Genome Assembly Tutorial

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Genome Assembly Workshop



Genome assembly with PacBio data using Canu

Why are we using this program?

- *Relatively* user-friendly and easy to install
- Not computationally intensive with small data sets
- Well maintained and documented

Genome assemblers

Other useful assemblers:

Illumina data:

w2rap-contigger

https://github.com/bioinfologics/w2rap-contigger

- can take lots of computer power to run
- works with single paired-end library

soapdenovo2

https://sourceforge.net/projects/soapdenovo2

- relatively easy to install and run
- works with large genomes

Genome assemblers

PacBio data:

falcon & falcon-unzip

https://github.com/PacificBiosciences/FALCON

http://profs.scienze.univr.it/delledonne/Papers/2016%20Chin%20NMethods.pdf Very powerful assemblers but not easy to install or use.

Quiver/Arrow/pbalign - genome alignment and polishing. Part of PacBio Genomic Consensus package.

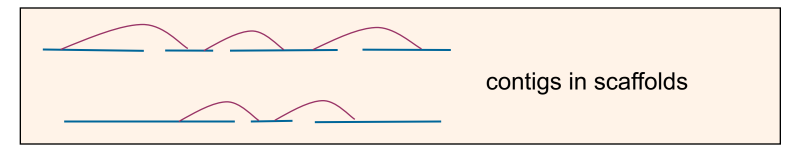
https://github.com/PacificBiosciences/GenomicConsensus

Again, powerful but not the easiest to use.

Canu

http://genome.cshlp.org/content/27/5/722

Linear assemblies

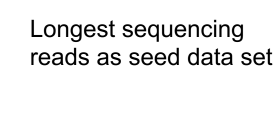


contig: a contiguous sequence of bases....

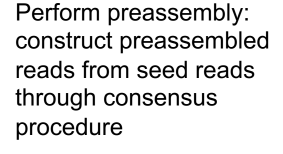
scaffold: a sequence of contigs, separated by gaps....

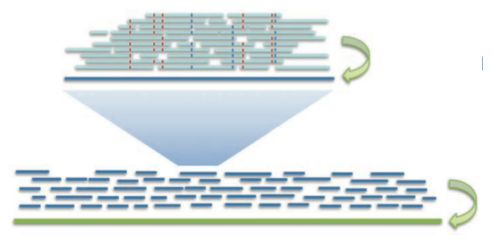
Number of Ns = predicted gap size, with error bars (can't be displayed in fasta format)

Long read sequencing



Use seed dataset to map shorter reads





Preassembled reads used for genome assembly

Error correction: mapping high-quality short PacBio reads to long reads

Canu: "unitigs"

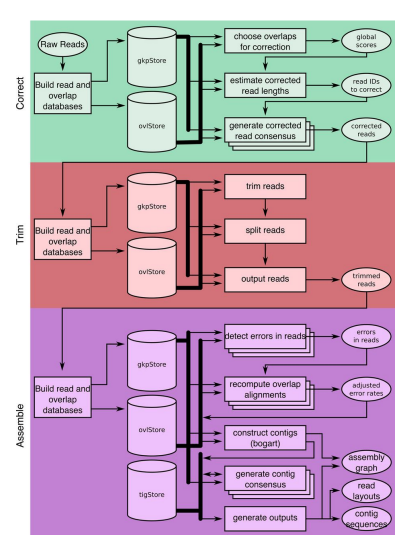
PB-only Correction & PolishingChin et al (2013) Nature Methods. 10:563–569

Using Canu

- Canu is an assembler that specializes in PacBio or Oxford Nanopore long-read data
- Derived from Celera assembler
- Corrects reads, trims suspect regions (e.g., adaptors)
 then assembles the corrected reads
- Can be run with one command to do all steps or each step can be run separately (correcting, trimming and assembling).
- Koren S, Walenz BP, Berlin K, Miller JR, Phillippy AM. Canu: scalable and accurate long-read assembly via adaptive k-mer weighting and repeat separation. Genome Research

A full Canu run includes three stages: correction (green), trimming (red), and assembly (purple).

