs and final count table

dna\_rna\_merge\_counts or dna\_rna\_mpranalyze\_merge Merge RNA/DNA count matrices per barcode

final\_merge (MPRAnalyze option only) Merge all DNA/RNA counts into one file

final\_label (MPRAnalyze option only) Label the barcodes

generate\_mpranalyze\_inputs (MPRAnalyze option only) Generate inputs for MPRAnalyze, counts tables and annotation tables for rna/dna

dna\_rna\_merge Merge each DNA and RNA file label with sequence and insert and normalize

calc\_correlations Calculate correlations between Replicates

make\_master\_tables Create tables of each CRS normalized across replicates

#### Output

The output can be found in the folder defined by the option --outdir. It is structured in folders of the condition as

#### Files

#### File tree

outdir
-Condition
-allreps.tsv
-average_allreps.tsv
-HepG2_1_2_correlation.txt
-HepG2_1_2_DNA_pairwise.png
-HepG2_1_2_Ratio_pairwise.png
-HepG2_1_2_RNA_pairwise.png
-HepG2_all_barcodesPerInsert_box.png
-HepG2_barcodesPerInsert.png
-Reps
-HepG2_1_counts.tsv
-HepG2_1_counts.tsv.gz
-HepG2_1_DNA_counts.tsv
-HepG2_1_DNA_raw_counts.tsv.gz
-HepG2_1_RNA_filtered_counts.tsv.gz
-HepG2_1_DNA_filtered_counts.tsv.gz
-HepG2_1_RNA_counts.tsv
-HepG2_1_RNA_raw_counts.tsv.gz

#### **Files for each Condition**

allreps.tsv TSV of normalized DNA and RNA count, ratio, log2ratio, and number of observed barcodes for each condition, replicate, of every CRS

average\_allreps.tsv mean ratio, log2 ratio, and observed barcodes per condidition normalized for all replicates

- HepG2\_1\_2\_correlation.txt correlation values for a condition and 2 replicates (ie: HepG2 replicate 1 vs replicate 2)
- HepG2\_1\_2\_DNA\_pairwise.png Correlation plot of DNA counts condition vs two reps (ie: HepG2 replicate 1 vs replicate 2)
- HepG2\_1\_2\_Ratio\_pairwise.png Correlation plot of normalized log2(RNA/DNA) condition vs two reps (ie: HepG2 replicate 1 vs replicate 2)
- HepG2\_1\_2\_RNA\_pairwise.png Correlation plot of RNA counts condition vs two reps (ie: HepG2 replicate 1 vs replicate 2)
- HepG2\_all\_barcodesPerInsert\_box.png Box plot of each CRS accross replicates for all barcodes in each condidtion. Colored by the label file.
- HepG2\_barcodesPerInsert.png Histogram of number of barcodes detected per CRS

HepG2\_group\_barcodesPerInsert\_box.png Boxplot of CRS normalized per insert, grouped by labels

#### Files for each replicate in each condidtion

HepG2\_1\_counts.tsv mean ratio, log2 ratio, and observed barcodes per condidition for each replicate

HepG2\_1\_counts.tsv.gz table of barcodes with DNA counts and RNA counts

HepG2\_1\_DNA\_counts.tsv table of barcodes with DNA counts

HepG2\_1\_DNA\_raw\_counts.tsv.gz table of barcodes, UMI, and DNA counts raw

- HepG2\_1\_DNA\_filtered\_counts.tsv.gz table of barcodes, UMI, and DNA counts raw, filtered for barcodes of correct length
- HepG2\_1\_RNA\_counts.tsv table of barcodes with RNA counts

HepG2\_1\_RNA\_raw\_counts.tsv.gz table of barcodes, UMI, and RNA counts raw

HepG2\_1\_RNA\_filtered\_counts.tsv.gz table of barcodes, UMI, and DNA counts raw, filtered for barcodes of correct length

#### 3.1.7 Association saturation mutagenesis

This workflow is about assocation variant calls with barcodes. Variants are introduced by an error-prone PCR. The workflow takes the sequencing of the region, with barcodes in index read and the reference sequence and maps the reads to the reference, calls variants and associates them with the corresponding barcode.

#### Input files

#### **Fastq Files**

3 Fastq files from library association sequencing –Reference sequencing, 1 forward and 1 reverse read –Barcode sequence, 1 read covering the barcode

#### **Reference file**

Fasta file of the referencesequence describing the mutated sequence

#### Example file:

>TERT

#### association\_saturationMutagenesis.nf

#### **Options**

With --help or --h you can see the help message.

Usage:

#### Mandatory arguments:

fastq-insert	Full path to library association fastq for insert			
fastq-insertPE	Full path to library association fastq for read2			
fastq-bc	Full path to library association fastq for bc			

	design	Full path to fasta of reference sequence (only one reference sequence) Name of the association. Files will be named after this.			
	name Name of the association. Files will be named after this.				
<b>Optional:</b>					
	<ul> <li>bc-length Barcode length (default 15)</li> <li>clipping-penalty bwa mem clipping penalty (default 80)</li> <li>min-ireads minimum number gapped reads for indel candidates (default 80)</li> </ul>				
	clipping-penalty	bwa mem clipping penalty (default 80)			
	min-ireads	minimum number gapped reads for indel candidates (default: 3)			
	split	Split up the fastq read into chunks with max limit of reads (default: 2000000)			
	outdir	The output directory where the results will be saved and what will be used as a prefix (default outs)			
	h,help	Print the help message			

#### **Processes**

Processes run by nextflow in the Association Saturation Mutagenesis Utility.

clean\_design Removes any illegal characters (defined by Piccard) in the reference file.

create\_BWA\_ref Creates a BWA reference based on the design file

get\_name Recieves the name written in the header of the reference fasta file

create\_BAM Merges the forward and reverse reads and stores them in BAM format. This is run for each chunk.

collect\_chunks combine the chunks (merge bams)

**PE\_mapping** Map the reads to the reference.

get\_count Get the barcodes and count for each barcode. Filter them

extract\_reads Create a BAM file for each barcode with the corresponding reads in it. This create a large number of files!

call\_variants: Call variants for each barcode seperately.

combine\_variants: Combine barcode/variant calls to one final output file.

