Genome Sequencing & Assembly Michael Schatz

Nov. 17, 2014 CSHL Adv. Sequencing Course





Outline

- I. Assembly theory
 - I. Assembly by analogy
 - 2. De Bruijn and Overlap graph
 - 3. Coverage, read length, errors, and repeats
- 2. Whole Genome Alignment
 - I. Aligning & visualizing with MUMmer
- 3. Genome assemblers
 - I. ALLPATHS-LG: recommended for Illumina-only projects
 - 2. Celera Assembler: recommended for long read projects
- 4. Summary & Recommendations

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 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

de Bruijn Graph Assembly



de Bruijn Graph Assembly



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 ...



Milestones in Genome Assembly

Nature Vol. 265 February 24 1977

articles

Nucleotide sequence of bacteriophage Φ X174 DNA

F. Sanger, G. M. Air', B. G. Barrell, N. L. Brown', A. R. Coulson, J. C. Fiddes, C. A. Hutchison III', P. M. Slocombe' & M. Smith' MR Labourso of Meladur Belag, Hill Read, Candidge C22 201, UK

a DNA supervise for the generator of heartrophysics 95/174 of opportunity, 53/57 melocitoshi has been diversiond using the rapid and simple 'plus and wisua' method. The apparent identifications of the protein source of the protein expansion, including initiation and accompaniens, founding initiation and accompaniens, founding initiation and accompaniens for the protein and RAAs. Two pairs of geness are cooled by the source regarding for the source regions of DAA using different reading frames.	straid DNA of SNA has the same sequence as the nRNA and certain conditions, with Bast indexees so that a pretex- fragment can be soluted and sequenced. Only one maps is associated, by comparison with the nervo acid sequence data without of the gene C protective (positions 2,262,243). At this stage sequencing conditions, using premet system with DNA polynemizes were being developed ² and Schor and other sequencing scheroses using premet system with DNA polynemizes were being developed ²¹ and Schor part of the robuscner beining site. The was used to pettern for part of the robuscner beining site. The was used to pettern for the state of the sequence being scheroses.
This generate of bacterisphage Φ Xi14 is a single-strended, densitie DNA of approximately 5.600 materials and the first sine known proteins. The order of these genes, as determined by genetic techniques ¹⁻¹ , is <i>i i i i i i i i i i</i>	the intercistronic region between the F and G genes, using DN, polymerase and ¹⁰ P-labelled triphophatters ¹¹ . The ribo-subbili icon sechesique ¹¹ facilitated the sequence determination of the labelled DNA produced. This decaracidestic-period systems was also used to develop the plus and misus method' Satable synthetic primers are, however, difficult to prepare and a

1977. Sanger *et al.* Ist Complete Organism 5375 bp



2000. Myers *et al.* Ist Large WGS Assembly. Celera Assembler. 116 Mbp



1995. Fleischmann *et al.* 1st Free Living Organism TIGR Assembler. 1.8Mbp



1998. C.elegans SC Ist Multicellular Organism BAC-by-BAC Phrap. 97Mbp





2001.Venter *et al.*, IHGSC Human Genome Celera Assembler/GigaAssembler. 2.9 Gbp

A CONTRACTOR OF CONTRACTOR OF

2010. Li *et al.* Ist Large SGS Assembly. SOAPdenovo 2.2 Gbp

Like Dickens, we must computationally reconstruct a genome from short fragments

Assembly Applications

Novel genomes





• Metagenomes





- Sequencing assays
 - Structural variations
 - Transcript assembly





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

Illumina Sequencing by Synthesis



1. Prepare

2. Attach

DNA fragment

Dense lawn of primers





4. Image

Metzker (2010) Nature Reviews Genetics 11:31-46 http://www.youtube.com/watch?v=I99aKKHcxC4









5. Basecall

Typical sequencing coverage



Imagine raindrops on a sidewalk

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









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.

