Reassortment Detection Manual

The purpose of this note is to document, in a first draft, how to re-run the code for the Reassortment Detection analysis.

Input files:

A CSV file called "[handle] Sequences.csv", which contains the following information downloaded from the IRD:

- Segment
- Protein Name
- Sequence Accession
- Complete Genome
- Sequence Length
- Collection Date
- Host Species
- Country
- State/Province
- Flu Season
- Strain Name
- A FASTA file containing all of the sequences downloaded, with only the Accession Number in the header.

Place both of these files under a single directory.

Make the following sub-directories:

- alignments
- distmats
- edges
- reassortant_edges
- shell scripts
- split_fasta
- sge_outputs

There is also a list of scripts present:

- align_sh.py
- align.py
- clean_affmats_sh.py
- clean_affmats.py
- compile_affmats_sh.py
- compile_affmats.py

- full_affmat.py
- graph_combiner.py
- graph_initializer.py
- max_edge_finder_sh.py
- max_edge_finder.py
- node_data_imputer.py
- preprocessing.py
- second_search.py
- sequence_splitter_sh.py
- sequence_splitter.py
- source_pair_combiner.py
- source_pair_manual_sh.py
- source_pair_sh.py
- tables_functions.py

Step 1: Pre-processing CSV file

- On the Rous server (or any system that has the SunGrid Engine installed, write a .sh file with the following bash command:
- python preprocessing.py [handle], where [handle] refers to the standard name that is used across all of your scripts.
- Wait a while for the preprocessing script to complete.
- If everything completes successfully, you will see a CSV file by the name [handle] Full Isolates.csv.
- Clean up the directory by running the following command in the terminal:

```
- mv *.sh.* sge_outputs/.
- cd ..
```

Step 2: Splitting FASTA file by segment

- To enable parallel running of the splitting script, which might take a while, I have provided a Python script (sequence_splitter_sh.py) which writes a series of shell scripts to submit.
- Modify this file, by replacing the handle string at the top with your handle. An example modification can be: '"20141103 All IRD"'
- Run the following bash commands:

```
python sequence_splitter_sh.pycd shell_scriptsqsub sequence_splitter.sh
```

• Wait until all of the jobs have been run.

- You should see all 8 segments being processed and written to disk. They will be in the directory /split_fasta/[Handle] Segment [Number].fasta
- While in the shell_scripts directory, clean up the directory by running the following commands:
- mv *.sh.* ../sge_outputs/.
- cd ..

Step 3: Alignment and Distance Matrix Generation

- To enable parallel running of the alignment script, I have provided align_sh.py.
- Modify this file by replacing the handle string at the top with your handle.
- Run the following bash commands:
 - python align_sh.py
 - cd shell_scripts
 - qsub align.sh
- Wait until all of the jobs have been run. On a large dataset, it'll take a while on the order of hours, depending on CPU usage and data size.
- While in the shell_scripts directory, clean up the directory by running the following commands:

```
- mv *.sh.* ../sge_outputs/.
- cd ..
```

Step 4: Convert Distance Matrix into Thresholded Similarity Matrix, and Rename the Index/Columns.

- To enable parallel running of the compilation scripts, I have provided the following scripts:
 - clean_affmats_sh.py <-- corresponding with clean_affmats.py</pre>
 - compile_affmats_sh.py <-- corresponding with compile_affmats.py</pre>
- Run the following bash commands:
 - python clean_affmats_sh.py
 - cd shell_scripts
 - qsub clean_affmats.sh
- Wait until all of the jobs have been run.
- When all of the jobs are done, clean up the shell_scripts directory as per before.
- Then, run the following bash commands:
 - python compile_affmats_sh.py

- cd shell_scripts
- qsub compile_affmats.sh
- Wait until the job is done.
- When the job is done, clean up the shell_scripts directory again.

Step 5: Make a summed affinity/similarity matrix

- Create a shell script in the shell_scripts directory, that includes the following commands:
 - cd ..
 - python full_affmat.py [handle]
- Run the following bash commands:
 - cd shell_scripts
 - qsub full_affmat.sh
- Wait until the job is done.
- When the job is done, similarly, clean up the shell_scripts directory.

Step 6: Search for max edges for each virus - full 'transmissions' only.

- Run the graph initializer script: python graph_initializer.py [handle]. This one should run fast. The expected output is a .pkl file which houses the initialized network with only nodes present.
- Run the script: python max_edge_finder_sh.py
- Then, run the following commands:
 - cd shell_scripts
 - qsub max edge finder.sh
- Wait until all the jobs are done.
- When done, clean up the shell_scripts directory.

Step 7: Combine found edges into a condensed graph.

- Run the graph combiner script: python graph_combiner.py [handle]
- If your "handle" is different, i.e. you have a different prefix for all of the files, then replace the text inside the quotation marks with your handle.
- This one should be fast, i.e. within a dozen minutes.

Step 8: Compile a list of nodes to perform source pair searches on.

- Run the script using: python second_search.py [handle] [percentile]
 - [handle]: the common prefix to all of your files.
 - [percentile]: the cutoff percentile of whole genome edges to try source pair searches. Our analysis used the 10th percentile, so the value input was "10".
- This script should run fast.
- The expected output is a .pkllist file with a list of nodes to perform source pairs on.

Step 9: Perform source pair searches.

- Run the following bash commands:
 - python source_pair_sh.py
 - cd shell_scripts
 - qsub source_pair.sh
- Wait for the job to finish completing. This one should take a while, depending on CPU availability. May take up to a few hours.
- When done, clean up the shell_scripts directory.

Step 10: Combine source pairs with full graph.

- Run the following bash commands: python source_pair_combiner.py [handle]
- This one should run fast (~minutes).

Step 11: Annotate graph with edge and node metadata

- Run the following bash commands:
 - python graph_pwi_finder.py [handle] (this should take a few minutes) - this will annotate edge PWIs
 - python graph_cleaner.py [handle] (also should be fast)
- At this point, the graph construction steps are complete.