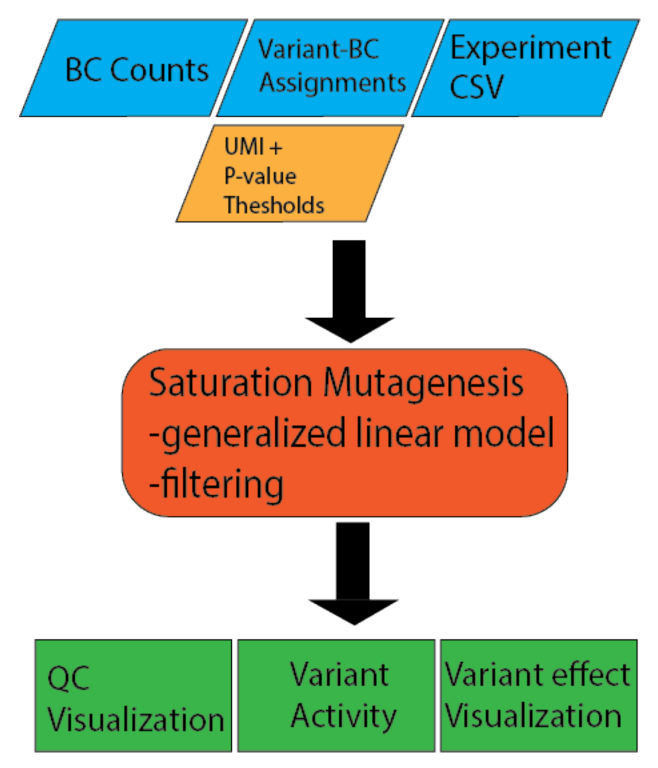
#### Output

The output can be found in the folder defined by the option --outdir and the subfolder defined by option --name. It is structured in folders of the condition as

#### **Files**

design\_rmIllegalChars.fa Reference file with remove illegal characters
\${datasetID}.variants.txt.gz Barcode to variant association. Named by the header in the reference.
counts\_\${datasetID}.filtered.tsv.gz Filtered barcode counts. Named by the header in the reference.
counts\_\${datasetID}.tsv.gz All barcode counts. Named by the header in the reference.

## 3.1.8 Saturation mutagenesis



## Input files

#### Variant-BC Assignments

The assignment file contains the variants, assigned to each barcode. It can be gzipped and it is set through the option --assignment. Each row starts with one barcode, e.g. AAAAACTAATACCA. The barcode is followed by multiple variants separated by a space. It can also be possible that a barcode has no variant. Then the barcode stands for the complete reference sequence. A variant has the format identifier:position:ref>alt, e.g. TERT:408:G>A. The *identifier* will be always the same and is use to mark the target of origin. Here the promoter TERT. The *position* is usually the position in the target transcript or any other coordinate system. *ref* is the reference base and *alt* the alternative base to the original target reference. Both can be a single or multiple nucleotide, e.g. TERT:75:GCCCC>GCCCCC is a valid variant and refers to an insertion of one C after position 79. For deletions a dot (.) can be used in the reference. So TERT:42:G>. is the deletion of the base G at position 42. It can also be written as TERT:41:TG>T.

#### Example file:

ААААААСТААТАССА	IRF6:104:A>G IRF6:408:G>A
AAAAAAGCAGGAACA	IRF6:66:A>G IRF6:373:T>A
AAAAAAGCATTCTGT	IRF6:371:G>T IRF6:510:G>A IRF6:560:C>T
AAAAACACTACTGGT	IRF6:326:C>T IRF6:509:T>A

#### **Experiment file**

Experiment file list the condition(s), replicate(s) and corresponding count files of the analysis. It is comma separated (without spaces in between) and must be set via --e or --experiment-file.

```
TERT-GBM,1,TERT-GBM_1_counts.tsv.gz
TERT-GBM,2,TERT-GBM_2_counts.tsv.gz
TERT-GBM,3,TERT-GBM_3_counts.tsv.gz
TERT-HEK,1,TERT-HEK_1_counts.tsv.gz
TERT-HEK,2,TERT-HEK_2_counts.tsv.gz
TERT-HEK,3,TERT-HEK_3_counts.tsv.gz
```

#### **DNA/RNA counts per barcode**

For each condition and replicate count file (gziped) is needed with the number of DNA and RNA counts per barcode. It is tab separated starting with the barcode, followed by the the DNA and then the RNA counts. Count files can be produced by the count workflow. See *Count*. All count files must be in the --dir folder.

Example file:

AAAAAAAAATGATAAGGAA	56	29		
AAAAAAAAATGGGAAGGCG	89	112		
AAAAAAAAATTTGCGTAAA	25	32		
AAAAAAAAATTTGCGTAAT	1	1		
AAAAAAAAATTTGGGGATA	5	26		
АААААААААСААТААGAAAT	21	55		
AAAAAAAAACCCAGAATACG	31	39		

#### SaturationMutagenesis.nf

#### **Options**

With --help or --h you can see the help message.

#### Mandatory arguments:

dir	Directory of count files (must be surrounded with quotes).
assignment	Variant assignment file.
e,experiment-file	Experiment csv file.

#### **Optional:**

outdir	The output directory where the results will be saved (default outs).
thresh	Minimum number of observed barcodes to retain variant (default 10).
pvalue	pValue cutoff for significant different variant effects. For variant effect plots only (default 1e-5).

#### Processes

Processes run by nextflow in the saturation mutagenesis utility.

- **calc\_assign\_variantMatrix** Creates the variant matrix for the linear model using only single base pair substitutions (for each condition and replicate).
- **calc\_assign\_variantMatrixWith1bpDel** Creates the variant matrix for the linear model using single base pair substitutions and 1 bp deletions (for each condition and replicate).
- fitModel Fit the matrix (variantMatrix and variantMatrixWith1bpDel) using a generalized linear model (for each condition and replicate).

#### summarizeVariantMatrix

Todo: describe summarizeVariantMatrix

- statsWithCoefficient Output of the log2 variant effects from the linear model combined with number of barcodes (for each condition and replicate).
- plotCorrelation Plots the correlation between replicates of one condition.
- **plotStatsWithCoefficient** Plots the variant effect plot of the target region using all variants larger than the threshold and the significnace level set by --p-value (for each condition and replicate).
- **fitModelCombined** Fit the matrix (variantMatrix and variantMatrixWith1bpDel) using a generalized linear model (for each condition the combined model).

#### combinedStats

**Todo:** describe combinedStats

- statsWithCoefficientCombined Output of the log2 variant effects from the linear model combined with number of barcodes (for each condition the combined model).
- **plotStatsWithCoefficientCombined** Plots the variant effect plot of the target region using all variants larger than the threshold and the significnace level set by --p-value (for each condition the combined model).