Formulation comes from the limit of the binomial equation

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.





Unitigging / Unipathing

- 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, 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



- If *n* reads are a uniform random sample of the genome of length *G*, we expect $k=n\Delta/G$ reads to start in a region of length Δ .
 - If we see many more reads than k (if the arrival rate is > A), it is likely to be a collapsed repeat

$$\Pr(X - copy) = \binom{n}{k} \left(\frac{X\Delta}{G}\right)^k \left(\frac{G - X\Delta}{G}\right)^{n-k} \qquad A(\Delta, k) = \ln\left(\frac{\Pr(1 - copy)}{\Pr(2 - copy)}\right) = \ln\left(\frac{\frac{(\Delta n/G)^k}{k!}e^{\frac{-\Delta n}{G}}}{\frac{(2\Delta n/G)^k}{k!}e^{\frac{-2\Delta n}{G}}}\right) = \frac{n\Delta}{G} - k\ln 2$$

The fragment assembly string graph Myers, EW (2005) Bioinformatics. 21 (suppl 2): ii79-85.

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



Scaffolding

- Initial contigs (aka unipaths, unitigs) terminate at
 - Coverage gaps: especially extreme GC
 - Conflicts: errors, repeat boundaries
- Use mate-pairs to resolve correct order through assembly graph
 - Place sequence to satisfy the mate constraints
 - Mates through repeat nodes are tangled
- Final scaffold may have internal gaps called sequencing gaps
 - We know the order, orientation, and spacing, but just not the bases. Fill with Ns instead





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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Whole Genome Alignment with MUMmer

Slides Courtesy of Adam M. Phillippy University of Maryland



• For two genomes, A and B, find a mapping from each position in A to its corresponding position in B



Not so fast...

 Genome A may have insertions, deletions, translocations, inversions, duplications or SNPs with respect to B (sometimes all of the above)



WGA visualization

- How can we visualize *whole* genome alignments?
- With an alignment dot plot T $-N \times M$ matrix G• Let i = position in genome A• Let j = position in genome B• Fill cell (*i*,*j*) if A_i shows similarity to B_j A



 A perfect alignment between A and B would completely fill the positive diagonal





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



Alignment of 2 strains of Y. pestis http://mummer.sourceforge.net/manual/



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Genome assembly with ALLPATHS-LG Iain MacCallum



How ALLPATHS-LG works



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**

*See next slide.

**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.
To close a read pair (red), we require the existence of another read pair (blue), overlapping perfectly like this:



More than one closure allowed (but rare).

I. Find 'seed' unipaths, evenly spaced across genome (ideally long, of copy number CN = 1)

II. Form neighborhood around each seed



and are extended by other unipaths



Population structure of Oryza sativa

Indica

Total Span: 344.3 Mbp Contig N50: 22.2kbp

Aus

Total Span: 344.9Mbp Contig N50: 25.5kbp

Nipponbare

Total Span: 354.9Mbp Contig N50: 21.9kbp

Whole genome de novo assemblies of three divergent strains of rice (O. sativa) documents novel gene space of aus and indica Schatz, MC, Maron, L, Stein, et al (2014) In press.

Pan-genomics of draft assemblies

Strategy:

- I. Align the genomes to each other (MUMmer)
- Identify segments of genome A that do not align anywhere to genome B (BEDTools)
- \rightarrow Megabases specific to each genome!!!!
- 3. Screen regions that fail to align with their k-mer frequencies (jellyfish)
 - In reality, "Genome specific regions" averaged over 10,000x kmer coverage while unique regions were ~50x
- \rightarrow 100s of KB specific to each genome!!!



