

De novo assembly of complex genomes

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Outline

1. 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 A Tale of Two Cities
 - Text printed on 5 long spools

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	best	of	times,	it	was	the	worst	of	times,	it	was	the	age	of	wisdom,	it	was	the	age	of	foolishness, ...	
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	best	of	times,	it	was	the	worst	of	times,	it	was	the	age	of	wisdom,	it	was	the	age	of	foolishness, ...	
It	was	the	best	of	times,	it	was	the	worst	of	times,	it	was	the	age	of	wisdom,	it	was	the	age	of	foolishness, ...

- 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

It was the best of
age of wisdom, it was
best of times, it was
it was the age of
it was the age of
it was the worst of
of times, it was the
of times, it was the
of wisdom, it was the
the age of wisdom, it
the best of times, it
the worst of times, it
times, it was the age
times, it was the worst
was the age of wisdom,
was the age of foolishness,
was the best of times,
was the worst of times,
wisdom, it was the age
worst of times, it was

It was the best of
was the best of times,
the best of times, it
best of times, it was
of times, it was the
of times, it was the
times, it was the worst
times, it was the age

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

Original Fragment

It was the best of

Directed Edge

It was the best → was the best of

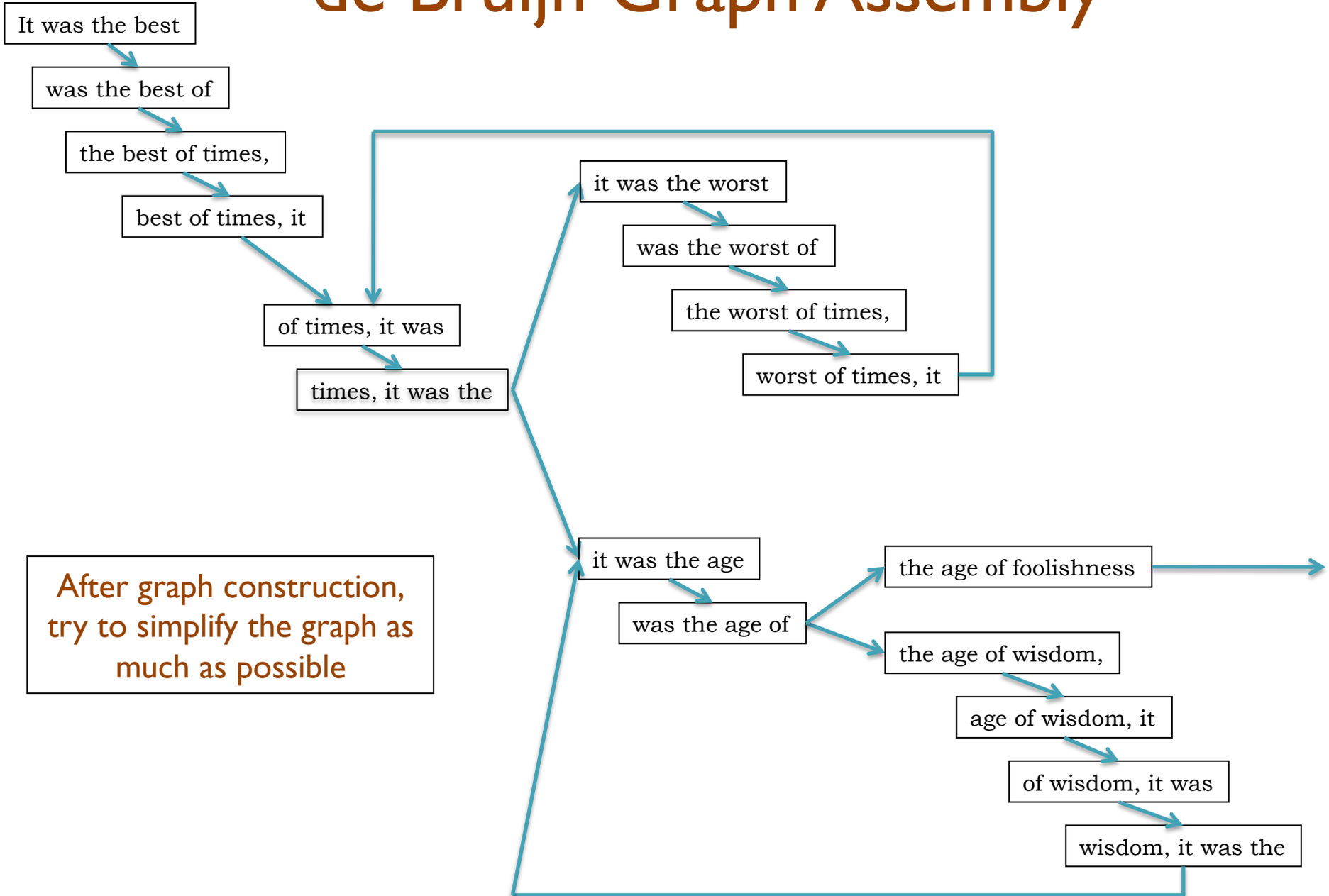
- Locally constructed graph reveals the global sequence structure
 - Overlaps between sequences implicitly computed