Generates k-mer histogram and conducts all-vs-all overlaps



Correct:

Select best overlaps to generate corrected reads. Uses longest 40X reads for correction

Trim:

Identifies unsupported regions and trims reads to longest supported range

Assemble:

Identify sequencing errors, constructs best overlap graph and outputs contigs.

Sergey Koren et al. Genome Res. 2017;27:722-736

Using Canu

Outputs contigs and unitigs.

 The unitig construction task finds sets of overlaps that are consistent, and uses those to place reads into a multialignment layout. The layout is then used to generate a consensus sequence for the unitig or a "high-confidence contig"

 We will use the sample Pac Bio E. coli data found on the canu homepage: http://canu.readthedocs.io
 25x E.coli fastq files with quality scores.

To install Canu:

Login as admin

Make a directory for installation

```
$ git clone <a href="https://github.com/marbl/canu.git">https://github.com/marbl/canu.git</a>
```

\$cd canu/src

\$ make -j 4 Compiler error?

Compiler error?

- \$brew install gcc
- \$which gcc

Try making again:

make -j 4

Where is it installed?

Executables found in PATHWAY/canu/Darwin-amd64/bin

Need gnuplot for plotting:

- \$ which gnuplot
- \$ brew install gnuplot@4

```
If you need to have gnuplot@4 first in your
PATH run:
   echo 'export
PATH="/usr/local/opt/gnuplot@4/bin:$PATH"'
>> ~/.bash profile
```

Getting Help

/canu/documentation/source - help docs

- \$ canu command line options
- \$ canu -options parameter options

Download canu test E.coli dataset:

http://canu.readthedocs.io/en/latest/quick-start.html?highlight=25x

- 25X subset (223Mb)

Where do you want them to download?

```
$ curl -L -o pacbio.fastq
http://gembox.cbcb.umd.edu/mhap/raw/ecoli_p6_
25x.filtered.fastq
```

- -o naming downloaded file pacbio.fastq
- -L if page has moved to a new location, curl will redo at the request at the new site

How many reads within the file? How many lines?

Running Canu

- screen unix command to run programs in multiple windows or "screens"
 - If your computer breaks connection, your program will continue run on server
- \$ screen to open a new screen
- \$ ctrl-a d to suspend screen
 - -Program will continue to run
- \$ screen -r to re-open existing screen session
- \$ screen -r #### use number to re-open if multiple sessions running

Running Canu

 We will run default with correcting, trimming and assembling all in one command:

```
$ /your pathway/Deb/canu/Darwin-amd64/bin/canu
  -p ecoli -d ecoli-pacbio genomeSize=4.8m -pacbio-raw \
your_path/canu/pacbio.fastq saveReads=true \
gnuplot=/usr/local/opt/gnuplot\@4/bin/gnuplot
2>CanuRun_20181022.log **to save output

*where are reads...include pathway
-p assembly-prefix
-d output directory
input types:
  -pacbio-raw -pacbio-corrected -nanopore-raw -nanopore-corrected fastq or fasta format

*gnuplot option specific to our exercise
```

**USE THE HELP DOCS!!!!

Running Canu - Manual assembly

Correct, Trim and Assemble Manually

 You can also do the three top-level tasks by hand. This would allow trying multiple construction parameters on the same set of corrected and trimmed reads.

Previous command to correct, trim & assemble:

```
canu -p ecoli -d ecoli-full genomeSize=4.8m -pacbio-raw \
your_path/canu/pacbio.fastq saveReads=true \
2>full_out_102517.log **to save output
```

Using Canu – Manual assembly

First correct the raw reads:

```
canu -correct -p ecoli -d ecoli-manual genomeSize=4.8m \
-pacbio-raw your_path/canu/pacbio.fastq \
saveReads=true 2>correct_out_102517.log
```

Then trim the output of the correction:

```
    canu -trim -p ecoli -d ecoli-manual genomeSize=4.8m \
        -pacbio-corrected ecoli-manual/ecoli.correctedReads.fasta.gz \
        saveReads=true 2>trim out 102517.log
```

