Armillaria Gallica Gene Analysis Tools

Methods

This section will brief explanation of all methods used over the course of this class. For all programs discussed here see https://github.com/ofbujak/Armillaria_gallica_gene_analysis_tools.

Read Depth Snapshots

The first work completed was the analysis of the read depth values for each of the 15 sequenced strains of the Armillaria gallica. This work was based on analysis which a former student, Hao Wang, had done previously to determine locations for each strain that had unusually high read depth. These locations are based on reads aligned to a reference. Unfortunally, we do not know the exact method which Hao had used to determine which locations he had deemed had significant read depth. Though we do not know the exact method he used, the regions he identified were those that had high read depth. He outputted his results into text files consisting of three columns, a scaffold number column, a location start column, and a location end column. These files had variable numbers of locations of high significance but they averaged approximatly 2636 locations per strain. With the goal to produce graphs of these locations of high read depth, in order to gain a understainding of the "landscape" about those locations, I first made up of the samtools depth command in a program called make_read_depth_files.sh

make_read_depth_files.sh will output a file which has three columns, scaffold, location,

and read depth. Using these results I wrote a program, read_depth_per_location.c, which will take in as arguments a specific samtools read depth file, a scaffold to search for, a start and end location (all space separated). read_depth_per_location.c will then print information (scaffold, location, read depth, etc...) on the locations, incrementing by 1 from the start location, for all locations which are within the range passed into it. This program was used in plot_all_read_depths.sh to produce graphs of each range of locations with high read depth \pm 500. The program used to create the graphs was plot_read_depth.R. Some examples of these graphs can be found in figures ?? and ??.

Read Depths For Whole Scaffolds

In order to produce depictions of the read depths for an entire scaffold the framework established to create the read depth snapshots was used with some slight changes in methodology. To produce these graphs the files produces from using the samtools depth command were searched via grep in order to separate each scaffold result into its own file. These files were then passed into plot_scaffold_read_depth.R and plot_scaffold_read_depth_histogram.R. plot_scaffold_read_depth.R is a program which graphs the read depth on the y axis and the location of that read depth on the x axis. plot_scaffold_read_depth_histogram.R creates a histogram of the read depths with binwidths of 5.

Read Depth Analyses

In order to gain insight into the different aspects of the scaffolds and the reads which were aligned I produced four metrics for each strain. I calculated the global average read depths, determined the total number of sites with reads aligned to them, calculated and graphed the average number of reads per scaffold, and graphed and determined the number of locations with reads aligned for each scaffold.

Unaligned Reads

To output all reads which are not aligned into a sam file the following samtools view command can be used: samtools view -f 4 file.bam > unmapped.sam. Following this The reads can be assembled via a de novo assembler. (The assembly of these has not been carried out yet).

New Assemblies

To produce new assemblies I attempted to do both *de novo* assemblies of five of the strains (Ar170, Ar174, Ar179, Ar196, Ar213). Following verification of quality using FastQC, both methods were used.

De Novo Assemblies

To produce new *de novo* assemblies of five of the strains (Ar170, Ar174, Ar179, Ar196, Ar213) were attempted. The program used for the assmeblies was Velvet, with the wrapper VelvetOptimiser used. The command used to run VelvetOptimiser was:

'perl VelvetOptimiser.pl -f "-shortPaired -fastq -separate

<Path To Forward Strand File>/<Forward_strand.fastq.gz>

<Path To Reverse Strand File>/<Reverse_strand.fastq.gz>" -t 32 -s 31 -e 31

-d <Output Directory>

Reference Based Assemblies

'To produce another reference based assembly the scipt fastqThroughNovoalign.sh was

used. Due to time constraints, the reference based assemblies have not been analyzed as of

yet.