de Bruijn, 1946

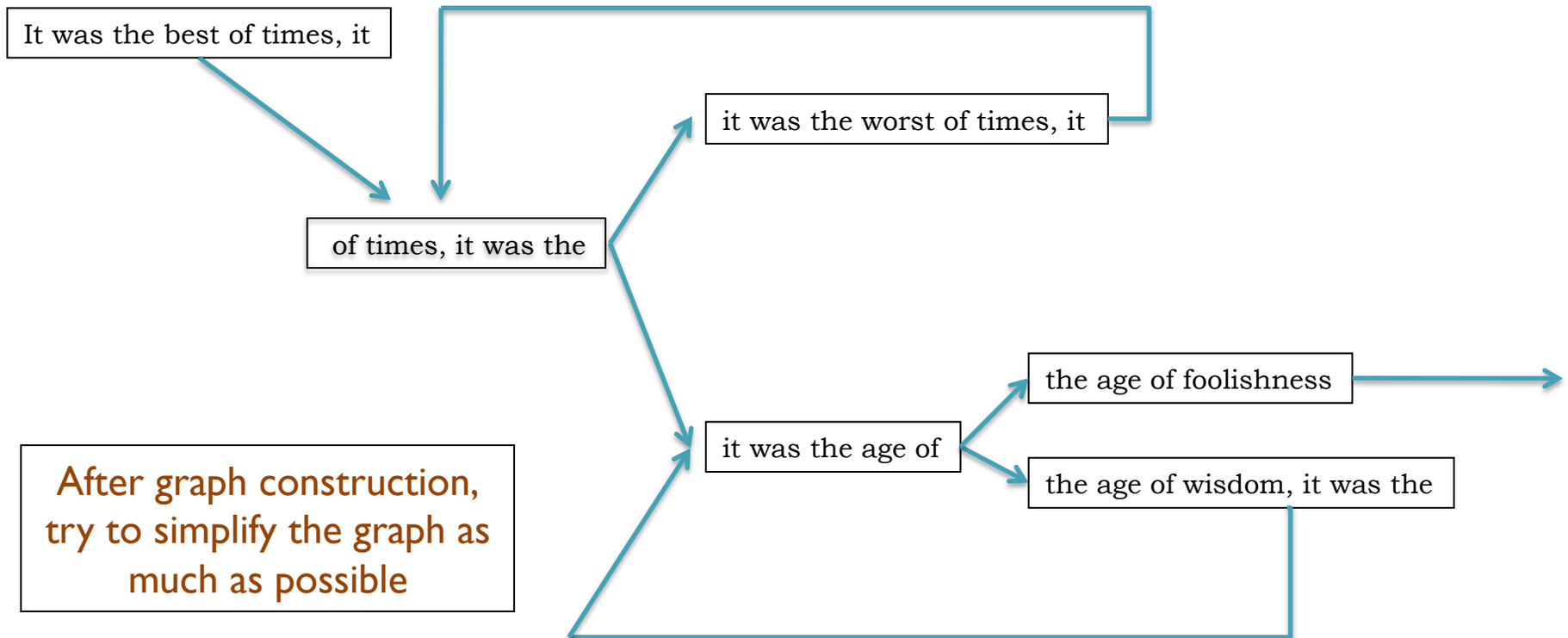
Idury and Waterman, 1995

Pevzner, Tang, Waterman, 2001

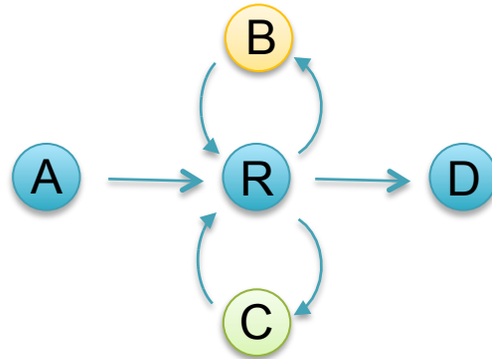
de Bruijn Graph Assembly



de Bruijn Graph Assembly



Counting Eulerian Tours



AR**B**RCRD
or
ARC**R**BRD

Generally an exponential number of compatible sequences

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

$$W(G, t) = (\det L) \left\{ \prod_{u \in V} (r_u - 1)! \right\} \left\{ \prod_{(u,v) \in E} a_{uv}! \right\}^{-1}$$

$L = n \times n$ matrix with $r_u - a_{uu}$ along the diagonal and $-a_{uv}$ in entry uv

$r_u = d^+(u) + 1$ 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

Example: 1 Mbp genome



N50 size = 30 kbp

$(300k + 100k + 45k + 45k + 30k = 520k \geq 500kbp)$

Note:

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

Assembly Applications

Novel genomes

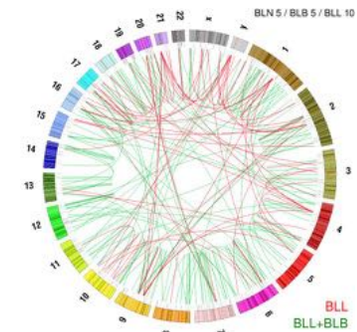
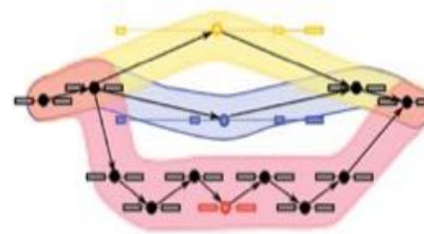


Metagenomes



Sequencing assays

- Transcript assembly
- Structural variations
- Haplotype analysis
- ...



Why are genomes hard to assemble?

1. Biological:

- (Very) High ploidy, heterozygosity, repeat content

2. Sequencing:

- (Very) large genomes, imperfect sequencing

3. Computational:

- (Very) Large genomes, complex structure

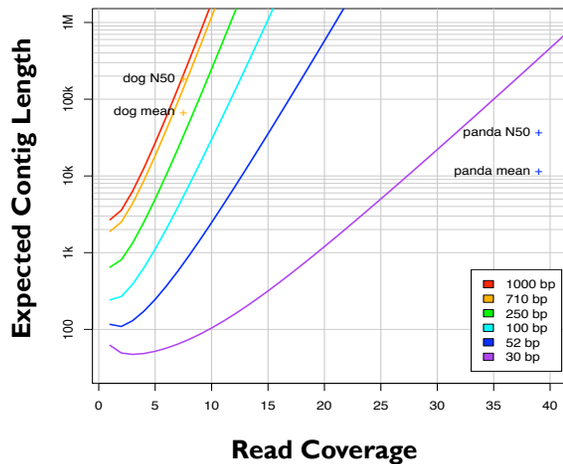
4. Accuracy:

- (Very) Hard to assess correctness



Ingredients for a good assembly

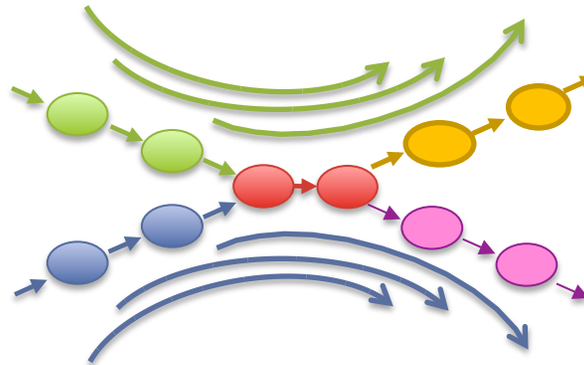
Coverage



High coverage is required

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

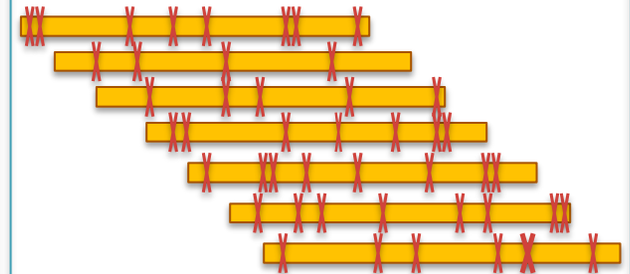
Read Length



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

Quality



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, WVR (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

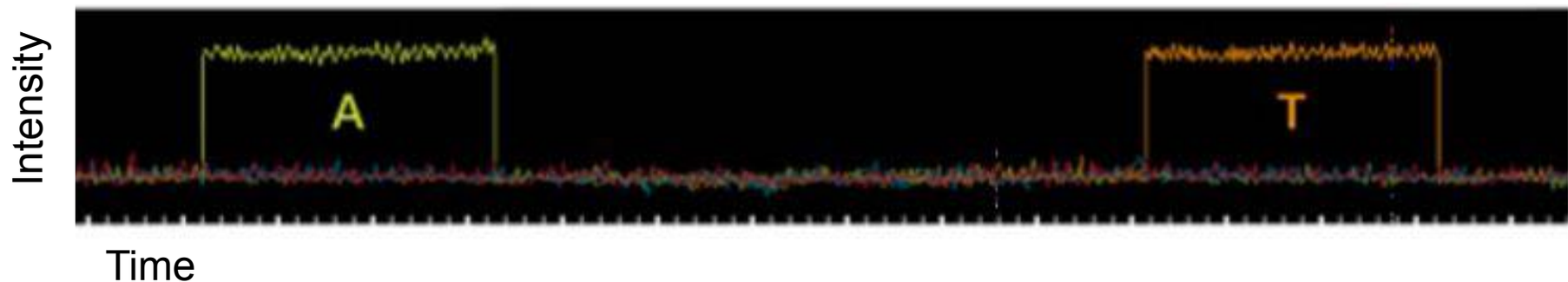
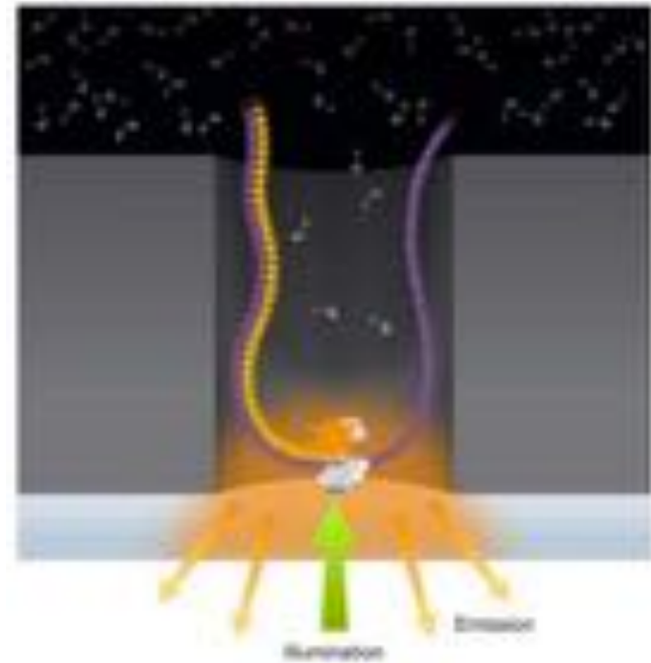
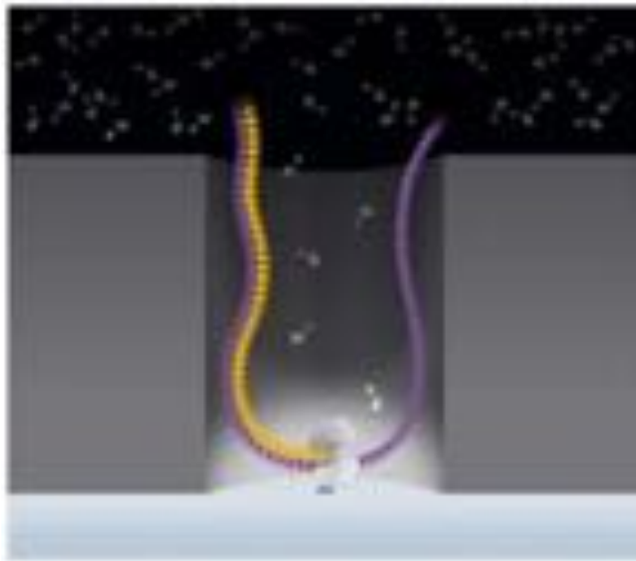
Lower throughput (600Mbp/day)