#### Output

The output can be found in the folder defined by the option --outdir. It is structured in folders of the condition as

**Files** 

Todo: Describe SatMut output files

## 3.1.9 Basic association workflow

This example runs the association workflow on 5'/5' WT MRPA data in the HEPG2 cell line from Klein J., Agarwal, V., Keith, A., et al. 2019.

#### Prerequirements

This example depends on the following data and software:

## Installation of MPRAflow

Please install conda, the MPRAflow environment and clone the actual MPRAflow master branch. You will find more help under *Installation*.

#### Meta Data

It is necessary to get the ordered oligo array so that each enhancer sequence can be labeled in the analysis and to trim any adaptors still in the sequence, in this case we trim off 15bp from the end of each sequence

#### Reads

There is one set of association sequencing for this data, which contains a forward (CRS-forward), reverse (CRS-reverse), and index (barcode) read for DNA and RNA. These data must be downloaded. All data is publically available on the short read archive (SRA). We will use SRA-toolkit to obtain the data.

Note: You need 10 GB disk space to download the data!

```
conda install sra-tools
cd Assoc_Basic/data
fastq-dump --gzip --split-files SRR10800986
cd ..
```

For large files and unstable internet connection we reccommend the comand *prefetch* from SRA tools before running *fastq-dump*. This command is much smarter in warnings when something went wrong.

```
conda install sra-tools
cd Assoc_Basic/data
prefetch SRR10800986
fastq-dump --gzip --split-files SRR10800986
cd ..
```

**Note:** Please be sure that all files are downloaded completely without errors! Depending on your internet connection this can take a while. If you just want some data to run MPRAflow you can just limit yourself to one condition and/or just one replicate.

#### With

tree data

the folder should look like this:

data

Here is an overview of the files:

Table 1: HEPG2 a	association data
------------------	------------------

Condition	GEO Ac-	SRA	SRA Runs
	cession	Accession	
HEPG2-association: HEPG2 library association	GSM423795	4 SRX7474872	SRR10800986

#### **MPRAflow**

Now we are ready to run MPRAflow and create CRS-barcode mappings.

#### **Run nextflow**

Now we have everything at hand to run the count MPRAflow pipeline. Therefore we have to be in the cloned MPRAflow folder. But we will change the working and output directory to the Assoc\_Basic folder. The MPRAflow count command is:

Note: Please check your conf/cluster.config file if it is correctly configured (e.g. with your SGE cluster commands).

If everything works fine the following 7 processes will run: count\_bc\_nolab create\_BWA\_ref, PE\_merge, align\_BWA\_PE, collect\_chunks, map\_element\_barcodes, filter\_barcodes.

#### Results

All needed output files will be in the Assoc\_Basic/output folder.

## 3.1.10 Basic Count workflow

This example runs the count workflow on 5'/5' WT MRPA data in the HEPG2 cell line from Klein J., Agarwal, V., Keith, A., et al. 2019.

#### **Prerequirements**

This example depends on the following data and software:

#### Installation of MPRAflow

Please install conda, the MPRAflow environment and clone the actual MPRAflow master branch. You will find more help under *Installation*.

#### Producing an association pickle

This workflow requires a python dictionary of candidate regulatory sequence (CRS) mapped to their barcodes in a pickle format. For this example the file can be generated using *Basic association workflow* or it can be found in the example folder of the GitHub repository.

#### **Design file**

File can be generated using the Basic association workflow or downloaded from the example folder.

#### Reads

There is one condition (HEPG2) with three technical replicates. Each replicate contains a forward (barcode-forward), reverse (barcode-reverse), and index (unique molecular identifier) read for DNA and RNA. These data must be downloaded. All data is publically available on the short read archive (SRA). We will use SRA-toolkit to obtain the data.

Note: You need 9 GB disk space to download the data and upwards of 50 GB to proccess it!

For large files and unstable internet connection we reccommend the comand *prefetch* from SRA tools before running *fastq-dump*. This command is much smarter in warnings when something went wrong.

conda install sra-tools cd Count\_Basic/data prefetch SRR10800881 SRR10800882 SRR10800883 SRR10800884 SRR10800885 SRR10800886 fastq-dump -gzip -split-files SRR10800986 cd ..

**Note:** Please be sure that all files are downloaded completely without errors! Depending on your internet connection this can take a while. If you just want some data to run MPRAflow you can just limit yourself to one condition and/or just one replicate.

#### With

tree data

the folder should look like this:

data

Here is an overview of the files:

Table 2: HEPG2 data			
Condition	GEO Ac-	SRA	SRA Runs
	cession	Accession	
HEPG2-DNA-1: HEPG2 DNA replicate 1	GSM423786	3 SRX7474781	SRR10800881
HEPG2-RNA-1: HEPG2 RNA replicate 1	GSM423786	4 SRX7474782	SRR10800882
HEPG2-DNA-2: HEPG2 DNA replicate 2	GSM423786	5 SRX7474783	SRR10800883
HEPG2-RNA-2: HEPG2 RNA replicate 2	GSM423786	6 SRX7474784	SRR10800884
HEPG2-DNA-3: HEPG2 DNA replicate 3	GSM423786	7 SRX7474785	SRR10800885
HEPG2-RNA-3: HEPG2 RNA replicate 3	GSM423786	8 SRX7474786	5 SRR10800886

## **MPRAflow**

Now we are close to starting MPRAflow and count the number of barcodes. But before we need to generate an environment csv file to tell nextflow the conditions, replicates and the corresponding reads.

#### Create experiment.csv

Our experiment file looks exactly like this:

```
Condition,Replicate,DNA_BC_F,DNA_UMI,DNA_BC_R,RNA_BC_F,RNA_UMI,RNA_BC_R
HEPG2,1,SRR10800881_1.fastq.gz,SRR10800881_2.fastq.gz,SRR10800881_3.fastq.gz,
→SRR10800882_1.fastq.gz,SRR10800882_2.fastq.gz,SRR10800882_3.fastq.gz
```

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```
HEPG2,2,SRR10800883_1.fastq.gz,SRR10800883_2.fastq.gz,SRR10800883_3.fastq.gz,

→SRR10800884_1.fastq.gz,SRR10800884_2.fastq.gz,SRR10800884_3.fastq.gz

HEPG2,3,SRR10800885_1.fastq.gz,SRR10800885_2.fastq.gz,SRR10800885_3.fastq.gz,

→SRR10800886_1.fastq.gz,SRR10800886_2.fastq.gz,SRR10800886_3.fastq.gz
```

Save it into the Count\_Basic/data folder under experiment.csv.

#### **Run nextflow**

Now we have everything at hand to run the count MPRAflow pipeline. Therefore we have to be in the cloned MPRAflow folder. But we will change the working and output directory to the Count\_Basic folder. The MPRAflow count command is:

