Assembly Tutorial

Michael Schatz

Oct 25, 2012 CSHL Sequencing Course





Outline

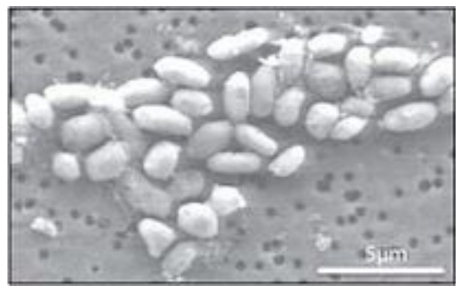
I. Sample Data for your mission!

2. ALLPATHS-LG

3. MUMmer

Halomonas sp. GFAJ-I





Library 1: Fragment

Avg Read length: 100bp

Insert length: 180bp

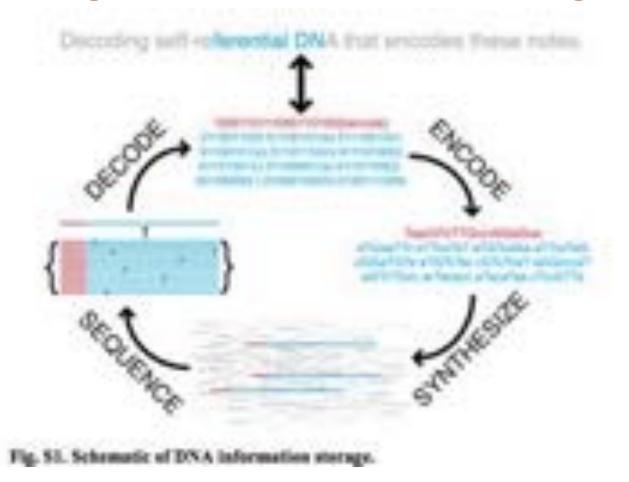
Library 2: Short jump

Avg Read length: 50bp

Insert length: 2000bp

A Bacterium That Can Grow by Using Arsenic Instead of Phosphorus Wolfe-Simon et al (2010) *Science*. 332(6034)1163-1166.

Digital Information Storage



Encoding/decoding algorithm implemented in dna-encode.pl from David Dooling.

Next-generation Digital Information Storage in DNA Church et al (2010) Science. 337(6102)1628

Mission Impossible

1. Setup virtual machine

sudo apt-get install gnuplot-x11 cd ~/allpathslg-43436 sudo make install

2. Download Reference Genome

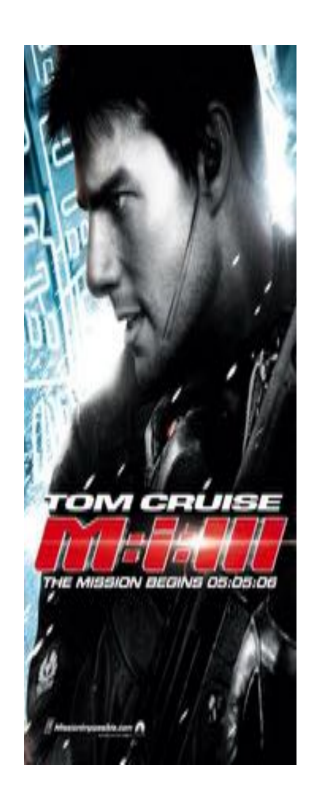
wget http://schatzlab.cshl.edu/data/AHBC01.fasta.gz gunzip AHBC01.fasta.gz

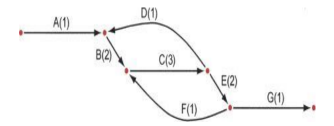
3. Download reads

wget http://schatzlab.cshl.edu/data/asm.tgz tar xzvf asm.tgz

4. Decode the secret message

- 1. Assemble the reads, Align to reference, Extract secret
- 2. dna-encode.pl -d





Running ALLPATHS-LG lain MacCallum



How to use ALLPATHS-LG

- 1. Data requirements (*** most critical thing ***)
- 2. Computational requirements & Installation
- 3. Preparing your data
- 4. Assembling
- 5. What is an ALLPATHS-LG assembly?

ALLPATHS-LG sequencing model

Libraries (insert types)	Fragment size (bp)	Read length (bases)	Sequence coverage (x)	Required
Fragment	180*	≥ 100	45	yes
Short jump	3,000	≥ 100 preferable	45	yes
Long jump	6,000	≥ 100 preferable	5	no**
Fosmid jump	40,000	≥ 26	1	no**

**For best results. Normally not used for small genomes.

However essential to assemble long repeats or duplications.

