#### De novo assembly of complex genomes Michael Schatz

Sept 18, 2012 Statistical Bioinformatics, Purdue University





#### Outline

- I. Genome assembly by analogy
- 2. Hybrid error correction and assembly
- 3. Very recent sequencing results

#### Shredded Book Reconstruction

Dickens accidentally shreds the first printing of <u>A Tale of Two Cities</u>
– Text printed on 5 long spools

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- How can he reconstruct the text?
  - 5 copies x 138, 656 words / 5 words per fragment = 138k fragments
  - The short fragments from every copy are mixed together
  - Some fragments are identical



# **Greedy Reconstruction**



The repeated sequence make the correct reconstruction ambiguous

• It was the best of times, it was the [worst/age]

Model sequence reconstruction as a graph problem.

#### de Bruijn Graph Construction

- $G_k = (V, E)$ 
  - V = All length-k subfragments (k < l)
  - E = Directed edges between consecutive subfragments
    - Nodes overlap by k-1 words



- Locally constructed graph reveals the global sequence structure
  - Overlaps between sequences implicitly computed

de Bruijn, 1946 Idury and Waterman, 1995 Pevzner, Tang, Waterman, 2001



### de Bruijn Graph Assembly



# Counting Eulerian Tours $A \rightarrow B \rightarrow D$ ARBRCRDor ARCRBRD

Generally an exponential number of compatible sequences

- Value computed by application of the BEST theorem (Hutchinson, 1975)

$$\mathcal{W}(G,t) = (\det L) \left\{ \prod_{u \in V} (r_u - 1)! \right\} \left\{ \prod_{(u,v) \in E} a_{uv}! \right\}^{-1}$$
  
L = n x n matrix with  $r_u$ - $a_{uu}$  along the diagonal and  $-a_{uv}$  in entry uv  
 $r_u = d^+(u) + l$  if  $u = t$ , or  $d^+(u)$  otherwise  
 $a_{uv}$  = multiplicity of edge from u to v

Assembly Complexity of Prokaryotic Genomes using Short Reads. Kingsford C, Schatz MC, Pop M (2010) *BMC Bioinformatics*.

### N50 size

Def: 50% of the genome is in contigs as large as the N50 value



```
N50 size = 30 \text{ kbp}
```

```
(300k+100k+45k+45k+30k = 520k \ge 500kbp)
```

Note:

N50 values are only meaningful to compare when base genome size is the same in all cases

# **Assembly Applications**

Novel genomes





#### Metagenomes





#### Sequencing assays

- Transcript assembly
- Structural variations
- Haplotype analysis





#### Why are genomes hard to assemble?

- **I.** Biological:
  - (Very) High ploidy, heterozygosity, repeat content

#### 2. Sequencing:

- (Very) large genomes, imperfect sequencing

#### **3.** Computational:

- (Very) Large genomes, complex structure

#### **4.** Accuracy:

- (Very) Hard to assess correctness



# Ingredients for a good assembly



#### High coverage is required

- Oversample the genome to ensure every base is sequenced with long overlaps between reads
- Biased coverage will also fragment assembly





#### Reads & mates must be longer than the repeats

- Short reads will have *false overlaps* forming hairball assembly graphs
- With long enough reads, assemble entire chromosomes into contigs

#### Errors obscure overlaps

- Reads are assembled by finding kmers shared in pair of reads
- High error rate requires very short seeds, increasing complexity and forming assembly hairballs

**Current challenges in de novo plant genome sequencing and assembly** Schatz MC, Witkowski, McCombie, WR (2012) *Genome Biology*. 12:243

### Hybrid Sequencing





**Illumina** Sequencing by Synthesis

High throughput (60Gbp/day) High accuracy (~99%) Short reads (~100bp)

#### Pacific Biosciences

SMRT Sequencing

Lower throughput (600Mbp/day) Lower accuracy (~85%) Long reads (1-2kbp+)

# **SMRT** Sequencing

Imaging of florescent phospholinked labeled nucleotides as they are incorporated by a polymerase anchored to a Zero-Mode Waveguide (ZMW).







Time

http://www.pacificbiosciences.com/assets/files/pacbio\_technology\_backgrounder.pdf



#### • Standard sequencing

- Long inserts so that the polymerase can synthesize along a single strand

#### • Circular consensus sequencing

- Short inserts, so polymerase can continue around the entire SMRTbell multiple times and generate multiple sub-reads from the same single molecule.

