#### **User Manual**

#### Introduction

rMATS-DVR is a convenient and user-friendly program to streamline the discovery of DVRs (Differential Variants in RNA) between two RNA-seq sample groups with replicates. rMATS-DVR combines a stringent GATK-based pipeline for calling SNVs including SNPs and RNA editing events in RNA-seq reads, with our rigorous rMATS statistical model for identifying differential isoform ratios using RNA-seq sequence count data with replicates.

## Availability

rMATS-DVR is a free software, which can be downloaded from https://github.com/Xinglab/rMATS-DVR.

# Requirements

- 1. Install Python 2.6.x or Python 2.7.x and corresponding versions of NumPy and SciPy.
  - 2. Install Java v1.8 (https://java.com/en/download/manual.jsp#lin).
  - 3. Add the Python and Java directories to the \$PATH environment variable.
  - 4. Download Picard v2.6.0 from

https://github.com/broadinstitute/picard/releases/tag/2.6.0.

- 5. Download GATK v3.6 from https://software.broadinstitute.org/gatk/download/.
- 6. Download SAMtools v1.3.1

(https://github.com/samtools/samtools/releases/tag/1.3.1) and add it to \$PATH environment variable.

Note: We have only tested rMATS-DVR in linux platform.

#### Installation

- 1. Create soft links of Picard and GATK into rMATS-DVR folder.
- cd rMATS-DVR
- In -s /path/to/picard/picard.jar ./
- In -s /path/to/GATK/GenomeAnalysisTK.jar ./
- 2. Then the source code can be directly called from Python.

## Required and optional external files

All the external files based on human hg19 genome can be downloaded from http://www.mimg.ucla.edu/faculty/xing/public data/rMATS-DVR/hg19 resource.tar.gz

To decompress and extract the files:

## tar -xvf hg19 resource.tar.gz

Alternatively, users can also prepare the external files under the following instructions:

1) Genome (required): we highly recommend the users use the genome sequence (together with a dictionary file such as "ucsc.hg19.dict" and index file such as "ucsc.hg19.fasta.fai") from GTAK bundle (https://software.broadinstitute.org/gatk/download/bundle). Alternatively, please follow the instructions in GATK (https://software.broadinstitute.org/gatk/guide/article?id=1204) to prepare the reference genome in proper format.

- 2) Known SNPs (optional): SNP annotation in VCF format. dbSNP annotation of human can be downloaded from GTAK bundle (<a href="https://software.broadinstitute.org/gatk/download/bundle">https://software.broadinstitute.org/gatk/download/bundle</a>). Alternatively, please follow the instructions in GATK (<a href="https://software.broadinstitute.org/gatk/guide/article?id=1204">https://software.broadinstitute.org/gatk/guide/article?id=1204</a>) to prepare the valid VCF file.
- 3) Known RNA editing sites (optional): table delimited txt file with the first two columns are chromosome and coordinates. The other columns are ignored. Header is optional. Users can download the file from RADAR dababase (http://rnaedit.com/download/).
- 4) Genome-wide repeat elements (optional): RepeatMasker Genomic Datasets downloaded from <a href="http://www.repeatmasker.org/genomicDatasets/RMGenomicDatasets.html">http://www.repeatmasker.org/genomicDatasets/RMGenomicDatasets.html</a>. For example: hg19.fa.out.gz
- 5) Gene annotation (optional): the gene annotation is in the GenePred (extended) format (see <a href="https://genome.ucsc.edu/FAQ/FAQformat.html">https://genome.ucsc.edu/FAQ/FAQformat.html</a> for detail). We recommend users to download it from UCSC. (<a href="http://hgdownload.soe.ucsc.edu/downloads.html">http://hgdownload.soe.ucsc.edu/downloads.html</a> ). For example, one can download hg19 RefSeq gene from: <a href="http://hgdownload.soe.ucsc.edu/goldenPath/hg19/database/refGene.txt.gz">http://hgdownload.soe.ucsc.edu/goldenPath/hg19/database/refGene.txt.gz</a>

### Run rMATS-DVR in one step

In one step mode, rMATS-DVR will first calibrate bam files one by one and then calculate Differential Variants of RNA using all samples. By the way, mapping with STAR is highly recommended.

Usage:

python rMATS-DVR.py --sample1 S1\_rep\_1.bam[,S1\_rep\_2.bam][,...,S1\_rep\_n.bam] --sample2 S2\_rep\_1.bam[,S2\_rep\_2.bam][,...,S2\_rep\_n.bam] --label S1,S2 --genome hg19.fa --output /Path/to/output/S1\_vs\_S2 [--known dbSNP147.vcf] [--editing RADAR2.txt] [--repeat repeats.txt] [--gene RefSeq.txt] [--minQ 20] [--minDP 5] [--thread 1] [--diff 0.0001] [--merge] [--ReadStranded] [--ReadPaired] [--skipBamCalibration] [--KeepTemp]

## Required Parameters:

-h, --help Show this help message and exit.

--sample1 <str> Bam (or sam) files of sample 1, replicates are separated by

comma.

--sample2 <str> Bam (or sam) files of sample 2, replicates are separated by

comma.

--label <str> Lable of sample 1 and sample 2, separated by comma, e.g.

Sample1,Sample2.