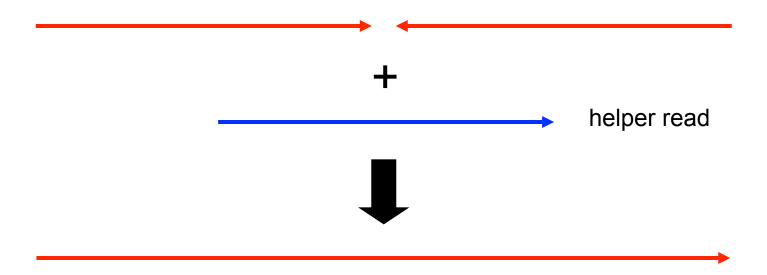
Cutting coverage in half still works, with some reduction in quality of results.

All: protocols are either available, or in progress.

^{*}See next slide.

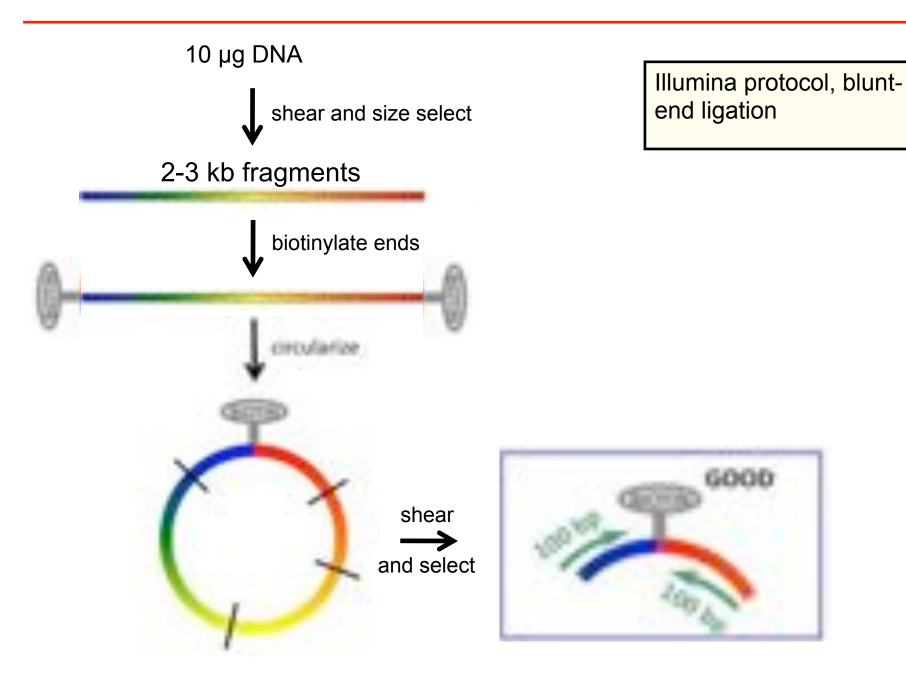
Libraries from 180 bp fragments

Pairs of 100 base reads from these libraries are merged to create 'reads' that are twice as long:



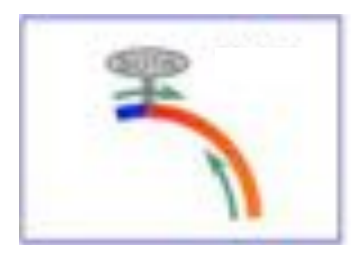
For longer reads, fragment size would be increased proportionally.

Short jumping libraries (2-3 kb)



Short jumping libraries (2-3 kb)

Problem 1. Read passes through circularization junction. This reduces the effective read length (and complicates algorithm).



What might be done to reduce incidence of this: shear circles to larger size and select larger fragments

Short jumping libraries (2-3 kb)

Problem 2. Reads come from nonjumped fragments and are thus in reverse orientation and close together on the genome. This reduces yield (and complicates algorithm).



Putative cause: original DNA is nicked or becomes nicked during process – biotins become 'ectopically' attached at these nicks



How to use ALLPATHS-LG

- 1. Data requirements
- 2. Computational requirements & installation
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- 5. What is an ALLPATHS-LG assembly?

