A user's manual for miR-island

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OVERVIEW

Next-generation sequencing of small RNAs have provided rich information on microRNAs profile of various plant species. However, few computational tools are presently available to analyze these data effectively. MiR-island is a local tool that analyzes plant small RNAs and quantifies small RNA genes. In this study, miR-island performs unprecedented speed and efficient memory-usage in plant microRNA annotations.

SUMMARY OF MIR-ISLAND FUNCTIONS

Based on ultra-deep sampling of small RNA libraries by next generation sequencing, miR-island has a lot of advantages in annotation and quantification of plant miRNA genes. MiR-island can be used to identify miRNA genes in plant species with or without annotations. MiR-island can identify miRNA genes in an ultra-fast speed, no matter at which assemble level the reference sequences are. MiR-island is properly for miRNA quantification and differentially miRNA expression. MiR-island offers user-friendly tabular outputs and publication-ready results.

IMPLEMENTATION AND ALGORITHM

MIR-Island is documented by Perl (Perl 5.10 or later versions) and other fundamental packages from Perl library. All the scripts have been tested on two Linux platforms, Centos 6.4 and RedHat 5.4, and should work on similar systems that support Perl.

The core algorithm of miR-Island was developed by modifying miRDeep (Friedlander et al., 2008), which is based on a modified probabilistic model of miRNA biogenesis and a series of plant-specific filtering criteria (Meyers *et al.*, 2008).

LICENSE AND AVAILABILITY

MiR-Island is freely available under a GNU Public License (Version 3) at: http://www.ncrna.net/tools/mir-island.html and http://sourceforge.net/projects/mir-island/ The miR-island scripts, demos and user manual can be obtained from both web sites.

CONVENTIONS AND RECOMMENDATIONS

All command lines, filenames and directory names are in italic. The command lines are in blue background. The attached demo package, *miR-island_demo*, include the files by which users can reproduce every step and gain familiarity with miR-island. Please note that some of the intermediate files are not included due to the size of these files. Users are recommended to generate the files by themselves.

INSTALLATION OF THIRD PARTY SOFTWARE

Note: To reduce time consuming, multiple threads specificity is applied in miR-Island. Your perl version must be equal to or greater than 5.10. You can check the perl version by running perl -v, and see if it is possible to use multiple threads by running perl -V and looking at the Platform

section. If you have useithreads-define you can use the specificity.

Several dependencies are required to run miR-Island.

- 1: bowtie (used for aligning small RNA-seq to reference genome). The bowtie package can be downloaded from the site: http://sourceforge.net/projects/bowtie-bio/files/bowtie/. Our test version is 0.12.9.
- 2: samtools (tools for alignments in the SAM format). Samtools can be downloaded from the site: http://samtools.sourceforge.net/. Our test version is 0.1.19.
- 3: ViennaRNA (RNAfold is used for predicting hairpin structure). The Vienna RNA package is currently at http://www.tbi.univie.ac.at/RNA/. Our test version is 2.0.0.
- 4: bioperl (used to improve the operation efficiency). Users are recommended to install bioperl according to http://www.bioperl.org/wiki/Installing_BioPerl. Our test version is 1.6.901.
- 5: Bio::DB::Sam (used to handle the SAM/BAM format alignment). When you install Bio::DB::Sam module, you should depress the tar.gz of samtools and change directory to the depressed folder, then run "make CXXFLAGS=-fPIC CFLAGS=-fPIC CPPFLAGS=-fPIC". At last, you install Bio::DB::Sam module.

INSTALL MIR-ISLAND

 $Download \ the \ .tar.gz \ file \ from \ \underline{http://www.ncrna.net/tools/miR-island_v1.0.tar.gz} \ and \ unpack.$

Then put the scripts into your PATH. This is done by typing the following lines:

echo 'export PATH=/path/to/miR-island:\$PATH' >> ~/.bashrc

source ~/.bashrc

Note: "/path/to/miR-island" should be changed to the real location of miR-island folder.

You can test by typing miR-island.pl on the command line with no parameters. You should get a help message.

INPUT DATA

- The reference genome in FASTA format is required. Any assemble level (contig level, scaffold level and chromosome level) is ok. If no genome is available, users can use EST/cDNA sequences instead.
- 2. RNA-seq data sets, which should be in FASTA format.
- 3. Known miRNAs in FASTA format is encouraged, which is not essential.
- 4. Tab-delimited text file that contains the location of known microRNA precursors (**only** separate the location and the name with a TAB), which is not essential:

Chr2 - 10676451 10676573 ath-MIR156a

Chr4 + 9888982 9889070 ath-MIR160b

ATTENTION: only characters of "A/T/C/G" are allowed in the above input FASTA files.

PREPROCESSING DATA SETS

If the raw reads is SRA format, you should install SRA Toolkit first. The current available version is at http://trace.ncbi.nlm.nih.gov/Traces/sra/sra.cgi?view=software. Then run fastq-dump SRRxxx.sra, while xxx stands for the SRR number. You should get a file in FASTQ

format named as SRRxxx.fastq. Detection and Removal of adapters need two scripts including "find_3p_adapter.pl" and "trim_illumina_sRNA_fastq.pl" at http://axtell-lab-psu.weebly.com/tools.html (Axtell, 2013).