Reference-free kmer analysis



IR64 - Sub1A (A-2) Kmer Coverage

Draft assemblies are difficult to conclusively analyze to determine if a given sequence is truly specific to one genome or another

- The sequence may be mis-assembled (or incompletely assembled in the other genome)
- Use k-mer analysis to rule out misassemblies
- Here we see the SubIA (A-2) locus present only in IR64

Strain specific regions





(B) IR64

IR_201	IR_299 IR	_382 IR_408	IR_479	IR_712	IR_918	IR_977	IR_1091	IR_1261
No No	N DE S	Red DI123 IND DI123 DI123	Dut23	NB INNA DU123	NIP IREA DU123	NIP IREA DU123	NIP IREA DJ123	Np Np
-			-	1		3	-	-

(C) DJ123

	D	J_1	82	_)J_	92	D	U_1	132	D.	1_1	85	D	U_2	289	D.	J_3	18	D	J_3	28	DJ	_90	03	DJ	_12	66	DJ	_13	92
111	đΝ	IREA	DJ123	Np	1P864	DJ123	NID	IREA	DU123	^o N	IBGA	DU123	Np	INN	DU123	2	1964	DU123	ş	R64	D1123	ł	1964	DU123	2	R64	D1123	2	R64	DU123
11															-															

Oryza sativa Gene Diversity



Overall sequence content

In each sector, the top number is the total number of base pairs, the middle number is the number of exonic bases, and the bottom is the gene count. If a gene is partially shared, it is assigned to the sector with the most exonic bases.

Genic content

In each sector, the top number is the median CDS length, the middle number is the average number of exons per gene, and the bottom is the percentage InterPro/homology.

Strain specific regions

- Very high quality representation of the "gene-space"
 - Overall identity ~99.9%
 - Less than 1% of exonic bases missing
- Genome-specific genes enriched for disease resistance
 - Reflects their geographic and environmental diversity
 - Detailed analysis of the S5 hybrid sterility locus, the Sub I submergence tolerance locus, the LRK gene cluster associated with improved yield, and the Pup I cluster associated with phosphorus deficiency
- Assemblies fragmented at (high copy) repeats
 - Missing regions have mean k-mer coverage >10,000x
 - Difficult to identify full length gene models and regulatory features



Genome assembly with the Celera Assembler

Assembly Complexity





Assembly Complexity



The advantages of SMRT sequencing Roberts, RJ, Carneiro, MO, Schatz, MC (2013) *Genome Biology*. 14:405

Long Read Sequencing Technology



PacBio SMRT Sequencing

Imaging of fluorescently 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

SMRT Sequencing Data



Match	83.7%
Insertions	11.5%
Deletions	3.4%
Mismatch	1.4%

TTGTAAGCAGTTGAAAACTATGTGT <mark>G</mark> GATTTAG <mark>A</mark> ATAAAGAACATG <mark>A</mark> AAG
ATTATAAA-CAGTTGATCCATT-AGAAGA-AAACGCAAAAGGC <mark>G</mark> GCTAGG
CAACCTTGAATGTAATCGCACTTGAAGAACAAGATTTTATTCCGCGCCCC
T <mark>A</mark> ACGAATC <mark>A</mark> AGATTCTGAAAACA <mark>C</mark> AT-AT <mark>AACA</mark> ACCTCCAAAA-CACAA
–AGGAGG <mark>GGAAAGGGGGG</mark> GAATATCT–AT <mark>A</mark> AAAGATTACAAATT <mark>A</mark> GA–TGA
ACT-AATTCACAA <mark>T</mark> A-AATAACACTTTTA-ACA <mark>G</mark> AATTGAT-GGAA-GTT
TC <mark>G</mark> GAGAGATCC <mark>A</mark> AAACAAT <mark>G</mark> GGC-ATCG <mark>C</mark> CTTTGA-GTTAC-AATCAAA
ATCCAGT <mark>G</mark> GAAAATATA <mark>AT</mark> TTATGC <mark>A</mark> ATCCA <mark>G</mark> GAACTTATTCACAATTAG

