### Introduction

There are some useful scripts.

## vcf2snpbinner.R

A R script for transforming vcf file to SNPBinner input file. The heterozygosity and missing rate will be calculated for output SNP marker.

#### Usage

```
Rscript ./vcf2snpbinner.R -h
```

```
usage: vcf2snpbinner.R [--] [--help] [--opts OPTS] [--input INPUT]
       [--out OUT] [--parent1 PARENT1] [--parent2 PARENT2] [--minDP_p1
      MINDP_P1] [--minDP_p2 MINDP_P2] [--max_missing MAX_MISSING]
a program for converting vcf to table of snpbinner. genotype same as
parent_1 is designated 'a', genotype same as parent_2 is designated
'b', heterozygous genotype is designated 'h', missing genotype is
designated '-'
flags:
  -h, --help
                    show this help message and exit
optional arguments:
  -x, --opts
                    RDS file containing argument values
  -i, --input
                    vcf or vcf.gz file containing two parents and
                    progeny lines
                    output file prefix
 -o, --out
  -p, --parent1
                    name of parent_1
 -P, --parent2
                    name of parent 2
 -d, --minDP_p1
                    Minimum depth of parent 1 [default: 5]
 -D, --minDP p2
                    Minimum depth of parent 2 [default: 5]
  -m, --max_missing Maximum missing rate of SNP [default: 0.3]
```

## Haplotype.R (GeneStructure with Variation)

A R script for drawing gene structure and the variation of this gene in a population. A gtf file containing target gene, a vcf file containing variation of this gene and phenotype data is needed. Hierarchical clustering algorithm was adopted to distinguish different haplotype, the number of haplotype can be designated

according clusting result. Some polymorphism may exsit within samples belonging to the same haplotype, you can divided them into different haplotypes by setting more haplotypes.



A figure containing gene structure and variation information

# ePCR.pl

A perl script for ePCR. Input is tsv (tab-separated values) file containing three columns (PrimerID, forwardPrimer, Reverse Primer).

### Requiement

ePCR

#### Preparation

```
# lower letter to UPPER letter
seqkit seq -u reference.fa > genome.fa
# prepare sequence database for re-PCR searches
famap -t N -b genome.famap genome.fa
fahash -b genome.hash -w 12 -f 3 genome.famap
```

#### Usage

```
perl ePCR.pl -h
```

```
# Usage: ePCR.pl --input primer.txt --output output.txt
# Required:
  --input <string>
                     input filename, one pair primer per line, tab
seperated, e.g.:
                     primerID
#
                             Left_primer_seq Right_primer_seq
  --output <string>
                     output filename.
```

# slidingWindow.R

A sliding window function in R. The R package tidyverse should be installed. values is a vector containing column names which need be calculted.

```
source("./slidingWindow.R")
# An example
sldWid <- slidingWindow(df = df, winSize = 1000000, winStep = 200000, groups =
"CHROM", position = "POS", values = c("R.R3.depth", "R.qY.depth"), fun = "mean")</pre>
```

## addUp.R

A R function for calculating accumulation value for a column of a table. For example, a data.frame contain two columns, "chromosome" and "position", this function will calculate the accumulation position of different chromosome, then a list containing a table with a new column "position\_add\_Up", a vector containing breaks position, a vector containing labels, a vector containing gaps position, will be returned.

```
source("./addUp.R")
# An example
addUp(df = df, len = len, group = "chromosome", pos = "position", band = 0.01)
addUp(df = df, len = len, group = "chromosome", pos = c("start", "end"), band =
0.01)
```

## run\_DESeq2.R

A R script for differential expression analysis using DESeq2 (with biological replication). You need to prepare three files:

### Requiement

- read count matrix file.
- samples file, tab-delimited text file indicating biological replicate relationships. e.g.

```
cond_A cond_A_rep1
cond_A cond_A_rep2
cond_B cond_B_rep1
cond_B cond_B_rep2
```

contrasts file, tab-delimited text file containing the pairs of sample comparisons to perform. e.g.

```
cond_A cond_B
cond_Y cond_Z
```