--output <str> Path and prefix of the output file.

--genome <str> Genome sequence in fasta format.

### Optional Parameters:

--known <str> Known SNPs in VCF format.

--editing <str> Known RNA editing sites.

--repeat <str> Repeat elements annotation.

--gene <str> Gene annotation.

--minQ <int> Minimum variant quality. [20]

--minDP <int> Minimum mean read coverage of both samples. [5]

--thread <int> Number of processors. [1]

--diff <float> Required level difference between the two samples. [0.0001]

--ReadStranded RNA-seq reads are Illumina strand-specific reads. Disable by

default.

--ReadPaired RNA-seq reads are paired-end reads. Disable by default.

--merge Merge the counts of all replicates. Enable by default when there

are less than 2 replicates in at least one sample group.

--skipBamCalibration Skip the step of calibrating bam files. Enable it when the input bam files have already been calibrated using bam\_calibration.py (see below). Disable by default.

--KeepTemp Keep the temporary files. Disable by default.

# Run rMATS-DVR in two steps

When there are a large number of replicates, one step mode, which calibrate bam files one by one, may take long time. In these cases, we recommend users to run bam calibration for all bam files in parallel at the first step. Then the users can run rMATS-DVR.py with --skipBamCalibration.

### Usage:

python bam\_calibration.py --bam sample.bam --output /Path/to/output/prefix --genome hg19.fa [--known dbSNP147.vcf] [--KeepTemp]

#### Parameters:

-h, --help Show this help message and exit.

--bam <str> Input bam (or sam) file.

--output <str> Path and prefix of the output file.

--genome <str> Genome sequence in fasta format.

--known <str> [Optional] Known SNPs in VCF format.

--KeepTemp [Optional] Keep the temporary files. Disable by default.

## Output

The final output files are in "Prefix\_rMATS-DVR\_results" folder, including "rMATS-DVR\_Result.txt" and "rMATS-DVR\_Result\_summary.txt".

"rMATS-DVR\_Result.txt" provides the variant information, read counts, P value, FDR, gene location, and multiple annotations based on known databases. "rMATS-DVR\_Result\_summary.txt" summarizes the frequencies of all types of total variants and DVRs respectively, they are also substratified into known SNPs, known RNA editing sites, and novel variants. All other files are temporary files.

# 1) rMATS-DVR\_Result.txt

Chroms: Chromosome of variant.

Site: 1-based coordinates of variant.

Ref\_allele: reference allele. Alt allele: alternative allele.

RNA-seqStrand: RNA strand from which the RNA-seq reads are originated. Only valid when --ReadStranded is applied in rMATS-DVR.

Variant quality: GATK reported Phred-scaled quality score of variant.

Sample1\_Alt: read counts of alternative allele in sample 1, replicates are separated by comma.

Sample1\_Ref: read counts of reference allele in sample 1, replicates are separated by comma.

Sample2\_Alt: read counts of alternative allele in sample 2, replicates are separated by comma.

Sample2\_Ref: read counts of reference allele in sample 2, replicates are separated by comma.

Pvalue: P value of differential allelic count ratios between two sample groups.

FDR: Benjamini-Hochberg corrected FDR of the above P value.

Sample1\_Alt\_allele\_fraction: fraction of alternative allele counts in sample 1, replicates are separated by comma.

Sample2\_Alt\_allele\_fraction: fraction of alternative allele counts in sample 2, replicates are separated by comma.

Alt\_allele\_fraction\_diff: average (Sample1\_Alt\_allele\_fraction) - average (Sample2 Alt allele fraction).

Genename: name of the gene in which the variant is located.

Strand: strand of the gene.

Ref\_onSense: reference allele on sense strand.

Alt onSense: alternative allele on sense strand.

Location: location of the variant on gene.

KnownSNP: rs ID of known SNP (dbSNP) hit.

KnownRNAediting: boolean variable to show whether the variant has a hit in known RNA editing database.

RepeatName: name of repeat element which covers the variant. RepeatName: family of repeat element which covers the variant.

## 2) rMATS-DVR Result summary.txt

Type (Ref-Alt) on sense strand: type of variants in the format of reference allelealternative allele on sense strand.

All Variants: frequency of each type of all called variants.

All DVRs (FDR<0.05): frequency of each type of all variants with FDR < 0.05.

SNP DVRs: frequency of each type of all known SNPs with FDR < 0.05.

RNA editing DVRs: frequency of each type of all known RNA editings with FDR < 0.05.

Novel DVRs: frequency of each type of all novel variants with FDR < 0.05.

### Run with the least input

For species without SNP or transcript annotation, users can run rMATS-DVR with only the RNA-seq alignments and the corresponding genome. In that case, rMAT-DVR still report the variant information and DVR information (column 1-15) in rMATS-DVR\_Result.txt. It is also helpful to provide a *de novo* transcript assembly in GenePred (extended) format as previously described.

#### Contacts

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If you find a bug or mistake in this project, we would like to know about it. Before you send us the bug report though, please check the following:

Are you using the latest version? The bug you found may already have been fixed. Check that your input is in the correct format and you have selected the correct options.

Please reduce your input to the smallest possible size that still produces the bug; we will need your input data to reproduce the problem, and the smaller you can make it, the easier it will be.

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