16S RNA Sequencing Data Management using SQLite

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Abstract

16S Ribosomal RNA Sequencing is used extensively in analyzing bacterial phylogeny and taxonomy. This project attempts to streamline the 16S sequencing pipeline using local file databases to replace multiple flat sequence files used in the pipeline, to ease logistical burdens on the researcher and enable greater metadata analysis and accountability of experiments.

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1 Introduction

The 16S small subunit of bacterial ribosomes are highly gene sequences found in all bacteria, which means that the differences within the 16S RNA profiles can be used as an analogue for species identity. Since the 16S gene contains both highly conserved and highly variable regions all interspersed together, the highly conserved regions can be amplified using PCR (Polymerase chain reaction), while the variable regions are used to classify the organism. HTS (High Throughput Sequencing) using Illumina sequencers are then used to sequence the PCR products from the prior step.

The output from HTS forms the start of the computational pipeline. In the preprocessing stage, poor quality reads are filtered and trimmed. Chimeras¹ and non-bacterial 16S reads are then removed. The pipeline then attempts to produce phylogenetic classification through the use of classification based on sequence similarity (i.e. OTU Analysis). By comparing the filtered output from HTS against the already existing taxonomies of over 2

¹a single organism composed of genetically distinct cells

million species, the genus, family, and order of the sample can be determined.

The majority of the pipeline is executed on the University of Oregons ACISS High-Performance Supercomputer Cluster, and utilizes PBS scripts (normal shell scripts with extra variables defined to manage job resources) to execute the different stages of the pipeline.

The FASTQ output produced by the Illumina sequencer is run through the preprocessing stage of filtering, trimming, and demultiplexing. The PBS script for the preprocessing stage calls on the Demultiplexer, a python script written by Rodger Voelker, which removes the primer attached to the sequences during the amplification process, and attaches barcodes (signifying sample origin) to both ends of the paired-end reads in the FASTQ file. After demultiplexing, the pipeline proceeds to use Trimmomatic v0.32, an open source tool to trim poor quality feeds from the reads. It uses a sliding window trimming, that cuts out sequences when the average quality within a window falls below a certain threshold. Finally, Bowtie 2.1.0 is used to align the reads to existing mitochondrial and phiX sequences and to remove them.

2 Data Management

Management of data in the pipeline is conceptually divided into four tasks: creating semantic structure within the database; storing sequence data in compressed form; transforming stored data into formats required by tools in the pipeline; logging settings and parameters of transformations and related

metadata.

2.1 Semantic Database Structure

Instead of juggling multiple input files such as raw Illumina FASTQ sequences, barcodes, and offsets, Pip creates several tables which impart a well-defined semantic structure to the data and contains them within a single file.

A Pip database is initialized with five tables:

primer holds all primer sequences used in the experiment

offset randomized offsets to aid 16S sequencing

barcode experiment identifier

log stores all metadata related to data, experiment, or pipeline

read1 stores the first part of paired-end sequence data

read2 stores the second part of paired-end sequence data

Thus, a Pip database provides a logical container for the data and stores them in a compressed, lossless form.

Every operation done in Pip can be stored as metadata in the log table, providing a detailed record of all settings and parameters used at the time of operation. Results of OTU Analysis can be affected by parameters used by the tools upstream in a pipeline, thus having a reliable record of operations and transformations to the data will aid in analysis downstream.

2.2 Data Compression

FASTQ files generated by Illumina NGS machines have this general format:

@HWI-ST0747:277:D1M96ACXX:6:1101:1232:2090 1:N:0:

GGATAGTACTAGGGTATCTAATCCTGTTTGCTCCCCACGCT...

+

ACCCFFDDFH#FHHIGIJJFJJJJJJJJJJJIIJJJGIJJG...

in ASCII text form, with each character in the file taking up 8 bits. The format uses five characters to represent the four DNA-bases (A - adenine, T - thymine, C - cytosine, G - guanine) and N to represent an errorneous read. Each base is paired with a quality score that has 42 values (0 to 41). Compression works by representing each base with an integer range (A: 0 to 49, T: 50-99, C: 100-149, G: 150-199, N: 200-255) and adding the quality score to the beginning of that range. Thus, base 'A' with a quality score of '#' (value 0) can be represented by the value 0, while base 'G' with a quality score of 'J' (value 41) can be represented by 191 (150 + 41). The resulting values fit within an 8-bit character, which is a 50% reduction in space usage. Since both compression and decompression require only addition operators, the scheme is fast and lossless.

2.3 Data Transformation

A central part of managing the sequence data in the 16S pipeline is the manipulation and transferring of sequences between the various tools. There

Tool	Expected input format
Trimmomatic	FASTQ/Gzipped FASTQ
Bowtie2	FASTQ/FASTA
QIIME	454-FASTA/454-Quality Scores

Table 1: Expected input formats for various tools

is often a need to modify the structure or format of the data to fit the varying input requirements of those tools. Table 1 shows the various input formats that tools in the pipeline require.