```
cd <path/to/MPRAflow>/MPRAflow
conda activate MPRAflow
nextflow run count.nf -w <path/to/Basic>/Count_Basic/work --experiment-file "<path/to/
Basic>/Count_Basic/data/experiment.csv" --dir "<path/to/Basic>/Count_Basic/data" --
outdir "<path/to/Basic>/Count_Basic/output" --design "<path/to/design/fasta>/design.
ifa" --association "<path/to/association/pickle>/SRR10800986_filtered_coords_to_
barcodes.pickle"
```

Note: Please check your conf/cluster.config file if it is correctly configured (e.g. with your SGE cluster commands).

If everything works fine the following 5 processes will run: create\_BAM (make idx) raw\_counts, filter\_counts, final\_counts, dna\_rna\_merge\_counts, calc\_correlations, make\_master\_tables.

#### Results

All output files will be in the Count\_Basic/output folder.

We expect the program to output the following status when complete:

```
start analysis
executor > sge (32)
[23/09474b] process > create BAM (make idx)
                                              [100%] 6 of 6 ✓
                                              [100%] 6 of 6 ✓
[Of/4ee034] process > raw_counts (6)
[01/6ac02f] process > filter_counts (6)
                                              [100%] 6 of 6 ✓
[4f/b23748] process > final_counts (6)
                                              [100%] 6 of 6 🗸
[86/4ded79] process > dna_rna_merge_counts (3) [100%] 3 of 3 ✓
                                              [100%] 3 of 3 ✓
[29/0813f8] process > dna_rna_merge (3)
[1d/4e7d56] process > calc_correlations (1)
                                              [100%] 1 of 1 ✓
[9c/4714cb] process > make master tables (1)
                                              [100%] 1 of 1 ✓
Completed at: 07-Jan-2020 04:29:07
Duration : 11h 28m 5s
CPU hours : 41.5
Succeeded : 32
```

## 3.1.11 Count for Saturation Mutagenesis of the TERT promoter

This example runs the count workflow on saturation mutagenesis data of the TERT promoter from Kircher et al. 2019. It will only do the count part. For unraveling single variant effects please go to the example: *Saturation mutagenesis of the TERT promoter*. The same saturation mutagenesis library was used in four different experiments. We will focus only on the experiments in HEK293T and in glioblastoma SF7996 (GBM) cells (two different conditions).

#### **Prerequirements**

This example depends on the following data and software:

#### Installation of MPRAflow

Please install conda, the MPRAflow environment and clone the actual MPRAflow master branch. You will find more help under *Installation*.

#### Reads

We have two conditions (HEK293T and GBM cells). Each of them has three technical replicates and one replicate exists of forward, reverse and index reads for DNA and RNA. These data has to be downloaded. All data is public available on the short read archive (SRA). We will use the SRA-tools to download the reads.

Note: You need 16 GB disk space to download the data!

```
conda install sra-tools
mkdir -p Count_TERT/data
cd Count_TERT/data
fastq-dump --gzip --split-files SRR8647059 SRR8647060 SRR8647061 SRR8647062_
→SRR8647063 SRR8647064 SRR8647119 SRR8647120 SRR8647121 SRR8647122 SRR8647123_
→SRR8647124 SRR8647125 SRR8647126 SRR8647127 SRR8647128 SRR8647129 SRR8647130
cd ..
```

**Note:** Please be sure that all files are downloaded completely without errors! Depending on your internet connection this can take a while. If you just want some data to run MPRAflow you can just limit yourself to one condition and/or just one replicate.

#### With

tree data

the folder should look like this:

```
data

SRR8647059_1.fastq.gz

SRR8647059_2.fastq.gz

SRR8647059_3.fastq.gz

SRR8647060_1.fastq.gz

SRR8647060_2.fastq.gz

SRR8647060_3.fastq.gz
```

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<pre>SRR8647061_1.fastq.gz SRR8647061_2.fastq.gz SRR8647062_1.fastq.gz SRR8647062_2.fastq.gz SRR8647063_1.fastq.gz SRR8647063_2.fastq.gz SRR8647064_1.fastq.gz SRR8647064_2.fastq.gz SRR8647064_3.fastq.gz SRR8647119_1.fastq.gz SRR8647119_1.fastq.gz SRR8647120_1.fastq.gz SRR8647120_3.fastq.gz SRR8647120_3.fastq.gz SRR8647120_4.fastq.gz SRR8647121_1.fastq.gz SRR8647122_1.fastq.gz SRR8647122_1.fastq.gz SRR8647122_1.fastq.gz SRR8647122_1.fastq.gz SRR8647122_1.fastq.gz SRR8647122_3.fastq.gz SRR8647122_3.fastq.gz SRR8647122_4.fastq.gz SRR8647123_1.fastq.gz SRR8647124_1.fastq.gz SRR8647124_1.fastq.gz SRR8647125_1.fastq.gz SRR8647125_1.fastq.gz SRR8647126_1.fastq.gz SRR8647130_1.fastq.gz SRR8647130_1.fastq.gz SRR8647130_1.fastq.gz SRR8647130_1.fastq.gz SRR8647130_1.fastq.gz SRR8647130_1.fastq.gz SRR8647130_1.fast</pre>				
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<ul> <li>SRR8647129_3.fastq.gz</li> <li>SRR8647130_1.fastq.gz</li> <li>SRR8647130_2.fastq.gz</li> <li>SRR8647130_3.fastq.gz</li> <li>SRR8647130_4.fastq.gz</li> </ul>				
<pre>SRR8647130_1.fastq.gz SRR8647130_2.fastq.gz SRR8647130_3.fastq.gz SRR8647130_4.fastq.gz</pre>			SKK864/129_2.fastq.gz	
SRR8647130_2.fastq.gz SRR8647130_3.fastq.gz SRR8647130_4.fastq.gz			SRR864/129_3.fastq.gz	
SRR8647130_3.fastq.gz SRR8647130_4.fastq.gz			SRR8647130_1.fastq.gz	
SRR8647130_4.fastq.gz				
0 directories, 60 files		L	SRR8647130_4.fastq.gz	
0 directories, 60 files				
	(	) di	rectories, 60 files	

Here is an overview of the files:

Table 5.	TERT data
Condition	GEO Ac- SRA SRA Runs
	cession Accession
TERT-GBM-DNA-1: TERT-GBM transfection DNA	GSM3604284 SRX5444854 SRR8647059
replicate 1	
TERT-GBM-DNA-2: TERT-GBM transfection DNA	GSM3604285 SRX5444855 SRR8647060
replicate 2	
TERT-GBM-DNA-3: TERT-GBM transfection DNA	GSM3604286 SRX5444856 SRR8647061
replicate 3	
TERT-GBM-RNA-1: TERT-GBM transfection RNA	GSM3604287 SRX5444857 SRR8647062
replicate 1	
TERT-GBM-RNA-2: TERT-GBM transfection RNA	GSM3604288 SRX5444858 SRR8647063
replicate 2	
TERT-GBM-RNA-3: TERT-GBM transfection RNA	GSM3604289 SRX5444859 SRR8647064
replicate 3	
TERT-HEK-DNA-1: TERT-HEK transfection DNA	GSM3604302 SRX5444888 SRR8647119,
replicate 1	SRR8647120
TERT-HEK-DNA-2: TERT-HEK transfection DNA	GSM3604303 SRX5444889 SRR8647121,
replicate 2	SRR8647122
TERT-HEK-DNA-3: TERT-HEK transfection DNA	GSM3604304 SRX5444890 SRR8647123,
replicate 3	SRR8647124
TERT-HEK-RNA-1: TERT-HEK transfection RNA	GSM3604305 SRX5444891 SRR8647125,
replicate 1	SRR8647126
TERT-HEK-RNA-2: TERT-HEK transfection RNA	GSM3604306 SRX5444892 SRR8647127,
replicate 2	SRR8647128
TERT-HEK-RNA-3: TERT-HEK transfection RNA	GSM3604307 SRX5444893 SRR8647129,
replicate 3	SRR8647130

Table 3: TERT data

**Note:** The runs SRR8647120, SRR8647122, SRR8647124, SRR8647126, SRR8647128, and SRR8647130 have two index reads (forward and reverse). In the publication by Kircher et al. 2019 merging and trimming is used to combine them. For simplification we will discard the reverse index reads: rm data/\*\_4.fastq.gz

Also two different sequencing runs where made in condition TERT-HEK using the same library. Therefore, we have to process both runs together. So we will combine the reads. But in this csase bot sequencing runs have different read lengths (20 and 50 bp). Different read length will make false assumption when merging paired-end reads. Therefore we cut them down to 20 bp.