*corrected reads created during first step

Using Canu - manual assembly

Finally, assemble the output of the trimming twice using two error rates (can use as many as you like):

```
canu -assemble -p ecoli -d ecoli-erate-0.013 genomeSize=4.8m \
correctedErrorRate=0.013 -pacbio-corrected ecoli-
manual/ecoli.trimmedReads.fasta.gz saveReads=true \
2>assemble-0.013_out_102517.log
```

```
canu -assemble -p ecoli -d ecoli-erate-0.025 genomeSize=4.8m \
correctedErrorRate=0.025 -pacbio-corrected ecoli-
manual/ecoli.trimmedReads.fasta.gz saveReads=true \
gnuplotImageformat=svg 2>assemble-0.025_out_102517.log
```

*trimmed reads created during second step

^{*}The error rate specifies the difference in overlap between two corrected reads which is typically <1% (canu default value 0.045) for PacBio data and <2% (canu default 0.144) for Nanopore data (<1% on newest chemistries). Higher rate, more sensitive.

^{*}Notice in the output there are separate directories for each error rate you specify.

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  -p ecoli -d ecoli-pacbio genomeSize=4.8m -pacbio-raw \
your_path/canu/pacbio.fastq saveReads=true \
gnuplot=/usr/local/opt/gnuplot\@4/bin/gnuplot
2>CanuRun_20181022.log **to save output

*where are reads...include pathway
-p assembly-prefix
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input types:
  -pacbio-raw -pacbio-corrected -nanopore-raw -nanopore-corrected fastq or fasta format

*gnuplot option specific to our exercise
```

**USE THE HELP DOCS!!!!

Canu output files - all found in helpdocs

ecoli*.report - assembly analysis log

READS

- ecoli*.correctedReads.fasta.gz The sequences after correction, trimmed and split based on consensus evidence. Typically >99% for PacBio and >98% for Nanopore but it can vary based on your input sequencing quality
- ecoli*.trimmedReads.fasta.gz The sequences after correction and final trimming. The corrected sequences above are overlapped again to identify any missed hairpin adapters or bad sequence that could not be detected in the raw sequences.

SEQUENCE

- ecoli*.contigs.fasta Everything which could be assembled and is part of the primary assembly, including both unique and repetitive elements.
- ecoli*.unitigs.fasta Contigs, split at alternate paths in the graph
- ecoli*.unassembled.fasta Reads and low-coverage contigs which could not be incorporated into the primary assembly.

Canu output files - all found in helpdocs

SEQUENCE

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- **ecoli*.unassembled.fasta** Reads and low-coverage contigs which could not be incorporated into the primary assembly.

The header line for each sequence provides some metadata on the sequence: >tig00000001 len=576432 reads=573 covStat=249.02 gappedBases=no class=contig suggestRepeat=no suggestCircular=no

tig # len=<integer> reads=<integer> covStat=<float> gappedBases=<yes|no> class=<contig|bubble|unassm> suggestRepeat=<yes|no> suggestCircular=<yes|no>

*covStat: Is the log of the ratio of the contig being unique vs it being two-copy, based on the number of reads Positive means more likely to be unique, negative means more likely to be repetitive

Canu output files - all found in helpdocs

GRAPHS

ecoli*.gkpStore - *svg

LAYOUT

The layout provides information on where each read ended up in the final assembly, including contig and positions. It also includes the consensus sequence for each contig.

- ecoli*.unitigs.layout.readToTig The position of each read in a contig
- ecoli*.contigs.layout.tigInfo, ecoli*.unitigs.layout.tigInfo A list of the contigs (unitigs), lengths, coverage, number of reads and other metadata.

How much coverage is enough?

4.5 Mb - 223Mb fastq reads (25X), 13,124 reads

Try some assemblies with filtered data sets.

Using canu - split reads

Count and split files:

\$wc -l pacbio.fastq