Lower accuracy (~85%)

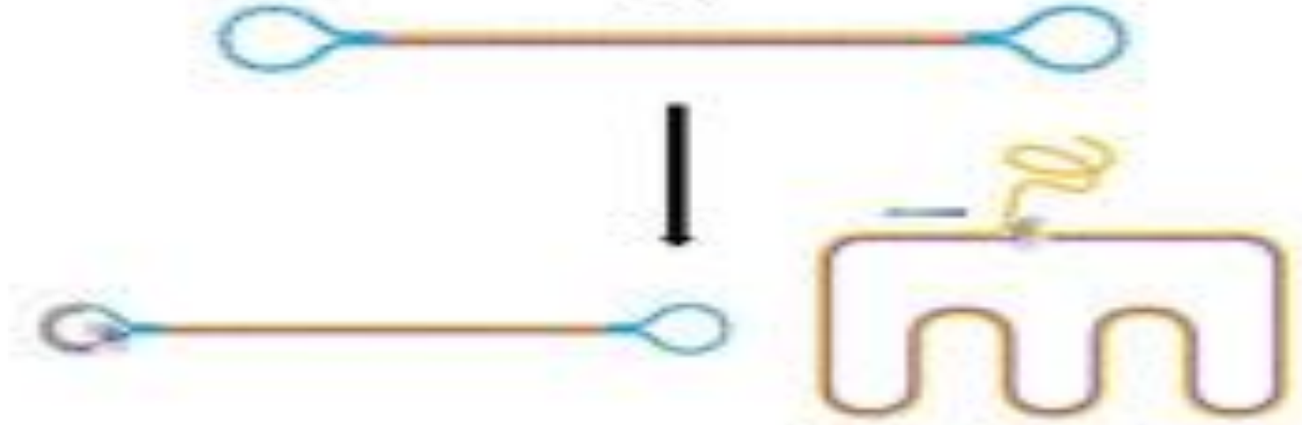
Long reads (1-2kbp+)

SMRT Sequencing

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

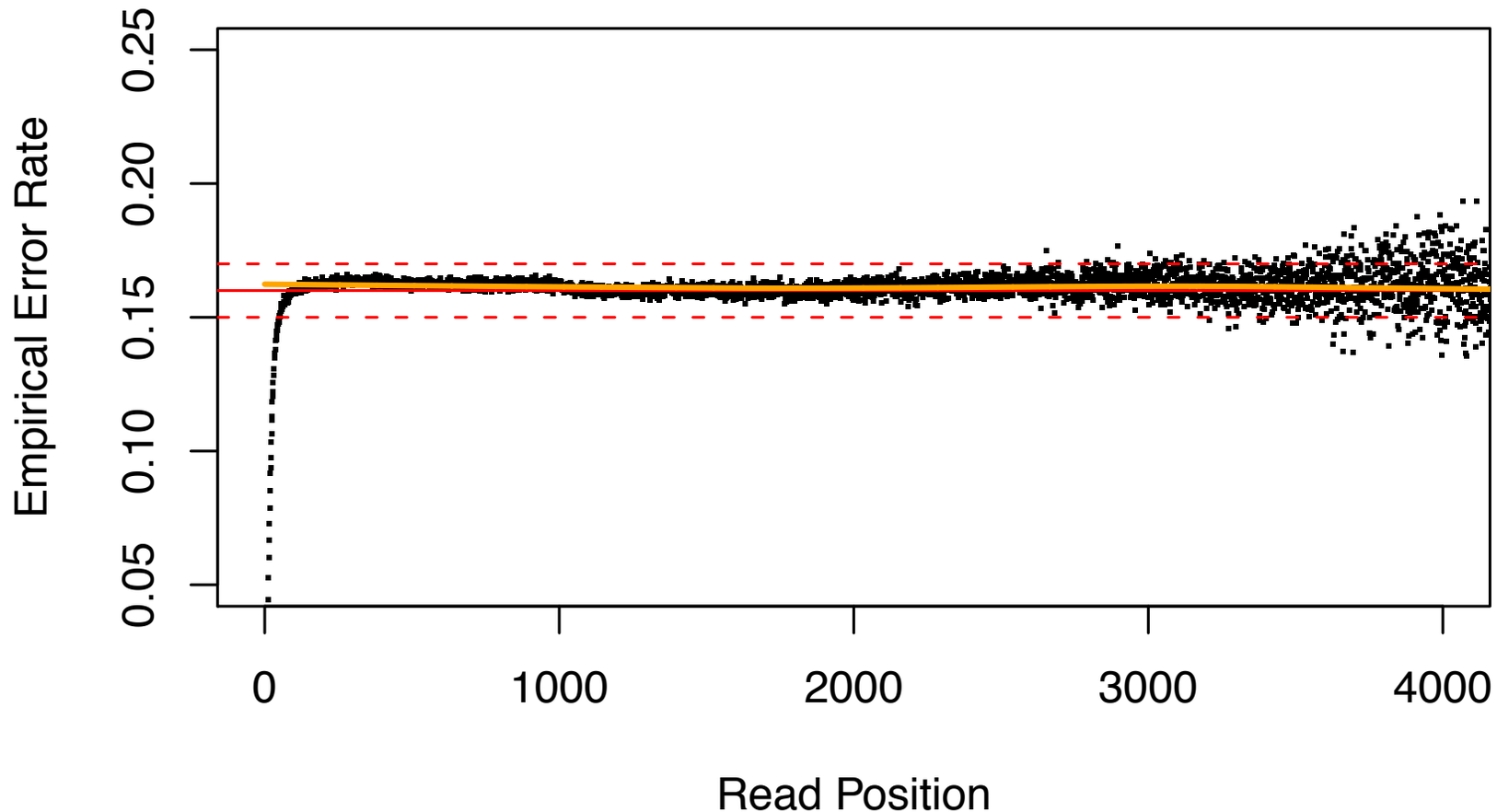


SMRT Read Types



- **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.

