Using NextGENe[®] Software to assemble Illumina[®] MiSeq[™] Paired End Reads with the Stepwise PE Assembler

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Introduction

Sequences that repeat throughout the genome can pose a problem for the assembly of short reads. Normally the repeat reads would assemble within the incorrect contig (figure 1). This causes two problems- the repeat regions elsewhere in the genome have reduced coverage and contigs terminate at repeat regions due to the formation of multiple ambiguous contigs.

NextGENe software solves this problem with a stepwise paired-end assembly. The end of a contig produced by assembly may indicate a repeat region. The software first calls in the reads paired to those assembled (using overlaps) at the end of the contig (up to 1 ½ times the library size from the end). These reads are then assembled and the software chooses the most complete assembly to continue the contig (figure 2). Shorter assemblies are not used because they are formed by repetitive sequences erroneously assembled together due to their similarity.



Figure 1 – Incorrect assembly of reads in a repeat region. Purple reads are from the same repeat sequence elsewhere in the genome.



Figure 2 – NextGENe calls in the paired reads and assembles them. Reads paired to those that do not belong in the original assembly will form a separate contig (purple). Only the longest contig is used in the assembly.





Procedure

In this analysis, a 150bp PE read dataset from a MiSeq instrument was used to assemble the E coli MG1655 genome. First, the format conversion tool is used to convert the raw data (FASTQ files) to FASTA format while performing some filtering and trimming based on the quality scores (Figure 3). Next, the data is loaded using NextGENe's Project wizard. The instrument type (Illumina) and application (de novo assembly) are selected (Sequence Condensation is not used). The PE assembler is selected, and settings are adjusted (Figure 4).

Settings				
Median Score Threshold >=	20	Max # of Uncalled Bases <= 3		
Called Base Number of Each Read >= 25				
✓ Trim or Reject Read when >=	3	Base(s) with Score <= 16		
✓ Paired Reads Data	Save As	FASTA File(*.fasta) 💌		

Figure 3 – Format Conversion Settings

The "word length" setting should be adjusted based on expected coverage. A word length of 23 is optimal for 20 to 30x coverage and a word length of 29 is sufficient for 50x coverage. The maximum suggested word length is 31. Longer word lengths allow for greater reduction of noise. The "high coverage limited" setting can be used to ignore coverage over a certain depth in order to improve processing speed. "Final contig merging" occurs after the PE assembly- the resulting contigs are checked for overlapping regions where they can be merged. "Reduce Memory Usage" will index only the 5' end of reads, which can result in a significant reduction in the amount of memory used for longer reads.

PE Assembly Options		
✓ Paired Read Data		
Library Size 250		
Long Library Size(>1000 Bases) Section Size 400		
Word Length 31 Minimum Scaffold Length 150		
₩ High Coverage Limited: Max Coverage = 100		
Final Contig Merging Reduce Memory Usage		

Figure 4 – Assembly Settings

Results

The original dataset consisted of 8,331,866 paired-end 150bp reads. After format conversion using default settings (with the paired reads option selected) there were 7,267,665 reads remaining. These reads were 137 bp on average, for an expected depth of coverage of 229x.

The assembly results are summarized in table 1.

Processing Time	1 hour, 48 min
Total Reads	7,267,665
Reads Used	7,101,508
% Reads Used	97.7%
Number of Contigs	65
Average Length	70,747 bp
Minimum Length	363
Maximum Length	494,530 bp
N50	212,730 bp

Table 1 – Assembly Results

The assembled contigs were aligned to the E. coli MG1655 reference genome. The coverage curve report was used to identify regions of the reference genome that were not covered at least once (Figure 5). Only 144,789 bp (3.12%) of the reference was not covered.





Figure 5 – Coverage Curve Report

Discussion

NextGENe's PE Assembler is a highly accurate and efficient tool for assembly of paired-end or mate-pair data. A small bacterial genome such as E. coli can be assembled into less than 70 contigs from a single MiSeq data run in less than 2 hours. The assembler can be run on relatively low-cost hardware- in this case the program needed less than 8 GB of RAM.

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