# SMRT Sequencing Data

#### Yeast (Pre-release Chemistry / 2010)

65 SMRT cells 734,151 reads after filtering Mean: 642.3 +/- 587.3 Median: 553 Max: 8,495





Sample of 100k reads aligned with BLASR requiring >100bp alignment Average overall accuracy: 83.7%, 11.5% insertions, 3.4% deletions, 1.4% mismatch





**Read Position** 

#### Consistent quality across the entire read

- Uniform error rate, no apparent biases for GC/motifs
- Sampling artifacts at beginning and ends of alignments

### Consensus Quality: Probability Review

Roll *n* dice => What is the probability that at least half are 6's

n	Min to Lose	Losing Events	P(Lose)
I		1/6	16.7%
2		P(lof 2) + P(2 of 2)	30.5%
3		P(2 of 3) + P(3 of 3)	7.4%
4		P(2 of 4) + P(3 of 4) + P(4 of 4)	13.2%
5		P(3 of 5) + P(4 of 5) + P(5 of 5)	3.5%
n	ceil(n/2)	$\sum_{i=\lceil n/2 \rceil}^{n} P(i \ of \ n) = \sum_{i=\lceil n/2 \rceil}^{n} \binom{n}{i} (p)^{i} (1-p)^{n-i}$	

# **Consensus Accuracy and Coverage**



#### Coverage can overcome random errors

- Dashed: error model from binomial sampling; solid: observed accuracy
- For same reason, CCS is extremely accurate when using 5+ subreads

$$CNS Error = \sum_{i=\lceil c/2 \rceil}^{c} \binom{c}{i} (e)^{i} (1-e)^{n-i}$$

# PacBio Error Correction

http://wgs-assembler.sf.net

- I. Correction Pipeline
  - I. Map short reads (SR) to long reads (LR)
  - 2. Trim LRs at coverage gaps
  - 3. Compute consensus for each LR



2. Error corrected reads can be easily assembled, aligned



Hybrid error correction and de novo assembly of single-molecule sequencing reads. Koren, S, Schatz, MC, et al. (2012) *Nature Biotechnology*. doi:10.1038/nbt.2280

#### **Error Correction Results**



Correction results of 20x PacBio coverage of E. coli K12 corrected using 50x Illumina

#### Celera Assembler

http://wgs-assembler.sf.net

- I. Pre-overlap
  - Consistency checks
- 2. Trimming
  - Quality trimming & partial overlaps
- 3. Compute Overlaps
  - Find high quality overlaps
- 4. Error Correction
  - Evaluate difference in context of overlapping reads
- 5. Unitigging
  - Merge consistent reads
- 6. Scaffolding
  - Bundle mates, Order & Orient
- 7. Finalize Data
  - Build final consensus sequences



# **SMRT-Assembly Results**



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Hybrid assembly results using error corrected PacBio reads Meets or beats Illumina-only or 454-only assembly in every case

### Improved Gene Reconstruction



FOXP2 assembled on a single contig

# Transcript Alignment



- Long-read single-molecule sequencing has potential to directly sequence full length transcripts
  - Raw reads and raw alignments (red) have many spurious indels inducing false frameshifts and other artifacts
  - Error corrected reads almost perfectly match the genome, pinpointing splice sites, identifying alternative splicing
- New collaboration with Gingeras Lab looking at splicing in human

#### PacBio Technology Roadmap



Internal Roadmap has made steady progress towards improving read length and throughput

Very recent improvements:

I. Improved enzyme: Maintains reactions longer

- "Hot Start" technology: Maximize subreads
  - . MagBead loading: Load longest fragments

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# **Preliminary Rice Assemblies**



In collaboration with McCombie & Ware labs @ CSHL

# Single Molecule Sequencing Summary

PacBio RS has capabilities not found in any other technology

- Substantially longer reads -> span repeats
- Unbiased sequence coverage -> close sequencing gaps
- Single molecule sequencing -> haplotype phasing, alternative splicing

Long reads enables highest quality de novo assembly

- Longer reads have more information than shorter reads
- Because the errors are random we can compensate for them
- One chromosome, one contig achieved in microbes

Exciting developments on the horizon

- Longer reads, higher throughput PacBio
- Nanopore Sequencing



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# Thank You!

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### Long Read Advantages



(a) Long reads close sequencing gaps

(b) Long readsassemble acrosslong repeats

(c) Long reads span complex microsatellites

#### **Theoretical Benefits of Hot Start Sequencing**



#### Magnetic Bead Enzyme-Template Complex Loading



Multiple complexes attached to magnetic beads that are much larger than individual ZMWs

Rotate magnet to evenly disperse beads across entire chip surface



Pre-Deposition: Complex loaded beads in solution

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(II) Introduce magnet: Bead complexes pulled to chip surface

#### MBS (MagBead Station)



Improvements to Sample Prep

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