```
for i in 1 2 3; do
    zcat data/{SRR8647119,SRR8647120}_$i.fastq.gz | cut -c 1-20 | gzip -c > data/
    SRR8647119_SRR8647120_$i.fastq.gz;
    zcat data/{SRR8647121,SRR8647122}_$i.fastq.gz | cut -c 1-20 | gzip -c > data/
    SRR8647121_SRR8647122,$i.fastq.gz;
    zcat data/{SRR8647123,SRR8647124}_$i.fastq.gz | cut -c 1-20 | gzip -c > data/
    SRR8647123_SRR8647124,$i.fastq.gz;
    zcat data/{SRR8647125,SRR8647126}_$i.fastq.gz | cut -c 1-20 | gzip -c > data/
    SRR8647125_SRR8647125,SRR8647126}_$i.fastq.gz | cut -c 1-20 | gzip -c > data/
    SRR8647125_SRR8647126,$i.fastq.gz;
    zcat data/{SRR8647126,SRR8647126}_$i.fastq.gz | cut -c 1-20 | gzip -c > data/
    SRR8647125_SRR8647126,$i.fastq.gz;
    zcat data/{SRR8647126,$i.fastq.gz;
    zcat data/{SRR8647127,SRR8647128}_$i.fastq.gz | cut -c 1-20 | gzip -c > data/
    SRR8647127_SRR8647128,$i.fastq.gz;
    zcat data/{SRR8647127,SRR8647128}_$i.fastq.gz;
    zcat data/{SRR8647127,SRR8647128}_$i.fastq.gz;
    zcat data/{SRR8647127_SRR8647128}_$i.fastq.gz;
    zcat data/{SRR8647127_SRR8647128}_$i.fastq.gz;
    zcat data/{SRR8647127_SRR8647128}_$i.fastq.gz;
    zcat data/{SRR8647127_SRR8647128}_$i.fastq.gz;
    zcat data/{SRR8647127_SRR8647128}_$i.fastq.gz;
    zcat data/{SRR8647127_SRR8647128}_$i.fastq.gz;
    zcat data/{SRR8647127_SRR8647128_$i.fastq.gz;
    zcat data/{SRR8647127_SRR8647128_$i.fastq.gz;
    zcat data/{SRR8647127_SRR8647128_$i.fastq.gz;
    zcat data/{SRR8647127_SRR8647128_$i.fastq.gz;
    zcat data/SRR8647128_$i.fastq.gz;
    zcat data/SRR8647128_$i.fastq.
```

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```
zcat data/{SRR8647129,SRR8647130}_$i.fastq.gz | cut -c 1-20 | gzip -c > data/

→SRR8647129_SRR8647130_$i.fastq.gz;

done
```

**Note:** If you combine multiple sequence runs (e.g. you need more reads) you have to combine the reads before. Otherwise barcodes with the same UMI can be count twice. But it is important that all read lengths are the same. The easiest workaround is to cut them down to the minimum length. If you have a different library (but with the same barcodes) you should run the count utility with both runs separately using different conditions. Later you have to combine the final counts of both conditions.

#### **MPRAflow**

Now we are close to start MPRAflow and count the number of barcodes. But before we need to generate an environment.csv file to tell nextflow the conditions, replicates and the corresponding reads.

#### Create experiment.csv

Our experiment file looks exactly like this:

```
Condition, Replicate, DNA_BC_F, DNA_UMI, DNA_BC_R, RNA_BC_F, RNA_UMI, RNA_BC_R
TERT-GBM,1,SRR8647059_1.fastq.gz,SRR8647059_3.fastq.gz,SRR8647059_2.fastq.gz,
→SRR8647062_1.fastq.gz,SRR8647062_3.fastq.gz,SRR8647062_2.fastq.gz
TERT-GBM, 2, SRR8647060_1.fastq.gz, SRR8647060_3.fastq.gz, SRR8647060_2.fastq.gz,
→SRR8647063_1.fastq.qz,SRR8647063_3.fastq.qz,SRR8647063_2.fastq.qz
TERT-GBM, 3, SRR8647061_1.fastq.gz, SRR8647061_3.fastq.gz, SRR8647061_2.fastq.gz,
→SRR8647064_1.fastq.gz,SRR8647064_3.fastq.gz,SRR8647064_2.fastq.gz
TERT-HEK,1,SRR8647119_SRR8647120_1.fastq.gz,SRR8647119_SRR8647120_3.fastq.gz,
→SRR8647119_SRR8647120_2.fastq.gz,SRR8647125_SRR8647126_1.fastq.gz,SRR8647125_
→SRR8647126_3.fastq.gz,SRR8647125_SRR8647126_2.fastq.gz
TERT-HEK,2,SRR8647121_SRR8647122_1.fastq.gz,SRR8647121_SRR8647122_3.fastq.gz,
→SRR8647121_SRR8647122_2.fastq.gz,SRR8647127_SRR8647128_1.fastq.gz,SRR8647127_
⇔SRR8647128_3.fastq.gz,SRR8647127_SRR8647128_2.fastq.gz
TERT-HEK, 3, SRR8647123_SRR8647124_1.fastq.gz, SRR8647123_SRR8647124_3.fastq.gz,
→SRR8647123_SRR8647124_2.fastq.qz,SRR8647129_SRR8647130_1.fastq.qz,SRR8647129_
→SRR8647130_3.fastq.gz,SRR8647129_SRR8647130_2.fastq.gz
```