Read Quality



Consistent quality across the entire read

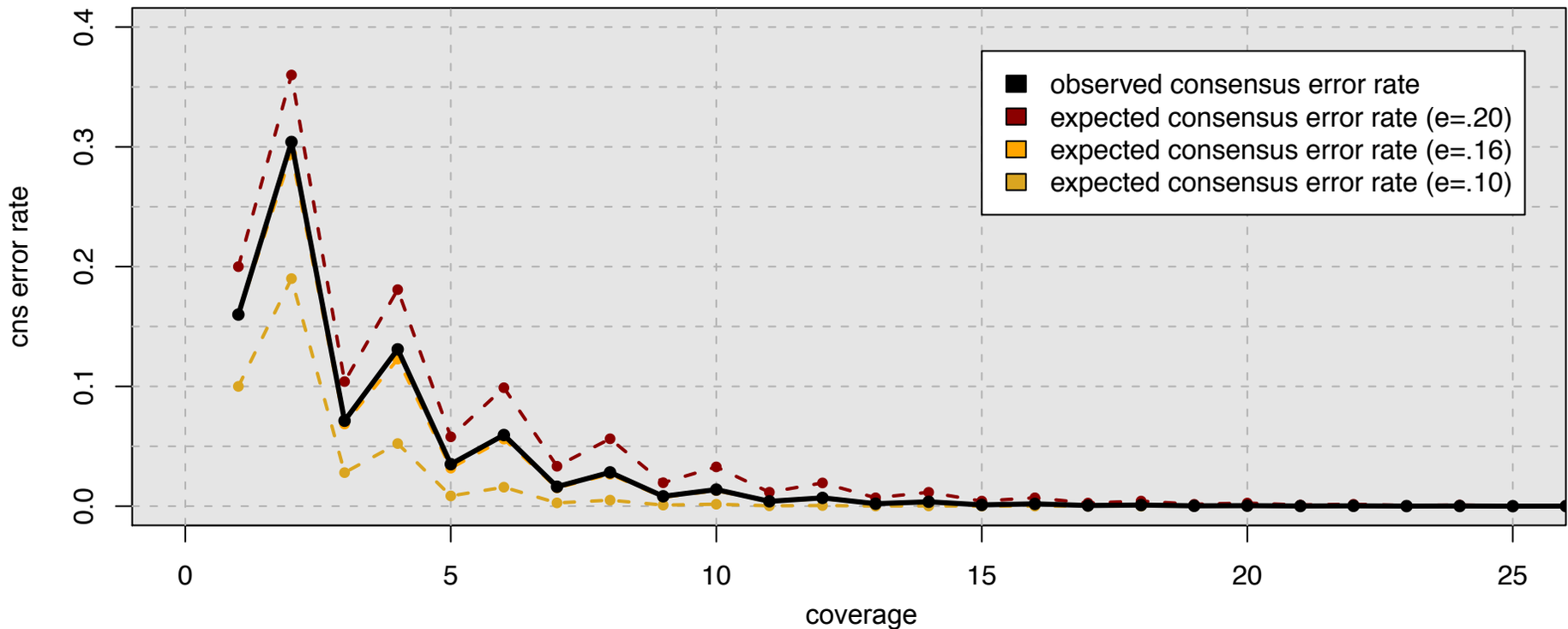
- Uniform error rate, no apparent biases for GC/motifs
- Sampling artifacts at beginning and ends of alignments

Consensus Quality: Probability Review

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

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

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

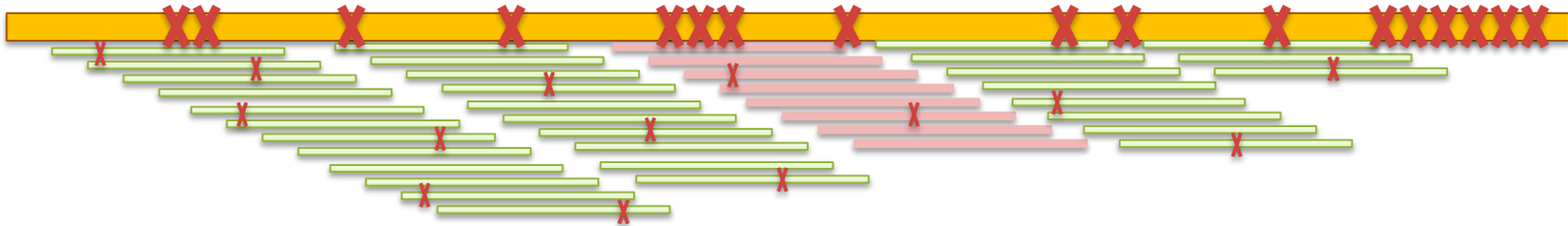
<http://wgs-assembler.sf.net>



I. Correction Pipeline

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

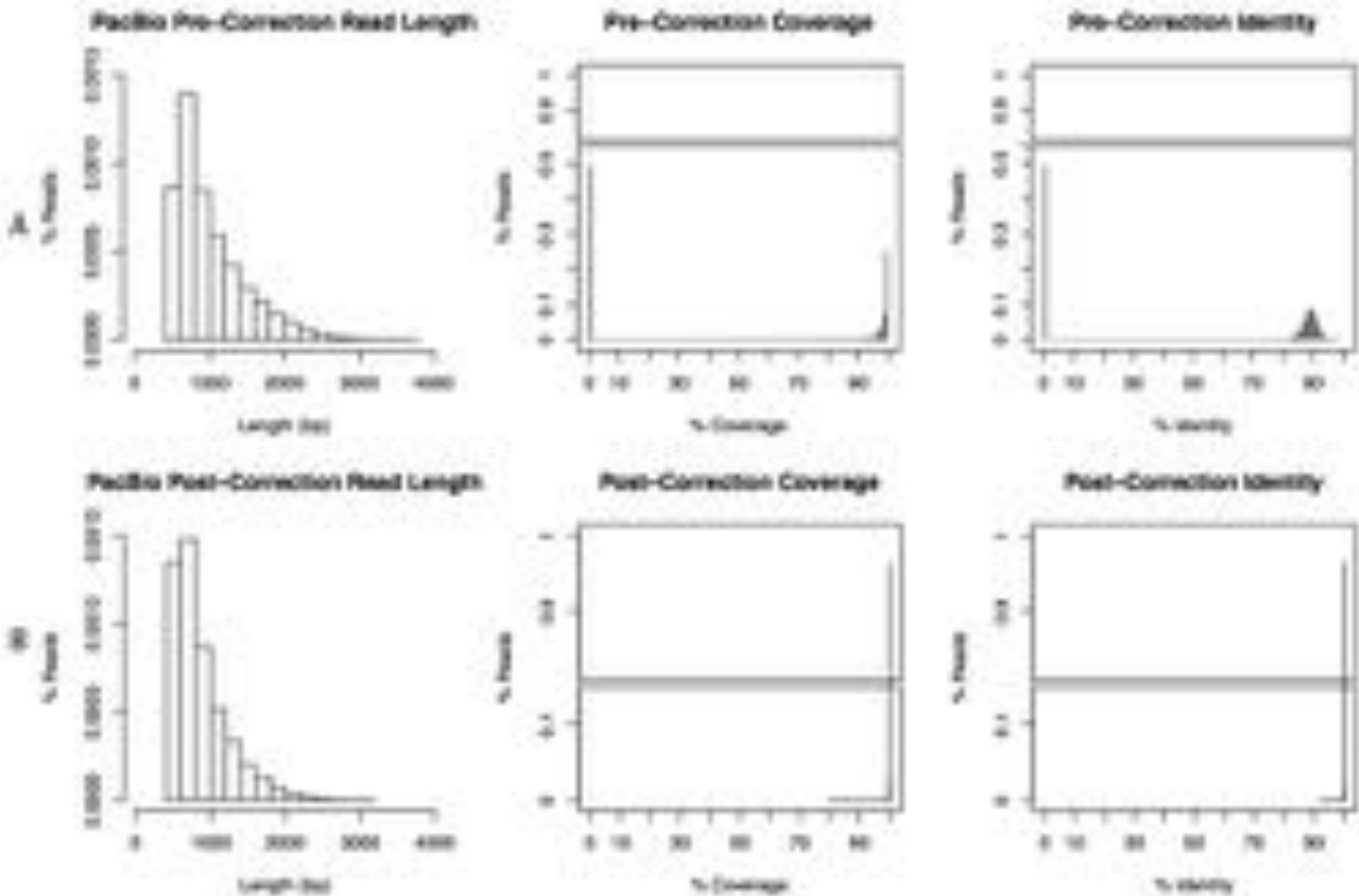
2. Error corrected reads can be easily assembled, aligned