Taking the Consensus of Aligned Read

Following identification of locations where read depth is significant it was useful to be

able to quickly get the actual DNA sequence which is aligned to a specific location. To do

this, custom programs were created to sort through the output from samtools view and to

take the consensus of the reads aligned. The total pipeline for these programs is encapsulated

within consensus_reads.sh.

To output sequences that were easier to work with, samtools view was used. The com-

mand used was:

samtools view -o <output_filename> <bam_file>

<scaffold_number>:<location_start>-<location_end>

Following outputting of the sam file, the program 'pull_reads.c' is used. This program

takes in the five columns listed above (in the order discussed) and outputs all the reads,

which fall within the range specified, into a file where each read occupies its own line with

each read offset by the difference between the start of the range of interest and the location

which the read starts.

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The output created by pull_reads.c is then passed into average_reads.c. average_reads.c takes in the output created by pull_reads.c and the difference between the start and end locations. This program creates 4 arrays of the size of the difference passed into the program and iterates over all lines in the file. When a base is encounteres at a location the array for that base is incremented by 1, at the location of that base. After all the lines in the file are iterated over, the mode of each base for each element of the array are then output, resulting in a consensus of the reads that were aligned between two locations.

Calculation of Average Read Depth and Number of Aligned Reads per Scaffold

The program avg_read_depth.c takes in a read depth file and the scaffold which the average of which will be calculated and outputs the average read depths for the locations with reads aligned to them and the total number of locations with reads aligned to them for that scaffold.

Determiation of Regions Of High Read Depth

To create a new method for deremining the regions of the assembly which had significant read depth high_read_depth_regions.c was created. This program takes in the average number of reads aligned to a scaffold, and the number of locations on a scaffold with reads aligned, and searches for all locations with read depth higher than a certian threshold. The threshold used for these analyses was a multiple of five fold the standard deviation plus the mean. Any regions which were found to be above that were identified as having significant read depth.

In addition to the search this program also allows for a 10 base grace peroid. If a region was found to have significant read depth had some locations immediatly following it which were below the threshold for less than 10 consecutive locations before the read depth rose above the level of significance than the entire region, including the up to 10 bases below that region would be reported as once contigious region of high read depth.

Compairisons of Read Depths

The program compare_read_depths.c takes in two read depth files, a reference file and a compairison file. This program iterates over all the lines in the file and comapires each location and calculates the difference between read depths.

Analysis of DNA Sequences Suspected to Result in High Read Depths

Using the locations found to have significant read depth and the method used to take the consensus of the the aligned reads between locations, the sequences at each location which had high read depth could be extracted. After these sequences were extracted they could be searched, via ncbi blastn, to attempt to determine if the sequence has any homologous in other species/ to determine the effect of the sequence.

Insertions and Deletions

Kassandras stuff here.

main

All the stuff Thomas did:

- read depth snapshots check!
- average read depth stuff
- total number of aligned reads
- program to take mode of reads
- scaffold graphing program
- read depth differences program
- assemblies
- blasting stuff
- RNA verification

Read Depth Snapshots

In order to gain a understanding of how the locaitons which Hao had determined had notably high read depth the regions Hao identified were graphed. There were approximatly between 2000 and 4000 regions for each of the 15 strains. Although there were many graphs four predomanant types of high read depth regions stood out: Normal, Rectangular, Right skewed, and Left skewed.

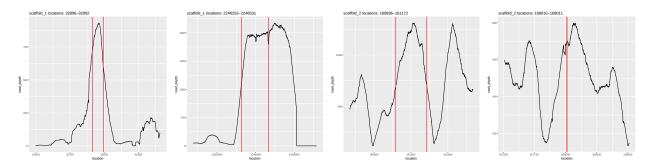


Figure 1. Examples of the four types of high read depth regions (left most) Normal, (left middle) Rectangular, (right middle) Left skewed, (right most)Right skewed

Average Read Depths and Number of Reads Aligned per Scaffold

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Read Depth Analyses

Unaligned Reads

Indel Analysis

Future Directions