Detect adapters by running find_3p_adapter.pl -m ugacagaagagagagagacac < SRRxxx.fastq, while "ugacagaagagagagacac" stands for the sequence of ath-miR156a. Users are welcome to read the script's help document first.

Remove adapters by running trim_illumina_sRNA_fastq.pl -a TCGTATGC -e 1 -q N -o SRRxxx.fa -p SRRxx_ps.fa < SRRxxx.fastq, while "TCGTATGC" is the adapter sequence. The RNA-seq data sets must be parsed into FASTA format.

PLANT MICRORNAS ANNOTATION AND QUANTIFICATION

Given that you have a tomato sRNA data set, whose cotyledon type is dicot, and there is a lot of known miRNAs referred to this species available in miRBase, miR-Island can run with the following command:

miR_island.pl -c dicot -g genome.fa -r sRNA_clean.fa -k miRNA.fa -p 4 -x coord.list

It will generate a directory names "*MiR_Island_run*" and a sub-directory names *run_xxxxxxx*, while xxxxxx is a series number returned by the function of localtime in perl.

You will get two miRNA expression lists: "denovo_miRNA_expression.list"; "known miRNA expression.list".

In the above two expression lists, NoN_mature_exp stands for the mature sequence raw reads count, NoN_star_exp stands for the star sequence raw reads count. NoN_mature_std stands for the mature sequence RPM, NoN_star_std stands for the star sequence RPM. The character N in red color represents integer, and it is in accordance with the order of the input sRNA data sets. When there are two small RNA data sets, "Ctl" or "Trt" will take the place of "NoN", "Ctl" stands for the first small RNA data set, and "Trt" stands for the second small RNA data set. The Log2 transformed Fold Change and P-value will be calculated (Audic and Claverie, 1997; Benjamini and Yekutieli, 2001).

However, if the known miRNAs referred to this species were not specified, miR-island can run with the following command:

miR_island.pl -c dicot -g genome.fa -r sRNA_clean.fa -p 4

This time no known miRNA expression list will exist.

In addition, an intact flat file named "result.out" will be generated, one sample is as follows:

```
>SL2.40ch02 2904
microRNA name
        sly-MIR171a
        ST.2.40ch02 - 41330642 41330741
precursor coord
        TGATTGAGCCGTGCCAATATC
mature sequence
        TATTGGCCTGGTTCACTCAGA
score total
          25.2
score for read counts
              18
            3.3
score for mfe
total read count
             47
42
mature read count
loop read count
star read count
                  obs
pri_seq
                  pri struct
                  #MM
t00475147_x5
                  ....uauuggccugguucacucaga.....
t01197946 x1
                  ugauugagccgugccaau......
t00343046 x1
                  t01687524 x40
                  С
           aca
                  au
  gaUAUUGGC UGGUUCA UCAGAc aaaaug aaacu auu g
  |||||| ||||| ||| a
  CUAUAACCG GCCGAGU AGUuug
             uuuugc uuuga
            C11-
```

If the coordinate of a predicted microRNA precursor is in accordance with that in coord.list, microRNA name will be specified. The precursor coordinate contains four parts: chromosome, strand, begin and end. The tail is a figure of the secondary structure (Shen et al., 2012), and the mature or star region are in Upper characters.

ABOUT MIR-ISLAND PACKAGE

MiR-island package has a total of six perl scripts:

- 1. miR_island.pl
- 2. transform_genome.pl
- $3.\ diff_miRNA_expression_analysis.pl$
- 4. excise potential precursors.pl
- 5. RNAfold_with_multi-threads.pl
- 6. miRNA_island_core.pl

You can get each help message by simply typing the script name.