Save it into the Count\_TERT/data folder under experiment.csv.

#### **Run nextflow**

Now we have everything at hand to run the count MPRAflow pipeline. Therefore we have to be in the cloned MPRAflow folder. But we will change the working and output directory to the Count\_TERT folder. For the TERT example the barcode length is 20 bp and the UMI length 10 bp. The MPRAflow count command is:

Note: Please check your conf/cluster.config file if it is correctly configured (e.g. with your SGE cluster commands).

If everything works fine the following 5 processes will run: create\_BAM (make idx) raw\_counts, filter\_counts, final\_counts, dna\_rna\_merge\_counts.

```
[fe/d8ac14] process > create_BAM (make idx) [100%] 12 of 12 ✓
[7d/b56129] process > raw_counts (12) [100%] 12 of 12 ✓
[06/2c938d] process > filter_counts (12) [100%] 12 of 12 ✓
[2d/ce1afe] process > final_counts (12) [100%] 12 of 12 ✓
[68/df8db0] process > dna_rna_merge_counts (6) [100%] 6 of 6 ✓
Completed at: 09-Jan-2020 15:38:32
Duration : 3h 45m 17s
CPU hours : 21.8
Succeeded : 54
```

#### **Results**

All needed output files will be in the Count\_TERT/output folder. In this tutorial we are only interested in the counts per barcode, because we can use these outputs in the *Saturation mutagenesis of the TERT promoter* tutorial.

```
cd <path/to/TERT>/Count_TERT
tree output -P "*[123]_counts.tsv.gz"
```

```
output
   - TERT-GBM
       - 1
         L____ TERT-GBM_1_counts.tsv.gz
        2
         - TERT-GBM_2_counts.tsv.gz
        3
         L

    TERT-GBM_3_counts.tsv.gz

  - TERT-HEK
       - 1
        L____ TERT-HEK_1_counts.tsv.gz
       - 2
        ____ TERT-HEK_2_counts.tsv.gz
        3
         L____ TERT-HEK_3_counts.tsv.gz
8 directories, 6 files
```

The count files are tab separated and contain the barcode, the number number of unique UMI DNA counts and the umber of unique RNA counts. E.g. this is an example count file:

zcat output/TERT-GBM/1/TERT-GBM\_1\_counts.tsv.gz | head

(continues on next page)

ΑΑΑΑΑΑΑΑΑΑΑΤΤ	1	2
ААААААААААААТGA	3	1
AAAAAAAAAAAATGG	1	3
АААААААААААТТА	1	2
1		

# 3.1.12 Association for Saturation mutagenesis of TERT example

This example runs the association saturation mutagenesis workflow of the TERT promoter from Kircher et al. 2019. The same saturation mutagenesis library was used in four different experiments.

### Prerequirements

This example depends on the following data and software:

### Installation of MPRAflow

Please install conda, the MPRAflow environment and clone the actual MPRAflow master branch. You will find more help under *Installation*.

#### **Reference file**

To know where to map to we have to use a reference sequence. Here we will download the used reference sequence of TERT. It was generated by sanger sequencing of the original template (before error-prone PCR).

### Reads

There is one set of association sequencing for this data, which contains a forward, reverse, and index (barcode) reads. Forward and reverse contains the reference sequence with mutations. The barcode read is the associated barcode with the sequence. These data must be downloaded. All data is publically available on the short read archive (SRA). We will use SRA-toolkit to obtain the data.

Note: You need 2 GB disk space to download the data!

```
conda install sra-tools
cd satMut_assoc/data
prefetch SRR8646911
fastq-dump --gzip --split-files SRR8646911
cd ..
```

**Note:** Please be sure that all files are downloaded completely without errors! Depending on your internet connection this can take a while.

### With

tree data

#### the folder should look like this:

```
data/
SRR8646911_1.fastq.gz
SRR8646911_2.fastq.gz
SRR8646911_3.fastq.gz
TERT.fa
```

#### **MPRAflow**

Now we are ready to run MPRAflow for mapping reads, calling variants and associate them with barcodes.

#### **Run nextflow**

Now we have everything at hand to run the association saturation mutagenesis MPRAflow pipeline. Therefore we have to be in the cloned MPRAflow folder. But we will change the working and output directory to the satMut\_assoc folder. The MPRAflow association saturation mutagenesis command is:

Note: Please check your conf/cluster.config file if it is correctly configured (e.g. with your SGE cluster commands).

If everything works fine the following 10 processes will run: clean\_design create\_BWA\_ref, get\_name, create\_BAM, collect\_chunks, PE\_mapping, get\_count, extract\_reads, call\_variants, and combine\_variants.

```
[100%] 1 of 1, cached: 1 🗸
[e4/cf3353] process > clean_design (count)
[70/31c1b2] process > create_BWA_ref (make ref) [100%] 1 of 1, cached: 1 ✔
[83/a75010] process > get_name
                                                [100%] 1 of 1, cached: 1 ✓
                                                [100%] 28 of 28, cached: 28 ✓
[4e/8bd490] process > create_BAM (28)
                                                [100%] 1 of 1, cached: 1 ✓
[0b/ff7cd0] process > collect_chunks
[7c/64c374] process > PE_mapping (align)
                                               [100%] 1 of 1, cached: 1 ✓
[f3/fa5fed] process > get_count
                                                [100%] 1 of 1, cached: 1 ✓
[a5/d3aac2] process > extract_reads
                                               [100%] 1 of 1, cached: 1 ✔
                                               [100%] 1024 of 1024, cached: 1024 🗸
[67/70c6e0] process > call_variants (1024)
                                                [100%] 1 of 1 ✓
[85/2cb7af] process > combine_variants
```

```
Completed at: 26-März-2021 08:43:10
Duration : 22m 53s
CPU hours : 43.0 (100% cached)
Succeeded : 1
Cached : 1'059
```

### **Results**

All needed output files will be in the satMut\_assoc/output/TERT/ folder.