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

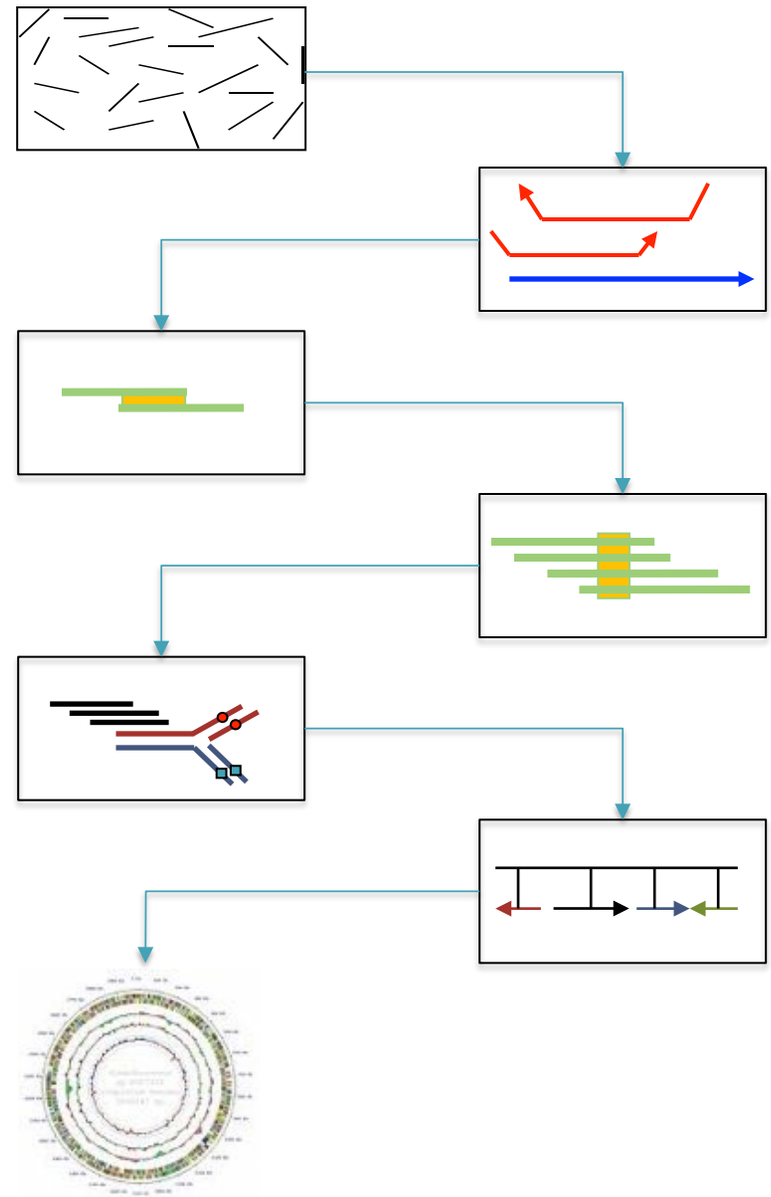


Correction results of 20x PacBio coverage of E. coli K12 corrected using 50x Illumina