Traditionally, the way to manage the various formats was to transform and store various copies of the data in multiple files. For example, the first stage of the pipeline usually involves using Trimmomatic to filter poor quality reads and to trim all the reads to a certain length. Doing this requires the user to provide 2 paired-end FASTQ sequence files to Trimmomatic, which will then produce 5 output files: unpaired and paired sequences 1 and 2, and a trimming log. The output files have sizes similar to the input, which results in the space usage nearly doubling after the first stage in the pipeline. Subsequent stages will then take the output of the first stage and create more output. Thus, both the number of files and their file size will increase very quickly due to the number of stages within the 16S pipeline.

Using a plug-in architecture, Pip can "stream" sequence data to tools in the pipeline without writing to files on disk. Pip uses UNIX Named Pipes to feed data to tools such as Trimmomatic and Bowtie while simultaneously consuming the output they produce back into the database. UNIX Named Pipes allows programs to transfer data through memory instead of disk, and therefore do not require reading and writing to disk. However, named pipes appear as size zero, regular files on a filesystem, thus tools can read and write from those files without needing any changes to the software. In the case of Trimmomatic, Pip provides a named pipe to capture the contents of the trimming log, and uses the details within it (sequence number, position of trim, number of characters trimmed) and stores them in a "trimmed" table. Since this table only stores the sequences which were operated on, instead of entire copies of the original sequences, the size increase of the Pip database after the first stage is almost negligible. Additionally, instead of needing to manage multiple output files and their naming, the data is now ready for the next stage of the pipeline.

3 Results

Pip was benchmarked on a 2.7 GHz Intel Core i7 processor with 16GB of RAM, running OS X 10.9. Table 2 shows the time taken for the specified number of sequence inserts into a new Pip database. Figure 1 plots the number of actual raw inserts per second into SQLite, showing consistent insertion rates between 96,000 and 102,000 raw inserts per second.

Number of sequences (paired-end)	Time taken (seconds)
500,000	10.35
1,000,000	19.57
2,000,000	39.22
4,000,000	78.67
8,000,000	160.84
16,000,000	319.38
32,000,000	644.90
64,000,000	1,286.26

Table 2: Insertion speeds into SQLite using Pip

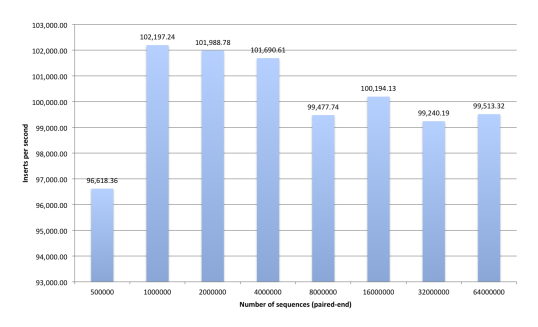


Figure 1: Inserts per second into SQLite using Pip

Table 3 shows the size of the raw FASTQ sequence file compared to the resulting database file after inserts. Figure 2 reveals a consistent 45% reduction in file size after insertions.

Number of sequences (paired-end)	Input FASTQ size (MB)	Pip database (MB)
500,000	371.40	206.90
1,000,000	743.00	413.90
2,000,000	1,486.00	828.00
4,000,000	3,051.52	1,699.84
8,000,000	6,082.56	3,389.44
16,000,000	12,165.12	6,789.12
32,000,000	24,350.72	13,568.00
64,000,000	48,701.44	27,146.20
91,000,000	69,754.88	35,061.76

Table 3: Comparison of input file sizes against Pip database sizes

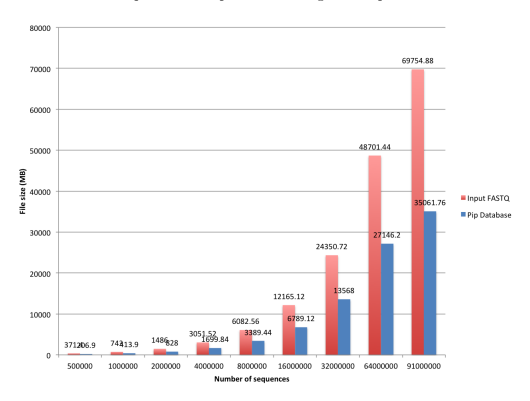


Figure 2: Chart of file sizes

4 Conclusion

The use of Pip within the 16S pipeline shows promise in reducing the logistical effort of researcher and interoperability headaches of multiple tools. Currently, Pip operates on the first two tools of the 16S pipeline (Trimmomatic, Bowtie) and shows promise. Pip can be extended to with tools later in the pipeline easily through a plug-in architecture. There appears to be no performance penalty in streaming data compared to disk access, and the underlying SQLite implementation is able to deal with datasets of up to 91 million rows without slowdowns.