## 3.1.13 Saturation mutagenesis of the TERT promoter

This example runs the saturation mutagenesis workflow on saturation mutagenesis data of the TERT promoter from Kircher et al. 2019. The same saturation mutagenesis library was used in four different experiments. We will use the experiments in HEK293T and in glioblastoma SF7996 (GBM) cells in this workflow to see differences between the two cell lines (conditions).

### **Prerequirements**

This example depends on the following data and software:

### Installation of MPRAflow

Please install conda, the MPRAflow environment and clone the actual MPRAflow master branch. You will find more help under *Installation*.

### **Assignment file**

This file is a tab separated files that assigns variants to barcodes. We will create a new working folder and download the file into it

```
mkdir -p SatMut_TERT/data
cd SatMut_TERT/data
wget http https://github.com/shendurelab/MPRAflow/raw/master/examples/
$\implies$ saturationMutagenesis/TERT.variants.txt.gz
cd ..
It is also possible to get using the workflow :ref:`Association for Saturation_
$\implies$ mutagenesis of TERT example`.
```

### **Count tables**

We need the count tables of the count workflow. Please go to the *Count for Saturation Mutagenesis of the TERT promoter* and run it first. Afterwards copy the count tables into the data folder or use symbolic links:

ln -s ../Count\_TERT/output/TERT-GBM/\*/TERT-GBM\_{1,2,3}\_counts.tsv.gz data/ ln -s ../Count\_TERT/output/TERT-HEK/\*/TERT-HEK\_{1,2,3}\_counts.tsv.gz data/ Now the data folder should have the following files:

tree data

data
 TERT-GBM\_1\_counts.tsv.gz
 TERT-GBM\_2\_counts.tsv.gz
 TERT-GBM\_3\_counts.tsv.gz
 TERT-HEK\_1\_counts.tsv.gz
 TERT-HEK\_2\_counts.tsv.gz
 TERT-HEK\_3\_counts.tsv.gz
 TERT.variants.txt.gz
0 directories, 7 files

### **MPRAflow**

Now we are close to start MPRAflow and find out individual variant effects. But before we need to generate an environment.csv file to tell nextflow the conditions, replicates and the count files.

#### Create environment.csv

Our experiment file looks exactly like this:

```
Condition, Replicate, COUNTS
TERT-GBM, 1, TERT-GBM_1_counts.tsv.gz
TERT-GBM, 2, TERT-GBM_2_counts.tsv.gz
TERT-GBM, 3, TERT-GBM_3_counts.tsv.gz
TERT-HEK, 1, TERT-HEK_1_counts.tsv.gz
TERT-HEK, 2, TERT-HEK_2_counts.tsv.gz
TERT-HEK, 3, TERT-HEK_3_counts.tsv.gz
```

Save it into the SatMut\_TERT/data folder under experiment.csv.

### **Run nextflow**

Now we have everything at hand to run the saturation mutagenesis MPRAflow pipeline. Therefore we have to be in the cloned MPRAflow folder. But we will change the working and output directory to the SatMut\_TERT folder. The MPRAflow saturation mutagenesis command is:

Note: Please check your conf/cluster.config file if it is correctly configured (e.g. with your SGE cluster commands).

If everything works fine the following 11 processes will run: calc\_assign\_variantMatrix calc\_assign\_variantMatrixWith1bpDel, fitModel, summarizeVariantMatrix,

statsWithCoefficient, plotCorrelation, plotStatsWithCoefficient, fitModelCombined, combinedStats, statsWithCoefficientCombined, and plotStatsWithCoefficientCombined.

[3c/835d00] process	> calc_assign_variantMatrix (1)	[100%] 6 of 6 🗸			
[7a/887135] process	<pre>&gt; calc_assign_variantMatrixWith1bpDel (1)</pre>	[100%] 6 of 6 🖌			
[ca/a90b00] process	> fitModel (8)	[100%] 12 of 12 🗸			
[67/3a3e8a] process	> summarizeVariantMatrix (12)	[100%] 12 of 12 🗸			
[56/846670] process	> statsWithCoefficient (12)	[100%] 12 of 12 🗸			
[74/466bfb] process	> plotCorrelation (1)	[100%] 12 of 12 🗸			
[a5/baf1ef] process	> plotStatsWithCoefficient (12)	[100%] 12 of 12 🗸			
[ac/d38378] process	> fitModelCombined (3)	[100%] 4 of 4 🗸			
[0b/600d8b] process	> combinedStats (2)	[100%] 4 of 4 🗸			
[32/80f6a6] process	> statsWithCoefficientCombined (2)	[100%] 4 of 4 🗸			
[2f/817e76] process	> plotStatsWithCoefficientCombined (1)	[100%] 4 of 4 🗸			
Completed at: 07-Jan-2020 11:31:00					
Duration : 22m 41	S				
CPU hours : 1.0					
Succeeded : 88					

## Results

All needed output files will be in the SatMut\_TERT/output folder.

## 3.1.14 Frequently Asked Questions

If you have more question please write us a ticket on github.

```
MPRAflow is not able to create a Conda environment If you get a message like:
```

```
Caused by: java.lang.IllegalStateException: Failed to create Conda environment
command: conda env create --prefix /home/user/MPRAflow/work/conda/mpraflow_py27-
→a6601743cee3b1029d4f3c810b7ebf02 --file /home/user/MPRAflow/conf/mpraflow_py27.
→yml`
```

Try to run conda separately using:

Afterwards try MPRAflow again. Please be sure that you are connected to the internet!

Can I use STARR-seq with MPRAflow? No. For more details have a look at this comment.

The pipeline is giving an error "QXcbConnection: Could not connect to display" and won't run. How can I fix this? Depending on your cluser configuration, the QT\_QPA\_PLATFORM variable may not be compatible with the plotting scripts in the pipeline. To resolve this error, add QT\_QPA\_PLATFORM='offscreen' to your .bash\_profile.

## 3.1.15 Contributing

Contributions are welcome, and they are greatly appreciated! Every little bit helps, and credit will always be given.