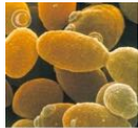
Celera Assembler

<http://wgs-assembler.sf.net>

1. 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



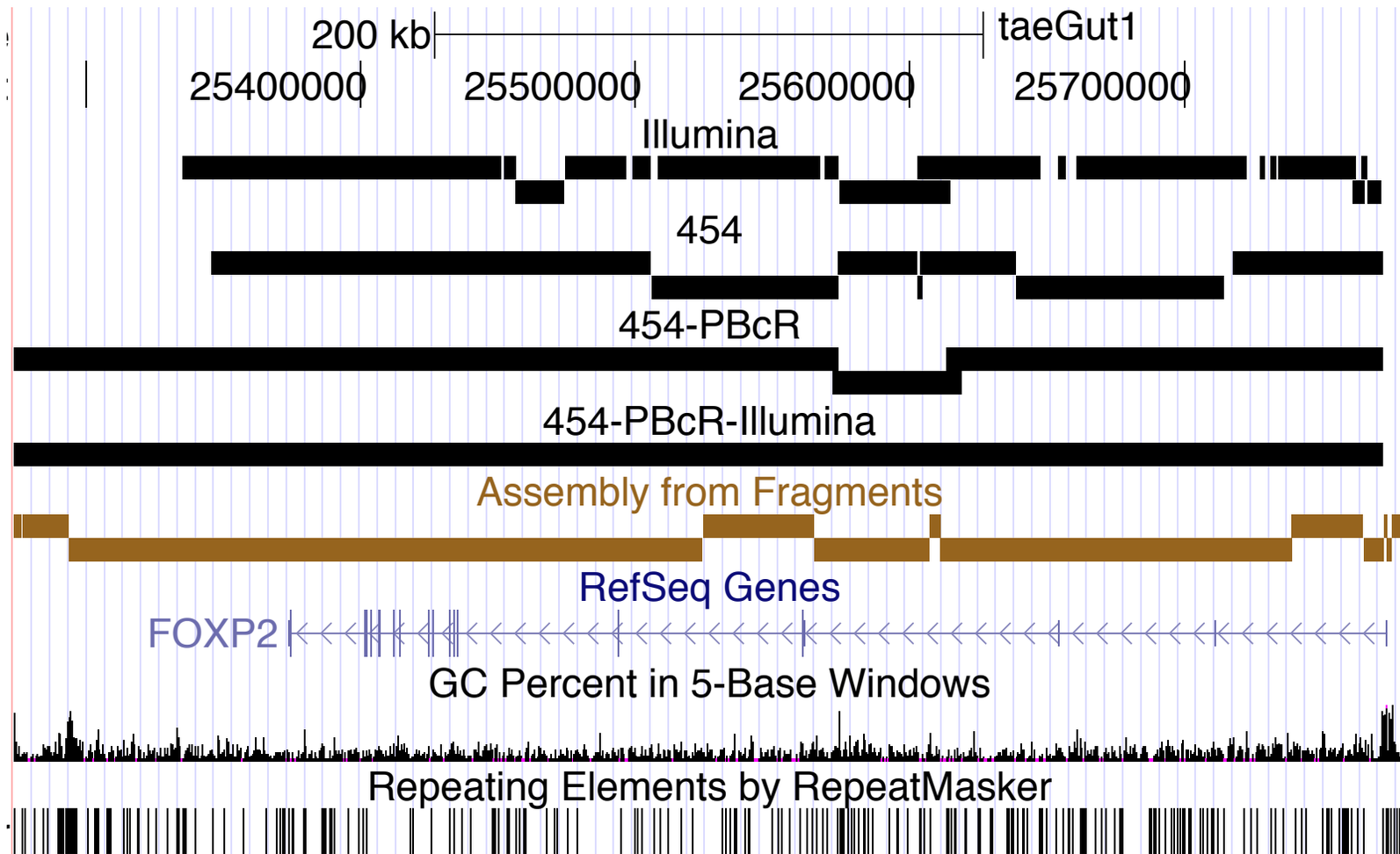
SMRT-Assembly Results



Organism	Technology	Reference Size	Assembly Size	NT Coverage	Max Contig Length	NGS
<i>Saccharomyces cerevisiae</i>	Illumina 150x 200bp	10 Mb	10 Mb	1	100,000 - 100,000	100,000 - 100,000 (100%)
	PacBio-PbA 10x		10 Mb	1	100,000 - 100,000	100,000 - 100,000 (100%)
<i>E. coli</i>	Illumina 150x 200bp	4.6 Mb	4.6 Mb	60	100,000 - 100,000	100,000 - 100,000 (100%)
	PacBio-PbA 10x		4.6 Mb	10	100,000 - 100,000	100,000 - 100,000 (100%)
	Both PacBio-PbA + Illumina 150x 200bp		4.6 Mb	60	100,000 - 100,000	100,000 - 100,000 (100%)
<i>E. coli</i> (2011)	PacBio-PbA 10x	4.6 Mb	4.6 Mb	10	100,000	100,000
	PacBio 20x PbA corrected by 20x CCS		4.6 Mb	60	100,000	100,000
	Both PacBio-PbA 20x + CCS 20x		4.6 Mb	60	100,000	100,000
	PacBio 50x PbA corrected by 50x CCS		4.6 Mb	60	100,000	100,000
	Both PacBio-PbA 50x + CCS 50x		4.6 Mb	60	100,000	100,000
	Manually corrected HiFi/CCS assembly		4.6 Mb	60	100,000	100,000
<i>S. cerevisiae</i> 1016a	Illumina 150x 200bp	12.1 Mb	12.1 Mb	100	100,000 - 100,000	100,000 - 100,000 (100%)
	PacBio-PbA 10x		12.1 Mb	10	100,000 - 100,000	100,000 - 100,000 (100%)
	Both PacBio-PbA 10x + Illumina 150x 200bp		12.1 Mb	100	100,000 - 100,000	100,000 - 100,000 (100%)
<i>Mycobacterium tuberculosis</i>	Illumina 150x 200bp paired end (150,000 read pairs)	4.2 Mb	4.2 Mb	24.5x	1,000,000	1,000,000
	40x 150x (PbA + HiFi PbA + SMRTLink paired-end)		4.2 Mb	14.5x	1,000,000	1,000,000
	40x 150x + PacBio-PbA 10x		4.2 Mb	14.5x	1,000,000	1,000,000

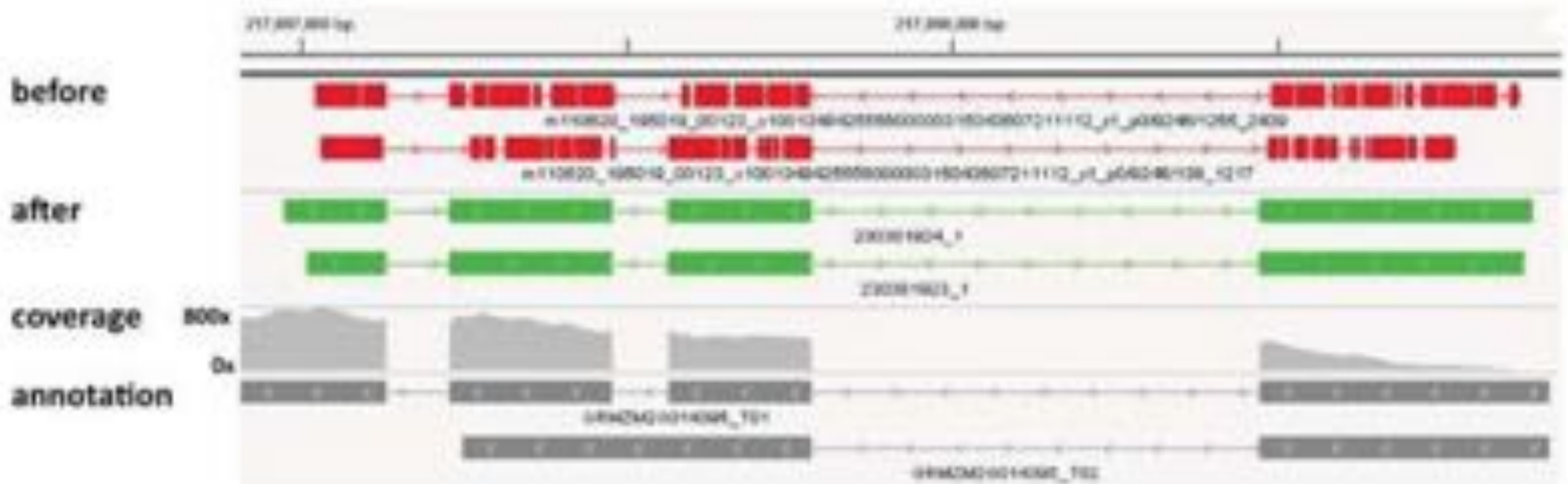
Hybrid assembly results using error corrected PacBio reads
Meets or beats Illumina-only or 454-only assembly in every case