miR_island.pl

```
MiRNA_Island: an ultrafast package for annotation and quantification of miRNA and MIRNA
genes with High throughput sequencing
WARNING: You did not provide enough information!
[-1 <haipin_length>] [-p <num_threads>] [-x <pre-miRNA_coord>]
REQUIRED:
-c <monocot|dicot>
                                cotyledon type: monocot | dicot
-g <genome.fa> reference genome file in multi-fasta format
-r <reads1.fa[,...,readsN.fa]> a comma-separated list of files with small RNA reads in
                                multi-fasta format
OPTIONAL:
                 maximum threshold MFE to be annotated as a potential precursor (def: -17.5)
-e <min_mfe>
-f < int>
                 minimum frequency of reads to trigger a precursor excising (def: 15)
-i <bowtie ebwt>
                 prefix of the reference genome's bowtie indexes file (def: same as option "g") known miRNAs of the related species in multi-fasta format
-k <miRNA.fa>
                 maximum length of the potential precursor to be excised (def: 277)
-1 < int>
-m < int>
                 maximum number of sites that a read could map to related genome (def: 10)
-p <int>
                 number of threads to use (def: all threads available)
-x <coord>
                 known pre-miRNA coordinates in tab-delimited file (only seperate
                  the location and the pre-miRNA name with a TAB (such as):
                  Chr2 - 10676451 10676573 ath-MIR156a
                 Chr4 + 9888982 9889070
                                           ath-MIR160b
-h
                 print intact help message
```

```
transform genome.pl
```

```
transform_genome.pl transfoms scaffold-level genome into chromosome-level and vice versa;
it can also be used to transform the genome to multi-fasta with sequences in same length
WARNING: You did not provide enough information!
Usage: transform_genome.pl -d <genome.fa> -i <index_file> -t <T> -o <pseudo_genome.fa>
        transform_genome.pl -d <data> -i <index_file> -t <F> -o <data_processed>
        transform_genome.pl -d <genome.fa> -t <O> -o <genome_with_same_length.fa>
OPTIONS:
-d
         data to transform or restroe
         index file that record scaffold info
-i
         specify the output file
-0
         function type to use: T/F/O
-t
         T to transform scaffold-level genome to chromosome-level
         F to restore chromosome-level info to scaffold-level
         O to transform genome with same sequence length
-h
         print intact help information
diff_miRNA_expression_analysis.pl
diff_miRNA_expression_analysis.pl aims at analyzing differential microRNA expression data. It will output a list with standard gene count, LFC (log2 transformed foldhange),
pvalue, FDR and significance, etc.
Usage:
     diff_miRNA_expression_analysis.pl -m <miRNA.fa> -c <combined_reads.fa> \
                                           -r file_1.fa[,...,file_N.fa] -o <file_out>
     Options:
 -c <reads.fa>
                                     a combined multi-fasta file of clean reads generated
                                     by format_clean_reads.pl
 -l <exp.list>
                                     a tab-delimited file contains microRNAs sequence info
 -m <miRNA.fa>
                                     a multi-fasta format file contains known microRNAs
 -r <file_1.fa[,...,file_N.fa>
                                     a comma-separated list of files with small RNA reads
                                     in multi-fasta format
                                     a tab-delimited output file
 -o <exp.out>
Note:
       option 'c' and option 'l' are mutually exclusive;
       The input reads should be formated by format_clean_reads.pl. The reads
       file should have each entry with unique sequence, the entry must be as '>CR_xN', while C represents one or more letters, R represents a non-redundant
       running number, and N represents current read count, _x is the separator.
Pvalues, FDR and significance will be calculated only in the case of two samples.
excise potential precursors.pl
excise_potential_precursors.pl excises potential miRNA precursors with high
throughput sequencing.
Version = 1.0
WARNING: You did not provide enough information!
Usage: excise_potential_precursors.pl -b <reads_vs_genome_sorted.bam> -r <genome.fa> \
                                          -o cursors.fa> [-m <min_freq>]
REQUIRED:
                     profile of reads mapped to the related genome in a formatted, sorted and indexed \ensuremath{\mathsf{BAM}} file
-b <sorted.bam>
-r <genome.fa> a related genome in multi-fasta format
-o recursors.fa> an output file in multi-fasta format
OPTIONAL:
                     maxmium length of the potential precursor to be excised (default: 277)
-1 < int >
-m <int>
                     minimum frequency of reads to trigger a precursor excising
```

RNAfold_with_multi-threads.pl

```
RNAfold_with_multi-threads.pl computes potential precursors using RNAfold with
multiple threads
Version = 1.0
WARNING: You did not provide enough information!
Usage: RNAfold_with_multi-threads.pl -i cursors.fa> -d <tmp_dir>
                                            -o cursor.struct> [-p <num_threads>]
REQUIRED:
-i cursors.fa>
-d <tmp_directory>
                               a multi-fasta file with each sequence in a line
                               a directory to store the temp files when running
                               an RNAfold output file with sequences and structures
-o cursors.struct>
OPTIONAL:
                               number of threads to use
-p <num_threads>
miR_island_core.pl
miR_island_core.pl is the miR-island package's core algorithm for plant microRNA annotation
Version = 1.0
WARNING: You did not provide enough information!
Usage: miR_island_core.pl -b <reads_vs_genome_sorted.bam> -c <monocot|dicot> -q <qenome.fa>
           _s <structure_file> -t <table_out> -r <result_out> [-m <miRNA.fa>] [-1 <col_width>]
REQUIRED:
-b <sorted.bam>
                       profile of reads mapped to the related genome in a formatted, sorted
                       and indexed BAM file
                       \verb|cotyledon| type: \verb|monocot|| dicot|
-c <monocot|dicot>
                       reference genome file in multi-fasta format
-g <genome.fa>
-s <structure_file>
                       secondary structure file produced by RNAfold
-r <result_out>
                       an intact output file that annotates miRNA and MIRNA genes
-t <table_out>
                       a brief tab-delimited text file that annotates miRNA and MIRNA genes
OPTIONAL.
                       coloum 1 width, if identifier is long, add this value (def: 40) known miRNAs of the related species in multi-fasta format
-1 <column width>
-m <miRNA.fa>
                       show the current version number
-h
                       print intact help message
```

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