#### Usage

```
Rscript run_DESeq2.R -h
```

```
usage: run_DESeq2.R [--] [--help] [--opts OPTS] [--matrix MATRIX]
       [--samples_file SAMPLES_FILE] [--min_reps MIN_REPS] [--min_cpm
      MIN_CPM] [--contrasts CONTRASTS]
Run differential expression analysis using DESeq2.
flags:
  -h, --help
                    show this help message and exit
optional arguments:
 -x, --opts
                   RDS file containing argument values
                     matrix of raw read counts (not normalized!)
  -m, --matrix
 -s, --samples_file tab-delimited text file indicating biological
                     replicate relationships.
                     At least min count of replicates must have cpm
  --min reps
                     values > min cpm value. [default: 2]
                     At least min count of replicates must have cpm
  --min_cpm
                     values > min cpm value. [default: 1]
                    file (tab-delimited) containing the pairs of
  -c, --contrasts
                     sample comparisons to perform.
```

# extractSeq.sh

If there is a gene/transcript id you are interested and the corresponding genomics data, then you want to abtain genomic, CDS, pep sequence and gene structure information of this gene/transcript, you can use this shll script.

### Usage

```
#chmod u+x extractSeq.sh
./extractSeq.sh --help
```

```
Usage: /home/wangpf/bin/extractSeq.sh [--genome genome_file] [--gff3 gff3_file] [--cds cds_file] [--pep pep_file] [--id gene/mRNA_id] [--up up] [--down down] [--gz] Options:

--genome Specify the genome fasta file
--gff3 Specify the gff3 file
--cds Specify the cds fasta file
--pep Specify the pep fasta file
--id Specify the gene/mRNA id
```

```
    --up Specify how many bp upstream for gene/mRNA
    --down Specify how many bp downstream for gene/mRNA
    --gz Compress all the result if this option is present
    --help Display this help message
```

## run\_edgeR.R

A R script for differential expression analysis using edgeR (without biological replication). You need to prepare three files:

#### Requiement

- read count matrix file.
- samples file, tab-delimited text file indicating biological replicate relationships. e.g.

```
cond_A sample_A
cond_B sample_B
```

• contrasts file, tab-delimited text file containing the pairs of sample comparisons to perform. e.g.

```
cond_A cond_B
cond_Y cond_Z
```

#### Usage

```
Rscript run_edgeR.R -h
```

```
usage: run_edgeR.R [--] [--help] [--opts OPTS] [--matrix MATRIX]
       [--samples_file SAMPLES_FILE] [--min_reps MIN_REPS] [--min_cpm
      MIN CPM] [--contrasts CONTRASTS] [--dispersion DISPERSION]
Run differential expression analysis using DESeq2.
flags:
 -h, --help
               show this help message and exit
optional arguments:
 -x, --opts
-m, --matrix
 -x, --opts
                     RDS file containing argument values
                     matrix of raw read counts (not normalized!)
 -s, --samples_file tab-delimited text file indicating biological
                      replicate relationships.
                      At least min count of replicates must have cpm
  --min_reps
                      values > min cpm value. [default: 1]
```

```
--min_cpm At least min count of replicates must have cpm
values > min cpm value. [default: 1]
-c, --contrasts file (tab-delimited) containing the pairs of
sample comparisons to perform.
-d, --dispersion edgeR dispersion value. [default: 0.1]
```

# get\_longest\_seq.pl

In some genomics data, there are multiple isoforms fo one gene because of alternative splicing. This perl script can get the longest CDS or pep sequence of genes.

#### Requiement

- perl module
  - o Bio::SeqIO
- data file
  - CDS or pep sequence file
  - gff file

### Usage

```
perl get_longest_seq.pl -h
```

### Output

- <outprefix>.longest.fa, the longest CDS or pep sequence in fasta format with gene id as sequence identifier
- <outprefix>.longest.list, a list contain gene id, longest mRNA id and the length of longest mRNA

# find\_gaps.py

There are gaps in majority genome fasta files except T2T genome. This find\_gaps.py script will find gaps position.

#### Requiement

- python3 module
  - o argparse
  - Biopython
- input data
  - o genome fasta file

#### Usage

```
chmod u+x find_gaps.py
find_gaps.py -h
```

### Output

A tab-separated text file containing three columns.

```
<chromosome_id> <gap_start> <gap_end>
```

A fasta file with split contigs.