Improved Gene Reconstruction



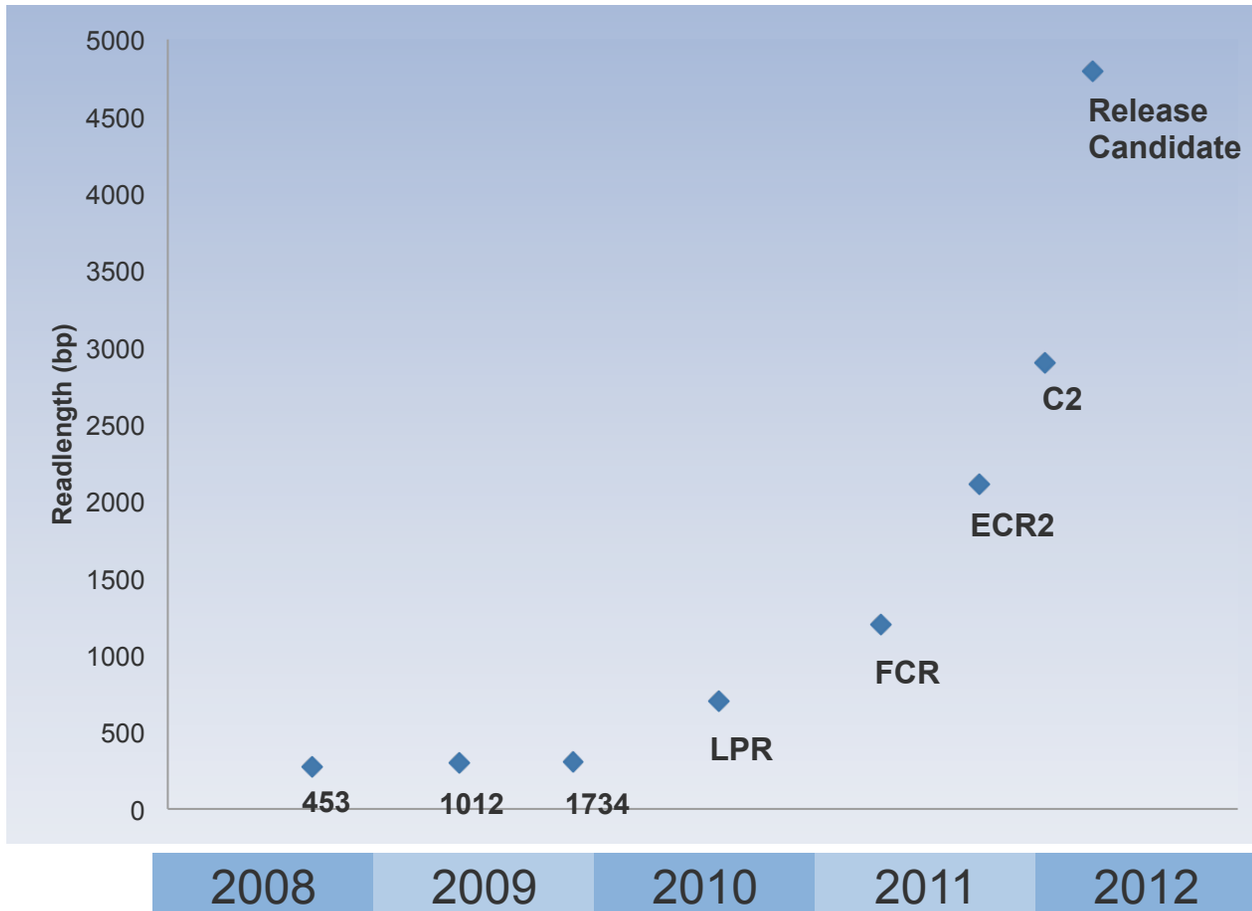
FOXP2 assembled on a single contig

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

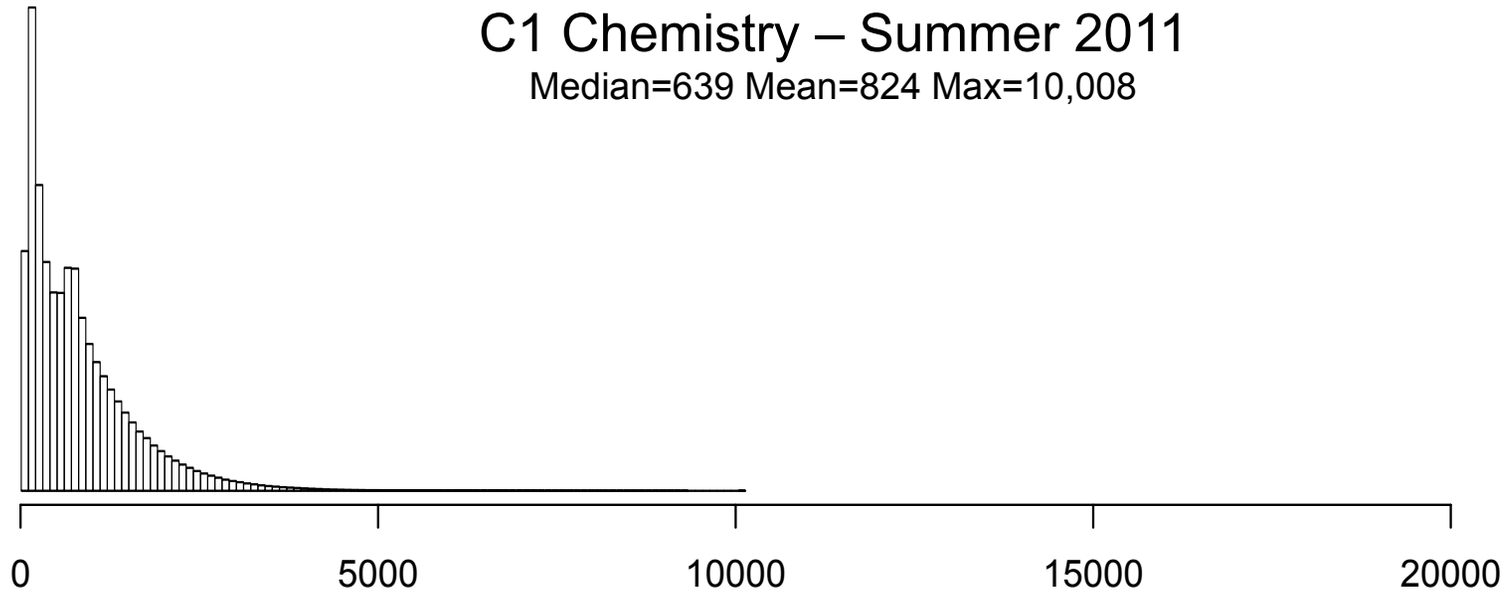
Very recent improvements:

1. Improved enzyme:
Maintains reactions longer
2. “Hot Start” technology:
Maximize subreads
3. MagBead loading:
Load longest fragments