Computational requirements

- 64-bit Linux
- runs multi-threaded on a single machine
- memory requirements
 - about 160 bytes per genome base, implying
 - need 512 GB for mammal (Dell R315, 48 processors, \$39,000)
 - need 1 GB for bacterium (theoretically)
 - 。 if coverage different than recommended, adjust...
 - potential for reducing usage
- wall clock time to complete run
 - 5 Mb genome → 1 hour (8 processors)

Installing ALLPATHS-LG

Web page:

http://www.broadinstitute.org/software/allpaths-lg/blog/

General instructions:

http://www.broadinstitute.org/science/programs/genome-biology/computational-rd/general-instructions-building-our-software

Getting the ALLPATHS-LG source

Our current system is to release code daily if it passes a test consisting of several small assemblies:

Download the latest build from:

ftp://ftp.broadinstitute.org/pub/crd/ALLPATHS/
Release-LG/

Unpack it:

% tar xzf allpathslg-39099.tar.gz

(substitute the latest revision id for 39099)

This creates a source code directory allpathslg-39099:

% cd allpathslg-39099

Building ALLPATHS-LG

Step one: ./configure

Options:

```
-prefix=<prefix path>
put binaries in <prefix path>/bin, else ./bin
```

Step two: make and make install

Options:

-j < n >

compile with n parallel threads

Step three: add bin directory to your path

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Preparing data for ALLPATHS-LG

Before assembling, prepare and import your read data.

ALLPATHS-LG expects reads from:

- At least one fragment library.
 - One should come from fragments of size ~180 bp.
 - This isn't checked but otherwise results will be bad.
- At least one jumping library.

<u>IMPORTANT</u>: use all the reads, including those that fail the Illumina purity filter (PF). These low quality reads may cover 'difficult' parts of the genome.

ALLPATHS-LG input format

ALLPATHS-LG can import data from: BAM, FASTQ, FASTA/QUALA or FASTB/QUALB files.

You must also provide two metadata files to describe them:

```
in_libs.csvdescribes the librariesin_groups.csvties files to libraries
```

FASTQ format: consists of records of the form

@<read name>

<sequence of bases, multiple lines allowed>

+

<sequence of quality scores, with Qn represented by ASCII code n+33, multiple
lines allowed>

Libraries – in_libs.csv (1 of 2)

For fragment libraries only

frag_size - estimated mean fragment size

frag_stddev - estimated fragment size std dev

For jumping libraries only

insert_size - estimated jumping mean insert size

insert_stddev - estimated jumping insert size std dev

These values determine how a library is used. If insert_size is ≥ 20000, the library is assumed to be a Fosmid jumping library.

paired - always 1 (only supports paired reads)read orientation - inward or outward.

Paired reads can either point towards each other, or away from each other. Currently fragment reads must be inward, jumping reads outward, and Fosmid jumping reads inward.

Libraries – in_libs.csv (2 of 2)

Reads can be trimmed to remove non-genomic bases produced by the library construction method:

```
genomic_start
genomic_end - inclusive zero-based range of read bases
to be kept; if blank or 0 keep all bases
```

Reads are trimmed in their original orientation.

Extra optional fields (descriptive only – ignored by ALLPATHS)

```
    project_name - a string naming the project.
    organism_name - the organism name.
    type - fragment, jumping, EcoP15I, etc.
```

EXAMPLE

```
library_name, type, paired, frag_size, frag_stddev, insert_size, insert_stddev, read_orientation, genomic_start, genomic_end Solexa-11541, fragment, 1, 180, 10, , , inward , Solexa-11623, jumping, 1, , , 3000, 500, outward 0, 25
```

Input files - in_groups.csv

Each line in in_groups.csv comma separated value file, corresponds to a BAM or FASTQ file you wish to import for assembly.

The library name must match the names in in_libs.csv.

```
- a unique nickname for this file
library_name
file_name
- the absolute path to the file
(should end in .bam or .fastq)
(use wildcards '?', '*' for paired fastqs)
```

Example:

```
group_name, library_name, file_name
302GJ, Solexa-11541, /seq/Solexa-11541/302GJABXX.bam
303GJ, Solexa-11623, /seq/Solexa-11623/303GJABXX.?.fastq
```

How to import assembly data files

```
PrepareAllPathsInputs.pl
IN_GROUPS_CSV=<in groups file>
IN_LIBS_CSV=<in libs file>
DATA_DIR=<full path of data directory>
PLOIDY=<ploidy, either 1 or 2>
PICARD_TOOLS_DIR=<picard tools directory>
```

- IN_GROUPS_CSV and IN_LIBS_CSV: optional arguments with default values ./in_groups.csv and ./in_libs.csv. These arguments determine where the data are found.
- DATA DIR: imported data will be placed here.
- PLOIDY: either 1 (for a haploid or inbred organism), or 2 (for a diploid organism) we have not tried to assemble organisms having higher ploidy!
- PICARD_TOOLS_DIR: path to Picard tools, for data conversion from BAM.