You can contribute in many ways:

## **Types of Contributions**

## **Report Bugs**

Report bugs at https://github.com/shendurelab/MPRAflow/issues

If you are reporting a bug, please include:

- Your operating system name and version.
- Any details about your local setup that might be helpful in troubleshooting.
- Detailed steps to reproduce the bug.

## **Fix Bugs**

Look through the Github issues for bugs. If you want to start working on a bug then please write short message on the issue tracker to prevent duplicate work.

### **Implement Features**

Look through the Github issues for features. If you want to start working on an issue then please write short message on the issue tracker to prevent duplicate work.

### Write Documentation

MPRAflow could always use more documentation, even on the web in blog posts, articles, and such.

MPRAflow uses Sphinx for the user documentation (that you are currently reading). See *doc\_guidelines* on how the documentation reStructuredText is used. See *doc\_setup* on creating a local setup for building the documentation.

### **Submit Feedback**

The best way to send feedback is to file an issue at https://github.com/shendurelab/MPRAflow/issues

If you are proposing a feature:

- Explain in detail how it would work.
- Keep the scope as narrow as possible, to make it easier to implement.
- Remember that this is a volunteer-driven project, and that contributions are welcome :)

### **Documentation Guidelines**

For the documentation, please adhere to the following guidelines:

- Put each sentence on its own line, this makes tracking changes through Git SCM easier.
- Provide hyperlink targets, at least for the first two section levels.
- Use the section structure from below.

```
.. heading_1:
_____
Heading 1
_____
.. heading_2:
Heading 2
_____
.. heading_3:
Heading 3
_____
.. heading_4:
Heading 4
 ____
.. heading_5:
Heading 5
.. heading_6:
Heading 6
: : : : : : : : :
```

## **Documentation Setup**

For building the documentation, you have to install the Python program Sphinx. This is best done in a virtual environment. We created a conda environment to work with the actual documentation.

Use the following steps for installing Sphinx and the dependencies for building the MPRAflow documentation:

```
cd MPRAflow/docs
conda env create -f environment.yml -n sphinx
conda activate sphinx
```

Use the following for building the documentation. The first two lines is only required for loading the virtualenv. Afterwards, you can always use make html for building.

```
cd MPRAflow/docs
conda activate sphinx
make html # rebuild for changed files only
make clean && make html # force rebuild
```

### **Get Started!**

Ready to contribute? First, create your Documentation development setup.

- 1. Fork the MPRAflow repo on GitHub.
- 2. Clone your fork locally:

git clone git@github.com:your\_name\_here/MPRAflow.git

3. Create a branch for local development:

git checkout -b name-of-your-bugfix-**or**-feature

Now you can make your changes locally.

5. When you're done making your changes, make sure that nextflow runs properly For nextflow:

```
nextflow run <your_nextflow_pipeline>
```

For documentation:

cd docs make clean && make html

6. Commit your changes and push your branch to GitHub:

```
git add <your_new_file> # or git stage <your_edited_file>
git commit -m "Your detailed description of your changes."
git push origin name-of-your-bugfix-or-feature
```

7. Submit a pull request through the GitHub website.

### **Pull Request Guidelines**

Before you submit a pull request, check that it meets these guidelines:

- 1. The pull request should include tests.
- 2. If the pull request adds functionality, the docs should be updated.

## 3.1.16 Authors

in alphabetical order

- Vikram Agarwal
- Tal Ashuach
- Gracie Gordon
- Martin Kircher
- Max Schubach
- Sean Whalen

## Contributors

• You name here

# 3.1.17 History

```
# MPRAflow Changelog
## v2.3.4
### count.nf
* increasing speed of `src/merge_label.py` speed for larger MPRAs.
## v2.3.3
### association.nf
* Bugfix in `src/nf_ori_map_barcodes.py` script (see issue #41). Now >= mapq instead_
\rightarrow of > mapq is used.
## v2.3.2
### count.nf
* Bugfix merge_all.py script (see issue #55)
## v2.3.1
### association.nf
* Bugfix empty design file (see issue #45)
## v2.3
### global changes
* Correcting typos in documentation
* adding new process label `highmem` to `conf/cluster.config`
### association_saturationMutagenesis.nf
New association saturation mutagenesis workflow. This workflow is about assocation,
-variant calls with barcodes. Variants are introduced by an error-prone PCR. The
-workflow takes the sequencing of the region, with barcodes in index read and the
-reference sequence and maps the reads to the reference, calls variants and
\rightarrowassociates them with the corresponding barcode. it is a pre-step of.
⇒`saturationMutagenesis.nf`.
### count.nf
* using BC threshold input for `plot_perInsertCounts_correlation.R` instead of hard-
-coded. Modify process `calc_correlations` to use the new input theshold.
### association.nf
```

```
* Remove "windows" characters in design fasta
## v2.2
No workflow changes. Only a few fixes and some restructuring of configs. Using
→nextflow version 20.01 now!
### global changes
* nextflow version 20.01 is needed because of multiMap() function
* introducing new config file `conf/global.config` with global variables like the min.
\hookrightarrow required nextflow version and the actual MPRAflow version.
* moving cluster config to a sepArate file: `conf/cluster.config`. Try to adapt the...
-times to the sort and longtime labels. Modify SLURM queue to SLUM not SGE options.
* improved documentation
### saturationMutagenesis.nf
* Bugfix of default out dir. It was not set to `params.outdir = "outs"` so it tries.
→to create a folder `null`. Now in `params.outdir`
* removing default `params.version` and `params.nf_required_version`. Now in `conf/
→global.config`
* Catching cases when barcode/p-value filtering produces 0 variants
* Change update depricated fork method. Now works with nextflow 20.01
### count.nf
* removing default `params.version`, `params.nf_required_version` and `params.outdir`.
→ Now in `conf/global.config`.
### association.nf
* removing default `params.version`, `params.nf_required_version` and `params.outdir`.
→ Now in `conf/global.config`.
## v2.1
Initial MPRAflow version for publication.
```

## 3.1.18 MPRAflow License

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# 3.1.19 Documentation TODO-list

Todo: describe summarizeVariantMatrix

(The original entry is located in /home/docs/checkouts/readthedocs.org/user\_builds/mpraflow/checkouts/v2.3.4/docs/saturation\_mutager line 95.)

Todo: describe combinedStats

(The original entry is located in /home/docs/checkouts/readthedocs.org/user\_builds/mpraflow/checkouts/v2.3.4/docs/saturation\_mutager line 105.)

### Todo: Describe SatMut output files

(The original entry is located in /home/docs/checkouts/readthedocs.org/user\_builds/mpraflow/checkouts/v2.3.4/docs/saturation\_mutager line 120.)