miRWoods Instruction Manual

 $\begin{array}{c} Author: \\ {\rm Jimmy~Bell} \end{array}$

June 23, 2019



Contents

1	Dependencies	2
2	Installation	2
3	Preparing Bam Files 3.1 Preparing Bam Files Using createBam.pl	
4	Creating a Bam File List	3
5	Creating a Config File	4
6	Running miRWoods 6.1 Running miRWoods using miRWoods.pl Script	
7	Viewing Output	5

1 Dependencies

miRWoods has the following dependencies:

- 1. bowtie
- 2. samtools
- 3. The Vienna RNA package with the RNA Perl Library
- 4. Cutadapt is used for adapter trimming and may be downloaded using the following command: pip3 install –user –upgrade cutadapt

The following perl libraries are required:

- 1. Bio::DB::Sam
- 2. Statistics::R
- 3. Math::CDF

you will also need to download R and install the following libraries:

- 1: ROCR
- 2: randomForest
- 3: mlbench

2 Installation

An install script is being worked on but not yet ready. For now do the following:

- 1. Add the Scripts directory from miRWoods into your path
- 2. Add the PerlModules directory to your PERL5LIB path

3 Preparing Bam Files

3.1 Preparing Bam Files Using createBam.pl

Example:

fastq-dump SRR326279

createBam.pl SRR326279.fastq ATCTCGTATGCCGTCTTCTGCTTGT 30 hg19

Directions:

createBam.pl is a script which runs each process described in section 3.2. createBam.pl is run in the following way:

createBam.pl (fastq file) (adaptor) (quality score) (bowtie index)

3.2 Preparing BamFiles without Using createBam.pl

Example:

fastq-dump SRR326279

 $cutadapt - e\ 0.2 - m\ 17 - q\ 10 - a\ ATCTCGTATGCCGTCTTCTGCTTGT\ SRR326279. fast q-o\ SRR326279_trimmed. fast q\ AvgQualityFilter. pl\ SRR326279_trimmed. fast q\ SRR326279_qualFilt. fast q\ 30$

bowtie -n 1 -e 50 -l 18 -m 10 -a -S -best -strata hg
19 SRR326279-qual Filt.fastq SRR326279.sam

addNHTags.pl SRR326279.sam SRR326279_NHTags.sam

 $trim Mir Reads.pl~SRR 326279_NH Tags.sam~SRR 326279_trimmed.sam$

samtools view -bS -o SRR326279_trimmed.bam SRR326279_trimmed.sam

samtools sort SRR326279_trimmed.bam SRR326279_sorted

Directions:

The following are instructions on a step by step process to creating the bam files. The user may use their own methods if they wish. If using your own method, please note that the bam file requires NH tags for each read showing the number of hits to the genome.

1. After the fastq file is downloaded reads are trimmed using cutadapt with the following options: cutadapt -e 0.2 -m 17 -q 10 -a $\langle adaptor \rangle \langle fastq file \rangle$ -o $\langle fastq output file \rangle$

Setting -e to 0.2 allows for 20% error in the adapter sequence to be cut. Setting -m to 17 removes all sequences less than 17 nucleotides in length. Setting -q to 10 removes any nucleotides from the 3 end with quality scores less than 10 prior to removing the adaptor. The -a option specifies the adaptor sequence.

- 2. Reads without an average quality score of 30 may be removed using the following command: fastqAvgQualityFilter.pl (input fastq\) (output fastq\) 30
- 3. Fastq files are aligned to the genome using bowtie with the options: bowtie -n 1 -e 50 -l 18 -m 10 -a -S -best -strata (bowtieIndex) (fastq file) (output sam)

Setting -l to 18 specifies a bowtie seed length of 18. Setting -n to 1 allows for 1 mismatch within the bowtie seed region. Setting -e to 50 allows for a total quality score of 50 for all mismatched nucleotides. Setting -m to 10 suppresses all mappings that hit more than 10 places in the genome. The -a options sets bowtie to report all valid alignments. The -S option sets bowtie to output the mapped reads in SAM format. The -best and -strata set bowtie to report all valid alignments based on the previously mentioned options.

- 4. A custom script is used to add NH tags to the sam file so that miRWoods will have information about the number of mappings for each read (Note: this script must be used prior to sorting): addNHTags.pl (input sam file) (output sam file)
- 5. If there are any mappings with several low quality mismatches on the end they may be trimmed down using the following script: trimMirReads.pl (input sam file) (output sam file)
- 6. The sam file is converted to a bam file using the following command: samtools view -bS -o (output bam file) (input sam file)
- 7. The bam file is sorted using: samtools sort (input bam file) (output prefix)
- 8. Finally, the bam file is indexed: samtools index (bam file)

