### De novo assembly of complex genomes Michael Schatz

Sept 10, 2013 UIUC



### Schatz Lab Overview





### Outline

- I. Genome assembly by analogy
- 2. Hybrid error correction and assembly
- 3. De novo mutations in autism



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### 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 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

## 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 ...





# Reducing Complexity



Longer reads span more repeats, simplifying the assembly problem

- Idealized assembly of B. anthracis reduces to a single contig with 5kb reads
- Exact improvement depends on the specific genome

#### The advantages of SMRT sequencing

Roberts, RJ, Carneiro, MO, Schatz, MC (2013) Genome Biology. 14:405

## 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:

A "good" N50 size is a moving target relative to other recent publications. 10-20kbp contig N50 is currently a typical value for most "simple" genomes.



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# **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



**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 (IGbp/day) Lower accuracy (~85%) Long reads (5kbp+)

# **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



#### • 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.
- Barbell sequence: ATCTCTCTCttttcctcctccgttgttgttgttGAGAGAGAT

# 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 
CAACCTTGAATGTAATCGCACTTGAAGAACAAGATTTTATTCCGCGCCCG 
TAACGAATCAAGATTCTGAAAACACAT-ATAACAACCTCCAAAA-CACAA 
–AGGAGG <mark>GGAAAGGGGGG</mark> GAATATCT–AT <mark>A</mark> AAAGATTACAAATT <mark>A</mark> GA–TGA 
ACT-AATTCACAATA-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 Error Correction

http://wgs-assembler.sf.net

- I. Correction Pipeline
  - I. Map short reads to long reads
  - 2. Trim long reads at coverage gaps
  - 3. Compute consensus for each long read



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

### Enhanced PacBio Error Correction

https://github.com/jgurtowski/pbtools

#### PacBioToCA fails in complex regions

- I. Simple Repeats Kmer Frequency Too High to Seed Overlaps
- 2. GC Rich Regions Known Illumina Bias
- 3. Error Dense Regions Difficult to compute overlaps with many errors



**Position Specific Coverage and Error Rate** 



### **Correction with Unitigs**



#### Unitigs:

High quality contigs formed from unambiguous, unique overlaps of reads

#### Illumina reads -> Illumina unitigs -> Map and error correct PacBio reads -> Assemble PacBio reads

Can Help us overcome:

- 1. Simple Repeats Kmer Frequency Too High to Seed Overlaps
- 2. GC Rich Regions Known Illumina Bias
- 3. Error Dense Regions Difficult to compute overlaps with many errors

# Population structure in Oryza sativa

3 varieties selected for de novo sequencing

#### **IR64**



High quality BAC-by-BAC reference

- ~370 Mbp genome in 12 chromosomes
- About 40% repeats:
  - Many 4-8kbp repeats
  - 300kbp max high identity repeat (99.99%)
- Useful model for other cereal genomes

#### Nipponbare



# **Preliminary Rice Assemblies**

Assembly	Contig NG50
HiSeq Fragments 50x 2x100bp @ 180	3,925
MiSeq Fragments 23x 459bp 8x 2x251bp @ 450	6,332
"ALLPATHS-recipe" 50x 2x100bp @ 180 36x 2x50bp @ 2100 51x 2x50bp @ 4800	18,248



In collaboration with McCombie & Ware labs @ CSHL

## Improved Gene Reconstruction

FOXP2 assembled in a single contig in the PacBio parrot assembly



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

## Long Read CNV Analysis

Aluminum tolerance in maize is important for drought resistance and protecting against nutrient deficiencies

- Segregating population localized a QTL on a BAC, but unable to genotype with Illumina sequencing because of high repeat content and GC skew
- Long read PacBio sequencing corrected by CCS reads revealed a triplication of the ZnMATEI membrane transporter



#### A rare gene copy-number variant that contributes to maize aluminum tolerance and adaptation to acid soils

Maron, LG et al. (2012) PNAS. doi: 10.1073/pnas.1220766110

# 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

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

#### PacBio Technology Roadmap



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

Very recent improvements:

Improved enzyme:
 Maintains reactions longer

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



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# Assembly Coverage Model







Simulate PacBio-like reads to predict how the assembly will improve as we add additional coverage

Only 8x coverage is needed to sequence every base in the genome, but 40x improves the chances repeats will be spanned by the longest reads

#### Assembly complexity of long read sequencing Lee, H, Gurtowski, J, Marcus, S., Schatz MC et al. (2013) In preparation

# Speculation for 2013



Doubling the average read length dramatically improves the assembly quality

 Able to span a larger repeats and lock contigs together

Expect to see contig N50 values over 1Mbp very soon, even in very complicated plant and animal species

Megabase contig N50 already routine in microbial assembly with PacBio sequencing

# Speculation for 2013



With PacBio-like reads averaging 11.2kbp (4x current), we should be able to assemble almost every chromosome arm of rice into single contigs

 The 300kbp near perfect repeat is the only exception

Even with the current assembly, we are seeing new genes and other sequences missing in the "high quality" BAC-by-BAC reference genome

