Protocol

Recommended programing languages and tools

- Perl 5.18.2
- Python 2.7.13
- EMBOSS:6.6.0.0 (use segret program in the package)
- Genbank_splice.py (error occurs when we run with python3)
 https://github.com/jrjhealey/bioinfo-tools/blob/master/Genbank_slicer.py
- sed
- Easyfig 2.2.2 (http://mjsull.github.io/Easyfig/)
- (optional) inkscape 0.92.2

Preparation of input files (Genbank format) for Easyfig

First, we extract annotation information for each single chromosome from the annotation file (annotation.gff3).

- \$ perl -ne 'print if/^1/' data/annotation.gff3 > 1.gff3
- \$ perl -ne 'print if/^2/' data/annotation.gff3 > 2.gff3
- \$ perl -ne 'print if/^3/' data/annotation.gff3 > 3.gff3
- \$ perl -ne 'print if/^4/' data/annotation.gff3 > 4.gff3
- \$ perl -ne 'print if/^5/' data/annotation.gff3 > 5.gff3

Next, we merge DNA sequence information (fasta format) and annotation information (GFF3 format) of each single chromosome (annotation.gff3) into a single genbank file using the 'seqret' program in EMBOSS package.

- \$ seqret -sequence data/1.fa -feature -fformat gff -fopenfile 1.gff3 -osformat genbank -outseq 1.gbk -osdirectory ./
- \$ seqret -sequence data/2.fa -feature -fformat gff -fopenfile 2.gff3 -osformat genbank -outseq 2.gbk -osdirectory ./
- \$ seqret -sequence data/3.fa -feature -fformat gff -fopenfile 3.gff3 -osformat genbank -outseq 3.gbk -osdirectory ./
- \$ seqret -sequence data/4.fa -feature -fformat gff -fopenfile 4.gff3 -osformat genbank -outseq 4.gbk -osdirectory ./
- \$ seqret -sequence data/5.fa -feature -fformat gff -fopenfile 5.gff3 -osformat genbank -outseq 5.gbk -osdirectory ./

Moreover, we extract DNA sequence and annotation information within a specific chromosomal region using the Genbank splice.py python script.

```
$ python2 scripts/Genbank_slice.py -g 1.gbk -o 1_pnr700.gbk -s 1842912 -e 2542912 -v
```

- \$ python2 scripts/Genbank_slice.py -g 2.gbk -o 2_pnr700.gbk -s 1817796 -e 2517796 -v
- \$ python2 scripts/Genbank_slice.py -g 3.gbk -o 3_pnr700.gbk -s 585879 -e 1285879 -v
- \$ python2 scripts/Genbank_slice.py -g 4.gbk -o 4_pnr700.gbk -s 1576680 -e 2276680 -v
- \$ python2 scripts/Genbank_slice.py -g 5.gbk -o 5_pnr700.gbk -s 1255217 -e 1955217 -v

Finally, we replace the "misc_RNA" feature in our annotation file for the "gene" feature so that the annotation information will be recognized in the later drawing process by Easyfig software.

```
$ sed -i " 's/misc RNA/gene/' 1 pnr700.gbk
```

- \$ sed -i " 's/misc RNA/gene/' 2 pnr700.gbk
- \$ sed -i " 's/misc_RNA/gene/' 3_pnr700.gbk
- \$ sed -i " 's/misc_RNA/gene/' 4_pnr700.gbk
- \$ sed -i " 's/misc_RNA/gene/' 5_pnr700.gbk

Drawing homologous regions between chromosomes using Easyfig

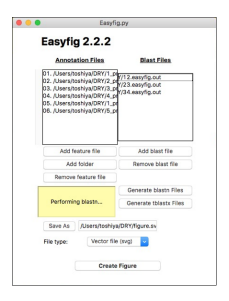
First, we launch Easyfig.

\$ /Applications/Easyfig_2.2.2_OSX/Easyfig

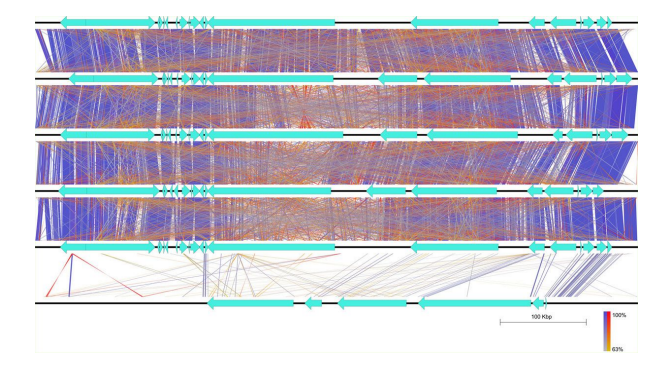
Then, we register input files used in drawing a figure by clicking the "Add feature file" button.



We can identify homologous regions between the chromosomal regions of interest using built-in blastn program by clicking the "Generate blastn files" button.



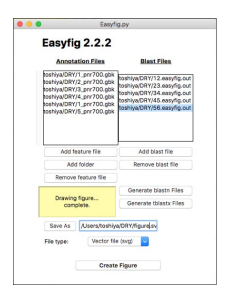
Raw blastn result files include repeat sequences that will generate noisy background in the final figure as in the figure below.



Therefore, we need to filter blast hits at repetitive sequences using the filter_repeat.pl Perl script.

- \$ perl scripts/filter_repeat.pl -i ./
- \$ perl scripts/filter_repeat.pl -s -i ./56.easyfig.out -m 250

After filtering blast hits at repeat sequences, we remove the unfiltered Blast result files by clicking the "Remove blast file" button. Then, we register the filtered Blast result files by clicking the "Add blast file" button.



Finally, we change drawing settings for the present dataset as below,

Menu bar: Image>Figure

Width of Figure: 5000 > 7000

Length of scale legend (in base pairs): 0 > 100000

Menu ba: Image>Blast

Choose minimum blast colour: inverted, Gray > Yellow

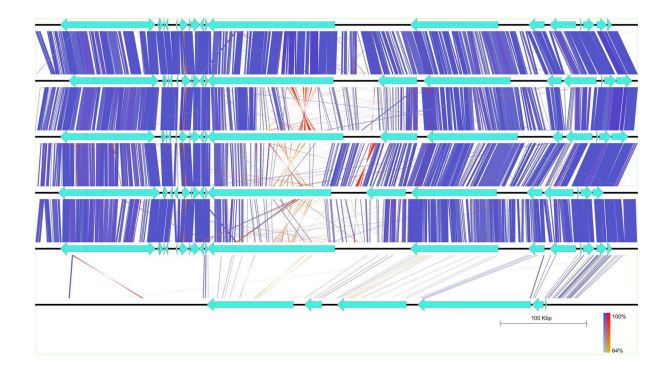
Choose minimum blast colour: normal, Dark_gray > Blue, inverted, Dark_grey > Red

Outline blast hits in black: checked > unchecked

Save As: File name (or File path)

File type: svg

and export the result figure by clicking the "Create Figure" button.



The result image file (svg format) can be imported to the free illustration software inkscape. We can manually modify colors of the different classes of genes, and insert annotations as below.