4 Creating a Bam File List

A list of all bam files is used as input for miRWoods. The bam file list is a tab delimited file containing sample names followed by their location.

```
Example:
hsa_cellLines1
                      /\(\rangle \text{path to directory}\)/hsa_cellLines1.bam
hsa_cellLines2
                      /\(\rangle \text{path to directory}\)/hsa_cellLines2.bam
hsa_cellLines3
                      /\(\rangle \text{path to directory}\)/hsa_cellLines3.bam
                      /\langle path to directory \rangle / hsa_cellLines4.bam
hsa_cellLines4
hsa_cellLines5
                      /\(\rangle path to directory\)/hsa_cellLines5.bam
hsa_cellLines6
                      /\(\rangle\) path to directory\/hsa_cellLines6.bam
(see example_bam.txt)
```

Creating a Config File 5

miRWoods takes its inputs in the form of a config file. This is a tab delimited file with the name of each parameter and its value separated by an equal sign

For example:

```
/\(\rangle\) path to miRWoods Dir\/miRWoods/Scripts
scriptDir
hairpinRF
                          /\(\rangle\) path to miRWoods Dir\/miRWoods/Models/hairpinRF.model
productRF
                         /\(\rangle\) path to miRWoods Dir\(\rangle\)/miRWoods/Models/productRF.model
SizesFile
                          /\langle path \rangle / hg19.chrom.sizes
                     =
                          /\langle path to Genome Dir \rangle / hg19
genomeDir
                          /\(\rangle \text{path to Annotations}\)/hg19_ensembl.gff3
geneModels
mirbaseGff
                          hsa.gff3
bamListFile
                          bamlist.txt
minReadLength
                          14
                          28
maxReadLength
HPDVCutoff
                          0.28
ARPMCutoff
                          0.11
```

(see example_config.txt)

Running miRWoods 6

Running miRWoods using miRWoods.pl Script

The easiest way to run miRWoods is to give feed it a config file using the -L option. This method runs all the scripts in section 6.2.

```
Example:
miRWoods -L config.txt
```

6.2Running miRWoods using One Script in the Pipeline at a Time

The following are instructions on how to run each script in the miRWoods pipeline. This is not necessary unless you're troubleshooting an error.

Steps in the pipeline:

1. checkMiRBaseGff.pl checks the miRBase for issues that might affect miRWoods and in some cases creates a new miRBase file to run with miRWoods:

checkMiRBaseGff.pl (configFile)

- 2. countMappedReads.pl counts the number of read mappings in each bam file and creates a tab delimited file of library sizes: countMappedReads.pl ⟨bam file list⟩
- 3. printReadRegions.pl find all regions of contiguous read loci and prints them out as a bed file: printReadRegions.pl -L ⟨configFile⟩
- 4. extract ProductFeatures.pl groups products together and scores them: extract ProductFeatures.pl -L \langle configFile \rangle
- 5. generateProductTrainData.pl creates training data in order to create a model for the mature product random forest (MPRF). This is only needed if you wish to build a new model: generateProductTrainData.pl -L ⟨configFile⟩
- 6. evaluateProductsWithRF.pl This evaluate products based on the product random forest model. if the trainModels parameter in the config is set to 1 it will also create a new model. If you are training a new model be careful not to set productRF to one to don't wish to have overwritten: evaluateProductsWithRF.pl -L ⟨configFile⟩
- 7. processReadRegions.pl the products that passed the mature product random forest (MPRF), combines them with surrounding products to create hairpins, and then scores each hairpin: processReadRegions.pl -L ⟨configFile⟩
- 8. generateAnnotations.pl Hairpins are annotated using this Script. A later script will apply these annotations after the hairpin precursor random forest (HPRF) scores each hairpin: generateAnnotations.pl -L (configFile)
- 9. extractHairpinFeatures.pl Hairpin features are further process by this file. It also counts neighboring loci and includes this score as a feature called neighborCount: generateAnnotations.pl -L (configFile)
- 10. generate HairpinTrainData.pl creates training data in order to create a model for the hairpin precursor random forest (HPRF). This is only needed if you wish to build a new model: generate HairpinTrainData.pl -L \(\)configFile \(\)
- 11. evaluateHairpinsWithRF.pl evaluates hairpins based on the hairpin precursor random forest model. If the trainModels parameter in the config is set to 1 it will also create a new model. If you are training a new model be careful not to set productRF to one to don't wish to have overwritten:

evaluateHairpinsWithRF.pl -L (configFile)

12. printMiRWoodsPredictions.pl gathers and prints the final output files of miRWoods: printMiRWoodsPredictions.pl positivePredictions.gff positivePredictions_hairpins.txt positivePredictions_products.txt

7 Viewing Output

The final output from miRWoods is in two files:

- $1.\ \operatorname{predictedMiRs.gff}\ \text{-}\ \operatorname{a}\ \operatorname{gff}\ \operatorname{file}\ \operatorname{containing}\ \operatorname{the}\ \operatorname{hairpins}\ \operatorname{and}\ \operatorname{products}\ \operatorname{predicted}\ \operatorname{by}\ \operatorname{miRWoods}$
- $2.\ \mathrm{predictedMiRs_productInfo.txt}$ a tab delimited file containing read count information for each product. This file includes loop and antisense products.