Putting it all together

1. Collect the BAM or FASTQ files that you wish to assemble. Create a in_libs.csv metadata file to describe your libraries and a in groups.csv metadata file to describe your data files.

2. Prepare input files

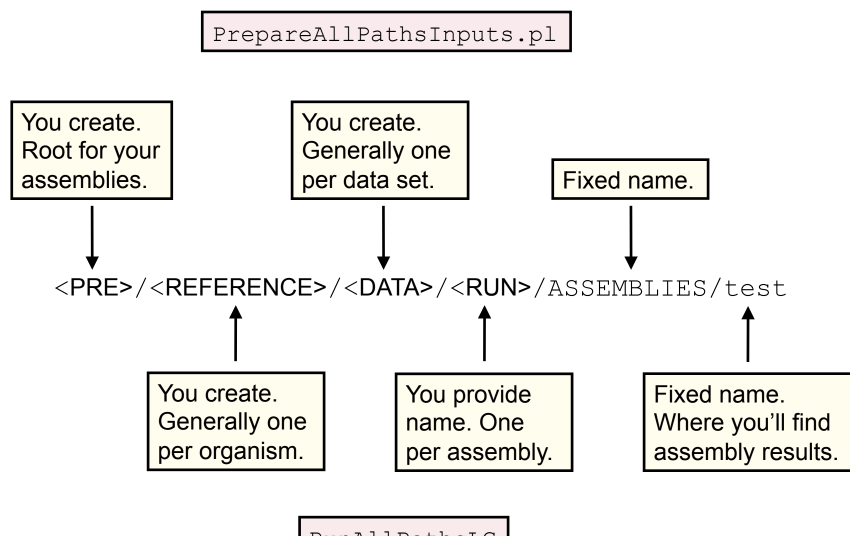
```
% cd asm
```

```
% PrepareAllPathsInputs.pl \
    DATA_DIR=`pwd` PLOIDY=1 >& prepare.log
```

How to use ALLPATHS-LG

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ALLPATHS-LG directory structure



RunAllPathsLG

How to assemble

Do this:

<u>Automatic resumption</u>. If the pipeline crashes, fix the problem, then run the same RunAllPathsLG command again. Execution will resume where it left off.

Results. The assembly files are:

```
    final.contigs.fasta - fasta contigs
    final.contigs.efasta - efasta contigs
    final.assembly.fasta - scaffolded fasta
    final.assembly.efasta - scaffolded efasta
```

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2. Prepare input files

```
% cd asm
% PrepareAllPathsInputs.pl \
    DATA DIR=`pwd` PLOIDY=1 >& prepare.log
```

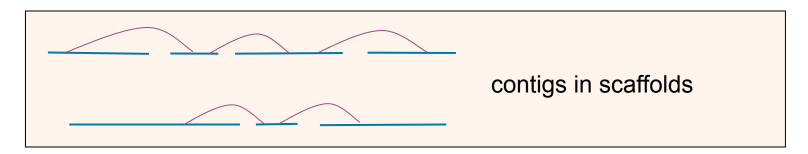
2. Assemble.

```
% RunAllPathsLG \
    PRE=/home REFERENCE_NAME=cshl \
    DATA_SUBDIR=asm RUN=default >& run.log
```

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1. 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)

1. Linear assemblies

Example of an assembly in fasta format

>scaffold 1

GGGATTTTGATGGGGTTTGCATTGAATCTGTAGATTGTCTTTGGTAAGATGGTTAGTTTTACTATGTTAATTCTGCCAAT CCACAAGCATGGGAGCGCTCTCCATTTTCTGAGATCTTCTTCAATTTCTTTGAGAAACTTGAAGTTATTGTCATACA >scaffold_2

3. Linearized graph assemblies

Efasta

...ACTGTTT{A,C}GAAAT... A or C at site

...CGCGTTTTTTTTT{,T,TT}CAT... 0 or 1 or 2 Ts at site

Example of an assembly in efasta format

>scaffold 1

GGGATTTTGATGGGGTTTGCATTGAATCTGTAGATTGTCTTTGGTAAGATGGTTAGTTTTACTATGTTAATTCTGCCAAT CCACAAGCATGGGAGCGCTCTCCATTTTCTGAGATCTTCTTCAATTTCTTTGAGAAACTTGAAGTTATTGTCATACA >scaffold 2

Putting it all together

1. Collect the BAM or FASTQ files that you wish to assemble. Create a in_libs.csv metadata file to describe your libraries and a in groups.csv metadata file to describe your data files.