```
$ split -1 output number of lines - full: 52,496
- 60% 31,498
$ split -1 31498 pacbio.fasta
$ mv xaa ecoli_filtered_0.60.fastq
$ rm the rest (xa*)
```

- confirm number of lines and reads

How much coverage is enough?

4.5 Mb - 223Mb fastq reads (25X), 13,124 reads

Try some assemblies with filtered data sets:

0.75 - 9,843 reads, 168 Mb, ~18X

0.60 - 7,874 reads, 133Mb, ~15 X

0.50 - 6,562 reads, 111Mb, ~12.5X

0.25 - 3,281 reads, 56Mb, ~ 6.3X

How much coverage is enough?

Full data set:

4.8 Mb - 223Mb fastq reads (25X), 13,124 reads

Percent Reads	Coverage	Contigs	Unitigs	Unassembled Reads	Genome Size (Mb)
I	25X	I	I	1,506	4.6
0.75	18X	I	I	1,109	4.6
0.6	I5X	8	8	959	4.6
0.5	12.5X	33	35	996	4.5
0.25	6.3X	67	75	1,188	1.9

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- **ecoli*.unassembled.fasta** Reads and low-coverage contigs which could not be incorporated into the primary assembly.

The header line for each sequence provides some metadata on the sequence:

tig # len=<integer> reads=<integer> covStat=<float> gappedBases=<yes|no> class=<contig|bubble|unassm> suggestRepeat=<yes|no> suggestCircular=<yes|no>

ecoli-0.25.contigs.fasta:

>tig0000001 len=102939 reads=69 covStat=1.36 gappedBases=no class=contig suggestRepeat=no suggestCircular=no

ecoli-full.contigs.fasta:

>tig00000001 len=4639282 reads=10263 covStat=3780.63 gappedBases=no class=contig suggestRepeat=no suggestCircular=no

Poisson Distribution

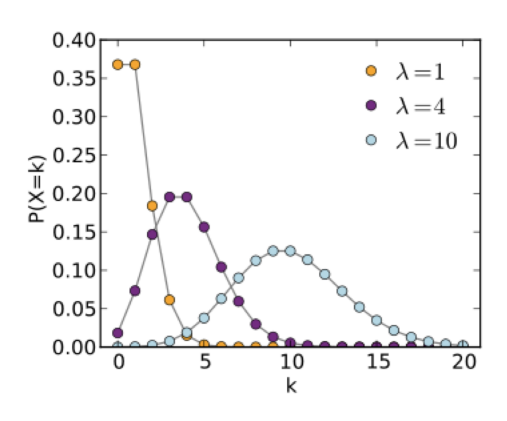
The probability of a given number of events occurring in a fixed interval of time and/or space if these events occur with a known average rate and independently of the time since the last event.

Resembles a normal distribution, but over the positive values, and with only a single parameter.

Key property:

The standard deviation is the square root of the mean.

$$P(k) = \frac{\lambda^k}{k!} e^{-\lambda}$$



Genome Assembly Problem Set

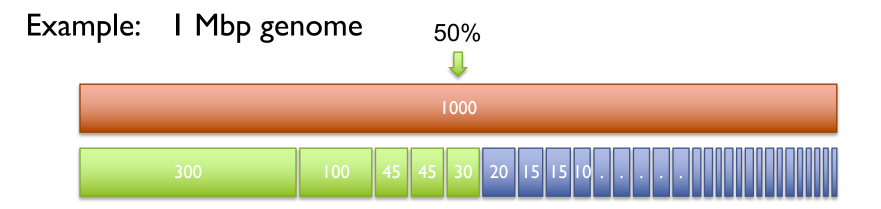
Using ecoli-0.25.contigs.fasta, write a script that reports:

- I. The number of contigs in the file
- 2. The shortest contig.
- 3. The longest contig.
- 4. The L50 size
- 5. The N50 size

The fasta.file is listed on github workshops/GenomeAssembly: ecoliPB-filtered_0.50.contigs.fasta.

N50 size

50% of the genome in contigs as long as or larger than N50 value



N50 size = 30 kbp
$$(300k+100k+45k+45k+30k = 520k >= 500kbp)$$

A greater N50 is usually a sign of assembly improvement

- Comparable with genomes of similar size
- Genome composition can bias comparisons
- High L50 vs Low N50

L50 size

Number of contigs that are as long or longer than the N50 value



L50 size = 30 kbp
$$(300k+100k+45k+45k+30k = 520k >= 500kbp)$$

- High L50 vs Low N50
 - longer sequences and fewer of them....in theory
 - lower stringency can inflate N50