# CGAL - 0.9.8 BETA Manual

### 1 Introduction

CGAL is a tool for comparing genome assemblies. It computes the likelihood of an assembly given a set of paired end reads which can be used as a metric for evaluating the assembly.

## 2 Installation

CGAL can be run on any environment that supports POSIX threads. To install it execute the following commands:

make

# 3 Running

To run CGAL the reads need to be mapped to the assembly using an external mapper. At present we support Bowtie 2 and BFAST. The output of the mapper need to be in SAM format. Once mapping is done, running CGAL consists of three steps:

• The first step converts the output generated by Bowtie 2 or BFAST to an internal format.

If you are using Bowtie 2, please use '-a -no-mixed' options to map and then run

./bowtie2convert out.sam maxFragmentLength

If BFAST is used run

./bfastconvert out.sam maxFragmentLength

where out.sam is the output file of the mapper in SAM format and maxFragmentLength is the maximum length of fragment i.e. the maximum insert size. Mappings with longer fragment length than maxFragmentLength will be ignored. If this value is set too high intermediate files will be quite large and runtime will be greatly increased. The default value is 5000.

• The tools for mapping are usually not able to align all the reads. To align the reads not mapped by the mapper, we have adapted the striped implementation of Smith-Waterman algorithm by Farrar. However, this step is time consuming. So, we align only a random subset of reads.

#### ./align contigFile toAlign numThreads

where contigFile is the assembly file in FASTA format, toAlign is the number of reads to be aligned and numThreads is the number of POSIX threads. In our experiments we have used values between 300 and 1000 for toAlign.

• cgal This step computes the likelihood value.

### ./cgal contigFile

where contigFile is the assembly file in FASTA format. The ouput is written to file out.txt.

### 3.1 Notes

- Each step generates intermediate files used by later steps. These files must not be deleted until final step is completed.
- CGAL does not filter out reads other than the ones with more than 80% of the read covered with N's.

# 4 Output

The ouput file contains following values separated by tabs:

- Number of contigs
- Total likelihood value
- Likelihood value of reads mapped by the mapping tool
- Likelihood value corresponding to reads not mapped
- Total number of paired-end reads
- Number of reads not mapped by the mapper