# Assembly Complexity of Long Reads



# De novo assembly of Arabidopsis

http://blog.pacificbiosciences.com/2013/08/new-data-release-arabidopsis-assembly.html



#### A. thaliana Ler-0 sequenced at PacBio

- Sequenced using the latest P4 enzyme and C2 chemistry
- Size selection using an 8 Kb to 50 Kb elution window on a BluePippin<sup>™</sup> device from Sage Science
- Total coverage >100x

Genome size:	124.6 Mb
GC content:	33.92%
Raw data:	II Gb
Assembly coverage:	15x over 9kbp

Sum of Contig Lengths:	149.5Mb
Number of Contigs:	1788
Max Contig Length:	12.4 Mb
N50 Contig Length:	8.4 Mb



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# Variation Detection Complexity



Analysis confounded by sequencing errors, localized repeats, allele biases, and mismapped reads

## Scalpel: Haplotype Microassembly

G. Narzisi, J. O'Rawe, I. Iossifov, Y. Lee, Z. Wang, G. Lyon, M. Wigler, and M. C. Schatz

DNA sequence **micro-assembly** pipeline for accurate detection and validation of *de novo* mutations (SNPs, indels) within exome-capture data.



- I. Combine mapping and assembly
- 2. Exhaustive search of haplotypes
- 3. De novo mutations



NRXN1 *de novo* SNP (auSSC12501 chr2:50724605)



#### **Scalpel Pipeline**



#### Repeats in the Genome



Variant Read: Large deletion or critical snp?

### Experimental Analysis & Validation

Selected one deep coverage exome for deep analysis

- 80% of the target at >20x coverage
- Evaluated with Scalpel, SOAPindel, and GATK Haplotype Caller

1000 indels selected for validation

- 200 Scalpel
- 200 GATK Haplotype Caller
- 200 SOAPindel
- 200 within the intersection
- 200 long indels (>30bp)



#### Scalpel Indel Discovery



**Detection of de novo mutations in exome-capture data using micro-assembly** Narzisi et al. (2013) In preparation

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# Exome sequencing of the SSC



Sequencing of 343 families from the Simons Simplex Collection

- Parents plus one child with autism and one non-autistic sibling
- Enriched for higher-functioning individuals

Families prepared and captured together to minimize batch effects

- Exome-capture performed with NimbleGen SeqCap EZ Exome v2.0 targeting 36 Mb of the genome.
- ~80% of the target at >20x coverage with ~93bp reads

**De novo gene disruptions in children on the autism spectrum** lossifov et al. (2012) Neuron. 74:2 285-299

#### De novo mutation discovery and validation

**Concept**: Identify mutations not present in parents.

**Challenge**: Sequencing errors in the child or low coverage in parents lead to false positive de novos



#### **Ref:** ... TCAGAACAGCTGGATGAGATCTTAGCCAACTACCAGGAGATTGTCTTTGCCCGGA...

- Father: ...TCAGAACAGCTGGATGAGATCTTAGCCAACTACCAGGAGATTGTCTTTGCCCCGGA...
- Mother: ...TCAGAACAGCTGGATGAGATCTTAGCCAACTACCAGGAGATTGTCTTTGCCCCGGA...
- Sib: ...TCAGAACAGCTGGATGAGATCTTAGCCAACTACCAGGAGATTGTCTTTGCCCGGA...
- Aut(1): ...TCAGAACAGCTGGATGAGATCTTAGCCAACTACCAGGAGATTGTCTTTGCCCGGA...
- Aut(2): ...TCAGAACAGCTGGATGAGATCTTA<u>C</u>C----CC<u>G</u>GGAGATTGTCTTTGCCCCGGA...

#### 6bp heterozygous deletion at chr13:25280526 ATP12A

#### De novo Genetics of Autism

- In 343 family quads so far, we see significant enrichment in de novo likely gene killers in the autistic kids
  - Overall rate basically I:I (432:396)
  - 2:1 enrichment in nonsense mutations
  - 2:1 enrichment in frameshift indels
  - 4:1 enrichment in splice-site mutations
  - Most de novo originate in the paternal line in an age-dependent manner (56:18 of the mutations that we could determine)
- Observe strong overlap with the 842 genes known to be associated with fragile X protein FMPR
  - Related to neuron development and synaptic plasticity
  - Also strong overlap with chromatin remodelers

De novo gene disruptions in children on the autism spectrum lossifov et al. (2012) Neuron. 74:2 285-299







- Hybrid assembly let us combine the best characteristics of 2<sup>nd</sup> and 3<sup>rd</sup> gen sequencing
  - Long reads and good coverage are the keys to a good de novo assembly
  - Single contig de novo assemblies of entire microbial chromosomes are now routine; Single contig de novo assemblies of entire plant and animal chromosomes on the horizon
- Assembly is the missing link towards high accuracy indel mutation discovery
  - Allows the algorithm to break free from the expectations of the reference
  - Pinpointing de novo mutations require both high sensitivity and specificity
- We are starting to apply these technologies to discover significant biology that is otherwise impossible to measure

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National Human Genome Research Institute





Thank You! http://schatzlab.cshl.edu @mike\_schatz