### mummerCoordsPlot.R

This script can draw a alignment plot according to mummer's show-coords program. In fact, this script is modeled after mummerCoordsDotPlotly.R in dotPlotly. mummerCoordsDotPlotly.R is a excellent program, but there are a few aspects that I am not entirely satisfied with:

- It will sort query ID by length, but I want keep query ID order in the query genome;
- Yellow is not very noticeable on a white background, so I use red and blue;
- Add the ability to color the alignment results based on whether they are forward or reverse complementary;
- IDs was not on the middle of chromosomes or contigs in figure;
- I typically work with the tidyverse ecosystem, so I have rewritten this script using tidyverse syntax for clarity and consistency;
- Some other modifications.

#### Requiement

- R package
  - argparser
  - o tidyverse
- input data
  - o Alignment result from mummer's show-coords program;

```
show-coords example.filter.delta > example.coords
```

Tab-separated text files containing reference and query IDs separately, first column is IDs in genome, the second is what you want to show in figure, only chromosomes or contigs in these files will be shown (optional). If not assigned, all IDs of chromomsomes or contigs longer than --min-query-length will be shown in figure, or you can just assign in the command line, e.g. "GWHERGL00000001:A01,GWHERGL00000002:A02,GWHERGL00000003:A03"

```
GWHERGL00000001 A01
GWHERGL00000002 A02
GWHERGL00000003 A03
GWHERGL00000004 A04
GWHERGL00000005 A05
```

 Tab-separated text files containing chromosome or contig length, first column is IDs in genome, the second is the corresponding length (optional);

```
GWHERGL00000001 59205763
GWHERGL00000002 39005548
GWHERGL00000003 44769757
GWHERGL00000004 28346059
GWHERGL00000005 46671336
...
```

#### Usage

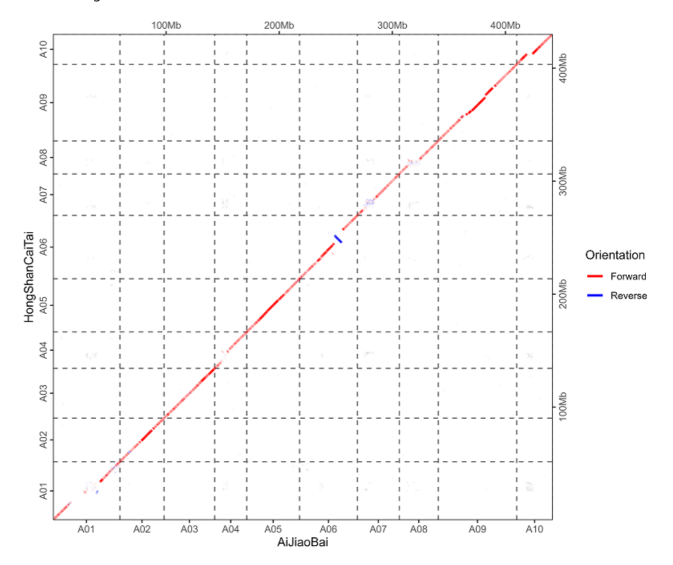
Rscript mummerCoordsPlot.R -h

```
usage: mummerCoordsPlot.R [--] [--help] [--opts OPTS] [--input INPUT]
       [--out OUT] [--ref REF] [--query QUERY] [--refIDs REFIDS]
       [--queryIDS QUERYIDS] [--reflen REFLEN] [--queryLen QUERYLEN]
       [--min-query-length MIN-QUERY-LENGTH] [--min-alignment-length
       MIN-ALIGNMENT-LENGTH] [--min-identity MIN-IDENTITY] [--color-by
       COLOR-BY] [--size SIZE] [--width WIDTH] [--height HEIGHT]
Generates plots of alignment data produced by show-coords.
flags:
  -h, --help
                              show this help message and exit
optional arguments:
  -x, --opts
                              RDS file containing argument values
 -i, --input
                              coords file from mummer program
                              'show-coords'
 -o, --out
                              outfile prefix
  -r, --ref
                              reference name shown in plot [default:
                              Reference]
  -q, --query
                              query name shown in plot [default: Query]
  -R, --refIDs
                              a tab-separated file containing two
                              columns, first column is IDs in reference
                              genome, second column is names to shown
                              in plot, or a string like
                              'scaffoldA01:A01,scaffoldA03:A03'. If
                              there is no file named that, we will
                              threat it as a string
  -Q, --queryIDs
                              a tab-separated file containing two
                              columns, first column is IDs in query
                              genome, second column is names to shown
                              in plot, or a string like
                              'scaffoldA01:A01,scaffoldA03:A03'. If
                              there is no file named that, we will
                              threat it as a string
  -1, --refLen
                              a tab-separated file containing two
                              columns, first column is IDs in reference
                              genome, second column is chr length
  -L, --queryLen
                              a tab-separated file containing two
                              columns, first column is IDs in query
                              genome, second column is chr length
  -M, --min-query-length
                              filter queries with total alignments less
                              than cutoff X bp [default: 4e+05]
  -m, --min-alignment-length
                              filter alignments less than cutoff X bp
                              [default: 2000]
  -s, --min-identity
                              filter alignments with identity less than
```

