### Evaluating Assembly Quality Michael Schatz

Dec 8, 2014 USDA/ARS Workshop





## Outline

- I. Assembly review
  - I. Assembly by analogy
  - 2. Causes of Mis-assemblies

### 2. Evaluating Assembly Quality

- I. Assemblathon
- 2. Size Statistics
- 3. Mate-pair Happiness
- 4. CEGMA

3. RNA-seq specific challenges

### Shredded Book Reconstruction

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

It was	s thevbesthef	bes <b>tinfes</b> nite	syais tilaes toloristor	of times,	it was the	a <b>ggebf</b>	v <b>isisolom</b> it	itwavashe	abe aga	ofistolistanes	as,
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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 the assembly problem as a graph problem

### de Bruijn Graph Construction

- $D_k = (V, E)$ 
  - 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







# 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

# Typical sequencing coverage



Imagine raindrops on a sidewalk

We want to cover the entire sidewalk but each drop costs \$1









# Genome Coverage Distribution



• Standard Deviation = sqrt(cov)

This is the mathematically model => reality may be much worse

- Double your coverage for diploid genomes
- Can use somewhat lower coverage in a population to find common variants





- After simplification and correction, compress graph down to its non-branching initial contigs
  - Aka "unitigs", "unipaths"
  - Unitigs end because of (1) lack of coverage, (2) errors, (3) heterozygosity/isoform differences, and (4) repeats





# Repetitive regions

Repeat Type	Definition / Example	Prevalence
Low-complexity DNA / Microsatellites	$(b_1b_2b_k)^N$ where $I \le k \le 6$ CACACACACACACACACACACA	2%
SINEs (Short Interspersed Nuclear Elements)	<i>Alu</i> sequence (~280 bp) Mariner elements (~80 bp)	13%
LINEs (Long Interspersed Nuclear Elements)	~500 – 5,000 bp	21%
LTR (long terminal repeat) retrotransposons	Ту I -copia, Ту3-gypsy, Pao-BEL (~100 – 5,000 bp)	8%
Other DNA transposons		3%
Gene families & segmental duplications		4%

- Over 50% of mammalian genomes are repetitive
  - Large plant genomes tend to be even worse
  - Wheat: I6 Gbp; Pine: 24 Gbp

### The full tale

... it was the best of times it was the worst of times ...
... it was the age of wisdom it was the age of foolishness ...
... it was the epoch of belief it was the epoch of incredulity ...
... it was the season of light it was the season of darkness ...
... it was the spring of hope it was the winder of despair ...



### Errors in the graph





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- Attempt to answer the question:
   "What makes a good assembly?"
- Organizers provided sequence data to assembly experts around the world
  - Assemblathon 1:~100Mbp simulated genome
  - Assemblathon 2:3 vertebrate genomes each ~IGB
- Results demonstrate trade-offs assemblers must make

Assemblathon I:A competitive assessment of de novo short read assembly methods. Earl, DA, et al. (2011) Genome Research. doi: 10.1101/gr.126599.111

Assemblathon 2: Evaluating de novo methods of genome assembly in three vertebrate species Bradnam, KR. et al (2013) GigaScience 2:10 doi:10.1186/2047-217X-2-10

# **Assembly Results**



### Scaffold Paths





# Final Rankings

ID	Overall	CPNG50	SPNG50	Struct.	CC50	Subs.	Copy. Num.	Cov. Tot.	Cov. CDS
BGI	36	\$						\$	
Broad	37		*	*	\$			1.0.0	
WTSI-S	46		\$		*	\$			
CSHL	52	*							2
BCCGSC	53								
DOEJGI	56		\$	$\overrightarrow{\mathbf{x}}$		*			
RHUL	58								
WTSI-P	64						1		
EBI	64						\$		
CRACS	64					23			

- ALLPATHS and SOAPdenovo came out neck-and-neck followed closely behind by Celera Assembler, SGA, and ABySS
- My recommendation for "typical" short read assembly is to use ALLPATHS
- Single molecule sequencing becoming extremely attractive if you have access

### N50 size

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



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

### A greater N50 is indicative of improvement in every dimension:

- Better resolution of genes and flanking regulatory regions
- Better resolution of transposons and other complex sequences
- Better resolution of chromosome organization
- Better sequence for all downstream analysis



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# Estimating coverage with Kmers



## Estimating coverage with Kmers



## QC: Read Coverage



### Wheat Genome (A. tauschi / CSHL)



### Heterozygous Genome



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## **Assembly Validation**



Automatically scan an assembly to locate misassembly signatures for further analysis and correction

### Assembly-validation pipeline

- I. Evaluate Mate Pairs & Libraries
- 2. Evaluate Read Alignments
- 3. Evaluate Read Breakpoints
- 4. Analyze Depth of Coverage



#### **Genome Assembly forensics: finding the elusive mis-assembly.** Phillippy, AM, Schatz, MC, Pop, M. (2008) *Genome Biology* 9:R55.