Sample of 100k reads aligned with BLASR requiring >100bp alignment

PacBio Assembly Algorithms

PacBioToCA

PBJelly

Gap Filling and Assembly Upgrade

English et al (2012) PLOS One. 7(11): e47768

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Hybrid/PB-only Error Correction

Koren, Schatz, et al (2012) Nature Biotechnology. 30:693–700



PB-only Correction & Polishing

Chin et al (2013) Nature Methods. 10:563–569

< 5x

PacBio Coverage



Consensus Accuracy and Coverage



Coverage can overcome random errors

- Dashed: error model from binomial sampling
- Solid: observed accuracy

Koren, Schatz, et al (2012) Nature Biotechnology. 30:693–700

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

S. pombe dg21

PacBio RS II sequencing at CSHL

 Size selection using an 7 Kb elution window on a BluePippin[™] device from Sage Science



S. pombe dg21

ASM294 Reference sequence

• 12.6Mbp; 3 chromo + mitochondria; N50: 4.53Mbp

PacBio assembly using HGAP + Celera Assembler

• 12.7Mbp; 13 non-redundant contigs; N50: 3.83Mbp; >99.98% id





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O. sativa pv Indica (IR64)

PacBio RS II sequencing at PacBio

 Size selection using an 10 Kb elution window on a BluePippin[™] device from Sage Science





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 (disca	53,652 over 10 <i>rded re</i>	kbp ads
"ALLPATHS-recipe" 50x 2x100bp @ 180 36x 2x50bp @ 2100 51x 2x50bp @ 4800	I8 kbp	20000 40000			Delov	w 8500k	<i>)</i> () (
HGAP 22.7x @ 10kbp	4.0 Mbp	0 -			Manua		
Nipponbare BAC-by-BAC Assembly	5.1 Mbp		10000	20000	30000	40000	50000

Human CHMI

http://blog.pacificbiosciences.com/2014/02/data-release-54x-long-read-coverage-for.html





Chromosome N50: 90.5 Mbp

Genome size:

Average read length:

3.0 Gb

7,680 bp

CHMI hert sequenced at PacBio

- Sequenced using the P5 enzyme and C3 chemistry
- Size selection using an 20kb elution window on a BluePippin[™] device from Sage Science
- Total coverage: 54x

Sum of Contig Lengths:	3.2 Gb
N50 Contig Length:	4.38 Mbp
Max Contig:	44 Mbp

High quality draft assembly Assembly Performance: 4.38Mbp/90.5Mbp = 4.5% Sanger HuRef assembly: 107kbp / 90.5Mbp = .1%

Current Collaborations



PacBio® Advances in Read Length





Lee, H*, Gurtowski, J*, Yoo, S, Marcus, S, McCombie, WR, Schatz, MC *http://www.biorxiv.org/content/early/2014/06/18/006395*

Oxford Nanopore MinION





- Thumb drive sized sequencer
 powered over USB
- Capacity for 512 reads at once
- Senses DNA by measuring changes to ion flow



What should we expect from an assembly?

Analysis of dozens of genomes from across the tree of life with real and simulated data

Summary & Recommendations

- < 100 Mbp: HGAP/PacBio2CA @ 100x PB C3-P5 expect near perfect chromosome arms
- HGAP/PacBio2CA @ 100x PB C3-P5 < IGB:high quality assembly: contig N50 over IMbp
- hybrid/gap filling > IGB: expect contig N50 to be 100kbp – 1Mbp
- Email mschatz@cshl.edu > 5GB:





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

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<u>CSHL</u>

Hannon Lab Gingeras Lab Jackson Lab Hicks Lab **Iossifov Lab** Levy Lab Lippman Lab Lyon Lab Martienssen Lab McCombie Lab Tuveson Lab Ware Lab Wigler Lab

Pacific Biosciences Oxford Nanopore





National Human Genome Research Institute



SFARI SIMONS FOUNDATION AUTISM RESEARCH INITIATIVE



Thank you http://schatzlab.cshl.edu @mike_schatz