2. Prepare input files

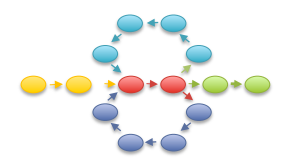
```
% cd asm
% PrepareAllPathsInputs.pl \
    DATA DIR=`pwd` PLOIDY=1 >& prepare.log
```

2. Assemble.

```
% RunAllPathsLG \
    PRE=/home REFERENCE_NAME=cshl \
    DATA_SUBDIR=asm RUN=default >& run.log
```

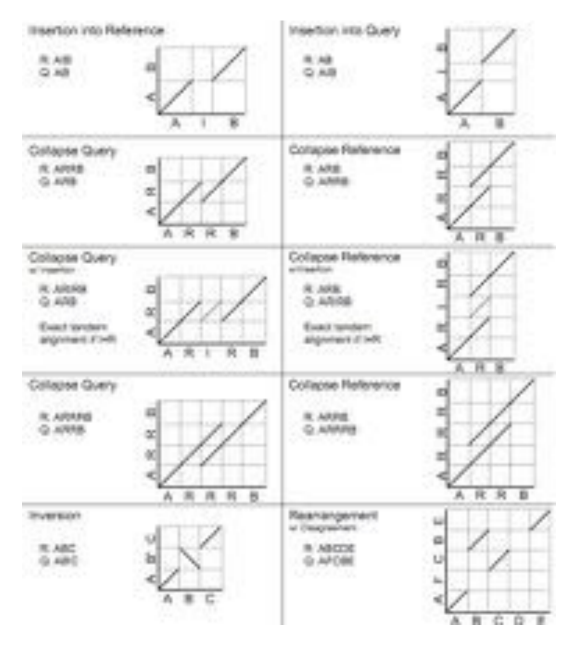
3. Get the results (four files).

```
% cd /default/ASSEMBLIES/test/
% less final.{assembly,contigs}.{fasta,efasta}
```



Whole Genome Alignment with MUMmer

SV Types



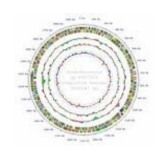
- Different structural variation types / misassemblies will be apparent by their pattern of breakpoints
- Most breakpoints will be at or near repeats
- Things quickly get complicated in real genomes

http://mummer.sf.net/manual/ AlignmentTypes.pdf

Find and decode

```
~/MUMmer3.23/nucmer -maxmatch AHBC01.fasta \
  default/ASSEMBLIES/test/final.contigs.fasta -p refctg
  -maxmatch Find maximal exact matches (MEMs) without repeat filtering
  -p refctg Set the output prefix for delta file
~/MUMmer3.23/mummerplot --layout refctq.delta
       Show the dotplot
~/MUMmer3.23/show-coords -rclo refctq.delta
       Sort alignments by reference position
  -c Show percent coverage
  -1 Show sequence lengths
  -o Annotate each alignment with BEGIN/END/CONTAINS
./samtools faidx default/ASSEMBLIES/test/final.contigs.fasta
./samtools faidx default/ASSEMBLIES/test/final.contigs.fasta \
   contig_XXX:YYY-ZZZ | ./dna-encode -d
```

Resources



Assembly Competitions

- Assemblathon: http://assemblathon.org/
- GAGE: http://gage.cbcb.umd.edu/

Assembler Websites:

- ALLPATHS-LG: http://www.broadinstitute.org/software/allpaths-lg/blog/
- SOAPdenovo: http://soap.genomics.org.cn/soapdenovo.html
- Celera Assembler: http://wgs-assembler.sf.net

Tools:

- MUMmer: http://mummer.sourceforge.net/
- Quake: http://www.cbcb.umd.edu/software/quake/
- AMOS: http://amos.sf.net

Questions?

http://schatzlab.cshl.edu/