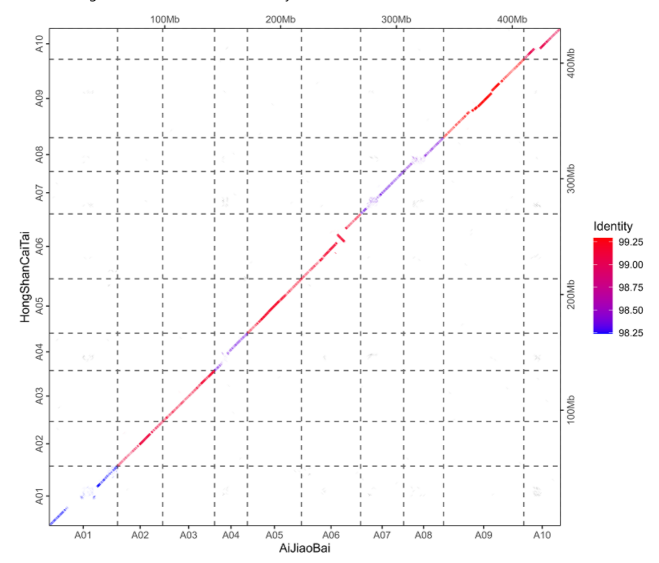
	X % [default: 90]	
-c,color-by	turn on color alignments by 'direction'	
	or 'identity', no color if not assign	
-S,size	line width of alignments in figure	
-W,width	<pre>plot width (inches) [default: 10]</pre>	
-H,height	<pre>plot height (inches) [default: 10]</pre>	

## Output

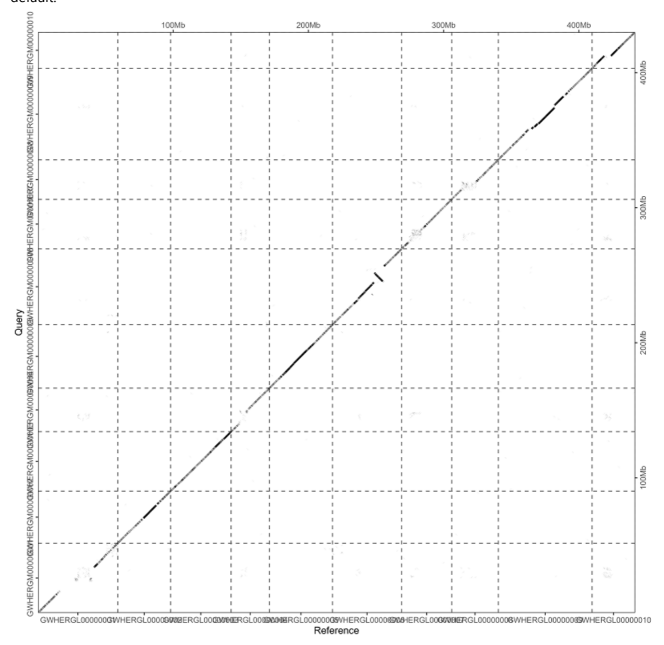
• Color the alignment results based on direction.



• Color the alignment results based on identity.



• default.



#### **Data Source**

These two genome is from this paper:

Zhou, Yifan et al. "The complexity of structural variations in Brassica rapa revealed by assembly of two complete T2T genomes." Science bulletin vol. 69,15 (2024): 2346-2351. doi:10.1016/j.scib.2024.03.030