# Paired-end and Mate-pairs

#### Paired-end sequencing

- Read one end of the molecule, flip, and read the other end
- Generate pair of reads separated by up to 500bp with inward orientation



### Mate-pair sequencing

- Circularize long molecules (1-10kbp), shear into fragments, & sequence
- Mate failures create short paired-end reads



### C/E Statistic

- The presence of individual compressed or expanded mates is rare but expected.
- Do the inserts spanning a given position differ from the rest of the library?
  - Flag large differences as potential misassemblies
  - Even if each individual mate is "happy"
- Compute the statistic at all positions
  - (Local Mean Global Mean) / Scaling Factor
- Introduced by Jim Yorke's group at UMD
Forensics

### Sampling the Genome





### C/E-Statistic: Expansion







### C/E-Statistic: Compression

**Forensics** 



8 inserts: 3.2 kb-4.8kb C/E Stat: (3488-4000) = -3.62 C/E Stat  $\leq$  -3.0 indicates Compression

6kb



Hawkeye & AMOS: Visualizing and assessing the quality of genome assemblies Schatz, M.C. et al. (2011) Briefings in Bioinformatics. doi: 10.1093/bib/bbr074

## Long Read Sequencing Technology



## O. sativa pv Indica (IR64)

Genome size: ~370 Mb Chromosome N50: ~29.7 Mbp



Assembly	Contig NG50	80000			HGAP F	Read Le	ngths
MiSeq Fragments 25x 456bp (3 runs 2x300 @ 450 FLASH)	I9 kbp	60000			22.7x <b>(disca</b>	53,652 over 10 <i>rded re</i>	kbp <b>ads</b>
"ALLPATHS-recipe" 50x 2x100bp @ 180 36x 2x50bp @ 2100 51x 2x50bp @ 4800	18 kbp	20000 40000			Delov	v 8500b	<i>(</i> <b>)</b>
HGAP 22.7x @ 10kbp	4.0 Mbp	0 -			Manna		
Nipponbare BAC-by-BAC Assembly	5.1 Mbp		10000	20000	30000	40000	50000

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### Detection and Correction with Quake

#### I. Count all "Q-mers" in reads

- Fit coverage distribution to mixture model of errors and regular coverage
- Automatically decide threshold for trusted k-mers

#### 2. Correction Algorithm

- Consider editing erroneous kmers into trusted kmers in decreasing likelihood
- Includes quality values, nucleotide/ nucleotide substitution rate



Quake: quality-aware detection and correction of sequencing reads. Kelley, DR, Schatz, MC, Salzberg, SL (2010) Genome Biology. 11:R116

# Gene Analysis with CEGMA

- Defined a set of 248 "core eukaryotic genes" (CEGs)
  - Highly conserved and in low copy numbers across all known eukaryotic species
  - House keeping genes and other basic functions

- Developed a robust alignmentbased search tool to seek out those genes in your new assembly
  - Your ability to discover these 248 CEGs is highly correlated with finding the rest of the genes in the genome



#### Assessing the gene space in draft genomes Parra, G, Bradnam, B, Ning, Z, Keane, T, Korf, I (2009) 37(1) 289–297. doi:10.1093/nar/gkn916



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## **RNA-seq** Overview



### **RNA-seq Overview**



# **RNA-seq Challenges**



#### Challenge I: Eukaryotic genes are spliced

Solution: Use a spliced aligner, and assemble isoforms

**TopHat: discovering spliced junctions with RNA-Seq.** Trapnell et al (2009) *Bioinformatics*. 25:0 1105-1111



Challenge 2: Read Count != Transcript abundance Solution: Infer underlying abundances (e.g. FPKM)

**Transcript assembly and quantification by RNA-seq** Trapnell et al (2010) *Nat. Biotech.* 25(5): 511-515



#### Challenge 3: Transcript abundances are stochastic

Solution: Replicates, replicates, and more replicates

**RNA-seq differential expression studies: more sequence or more replication?** Liu et al (2013) *Bioinformatics*. doi:10.1093/bioinformatics/btt688

### Assembly Summary



Assembly quality depends on

- I. Coverage: low coverage is mathematically hopeless
- 2. Repeat composition: high repeat content is challenging
- 3. Read length: longer reads help resolve repeats
- 4. Error rate: errors reduce coverage, obscure true overlaps
- Assembly is a hierarchical, starting from individual reads, build high confidence contigs/unitigs, incorporate the mates to build scaffolds
  - Extensive error correction is the key to getting the best assembly possible from a given data set
- Watch out for collapsed repeats & other misassemblies
  - Globally/Locally reassemble data from scratch with better parameters & stitch the 2 assemblies together