PacBio Rice Sequencing

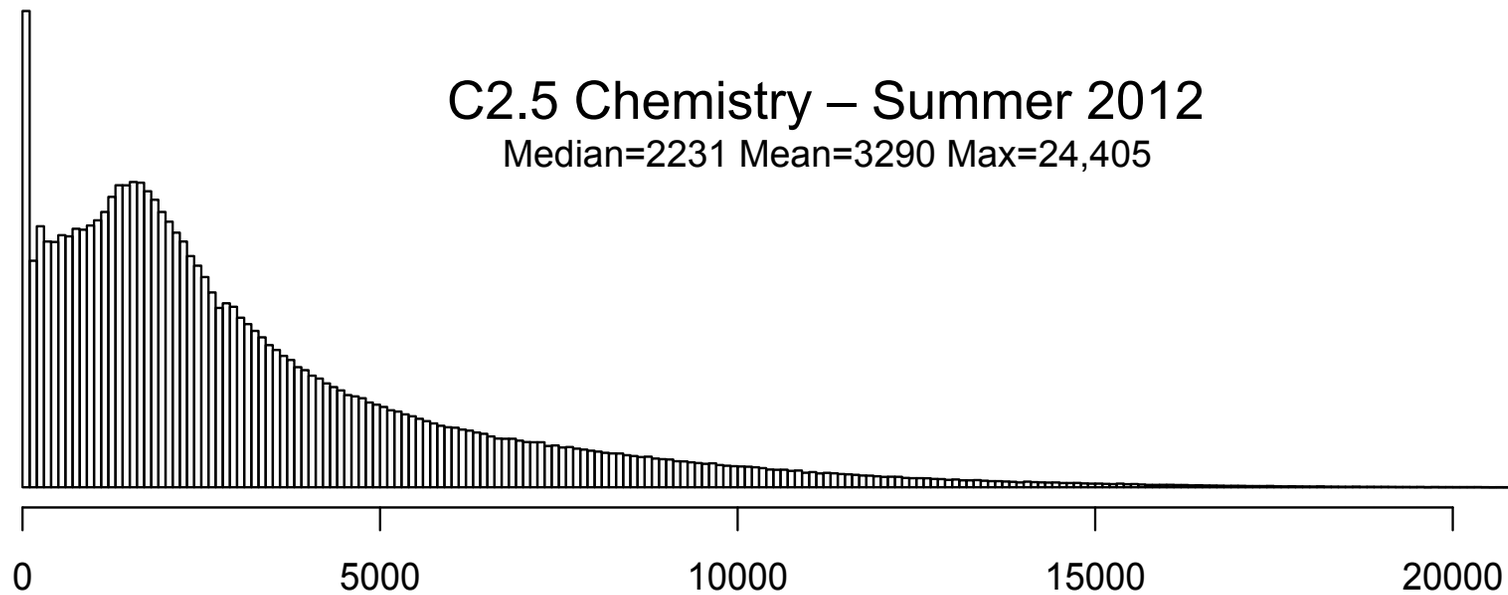
C1 Chemistry – Summer 2011

Median=639 Mean=824 Max=10,008

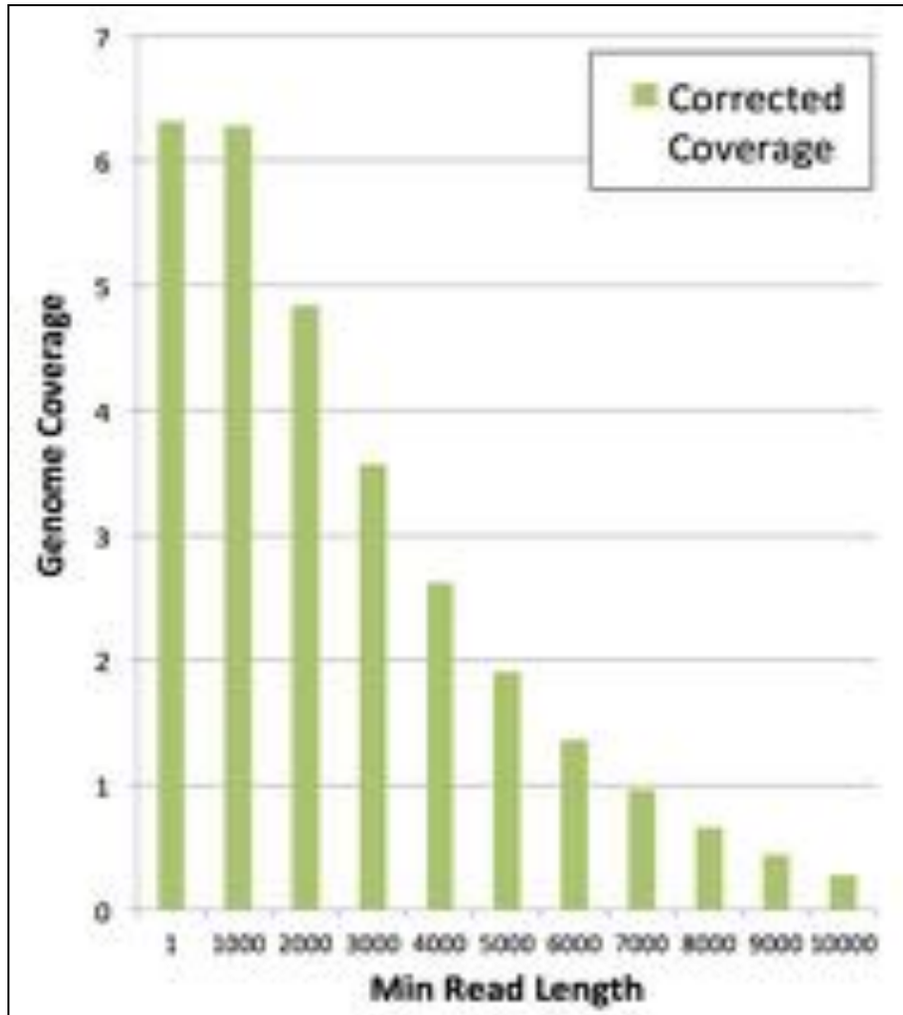


C2.5 Chemistry – Summer 2012

Median=2231 Mean=3290 Max=24,405



Preliminary Rice Assemblies



Assembly	Contig N50
Illumina Fragments 50x 2x100bp @ 180	3925
Illumina Mates 50x 2x100bp @ 180 36x 2x50bp @ 2100 51x 2x50bp @ 4800	13696
MiSeq Fragments 23x 459bp 8x 2x251bp @ 450	6444
PBeCR Reads 6.3x 2146bp ** MiSeq for correction	13600
PBeCR + Mates 6.3x 2146bp ** MiSeq for correction 51x 2x50bp @ 4800	In Progress

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



Acknowledgements

Schatz Lab

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Varrun Ramani
Eric Biggers

CSHL

Hannon Lab
Iossifov Lab
Levy Lab
Lippman Lab
Lyon Lab
Martienssen Lab
McCombie Lab
Ware Lab
Wigler Lab

NBACC

Adam Phillippy
Sergey Koren

JHU/UMD

Steven Salzberg
Mihai Pop
Ben Langmead
Cole Trapnell





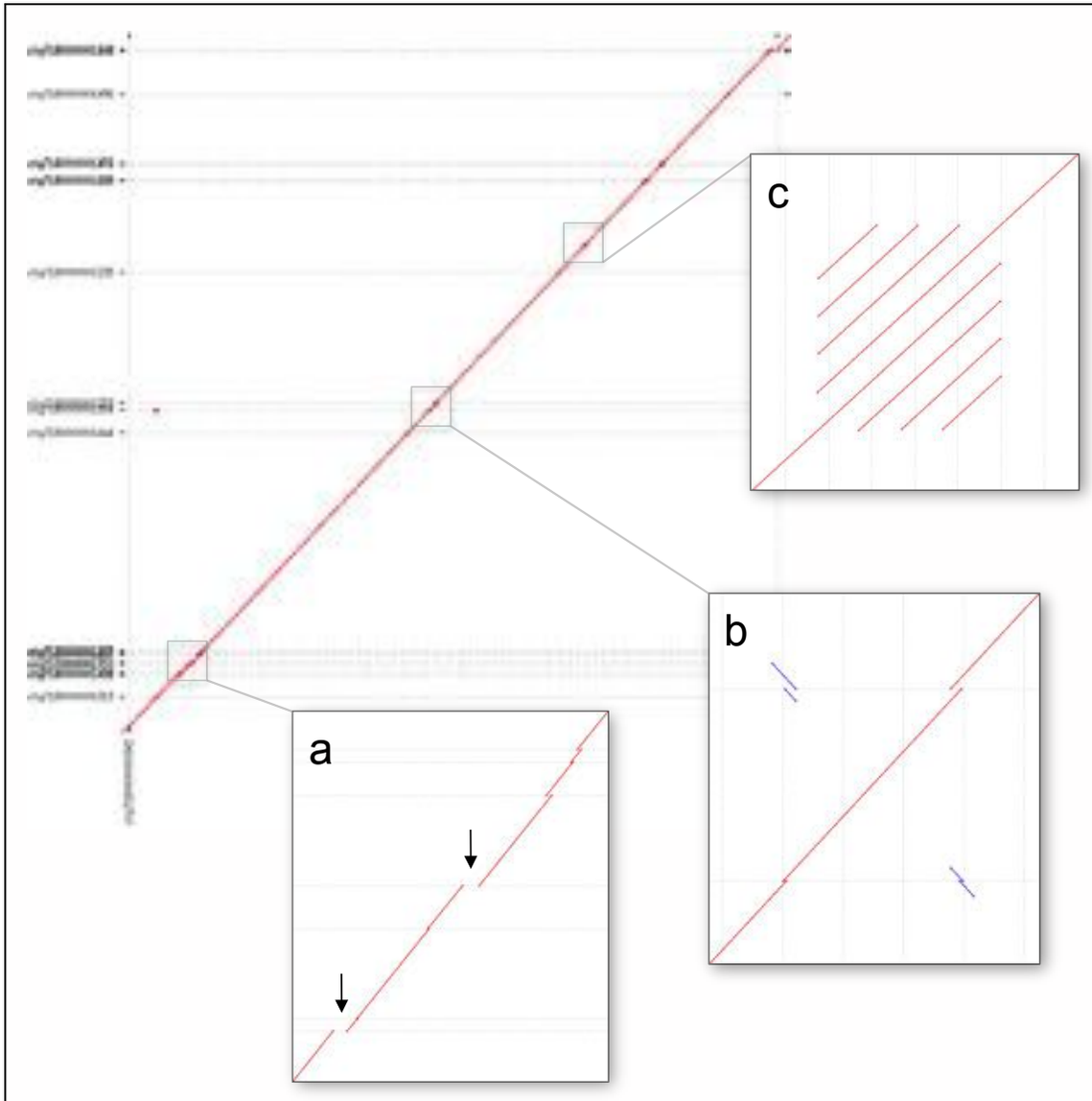
Thank You!

Want to push the frontier of bioinformatics, biotechnology, & genetics?

<http://schatzlab.cshl.edu/apply/>



Long Read Advantages

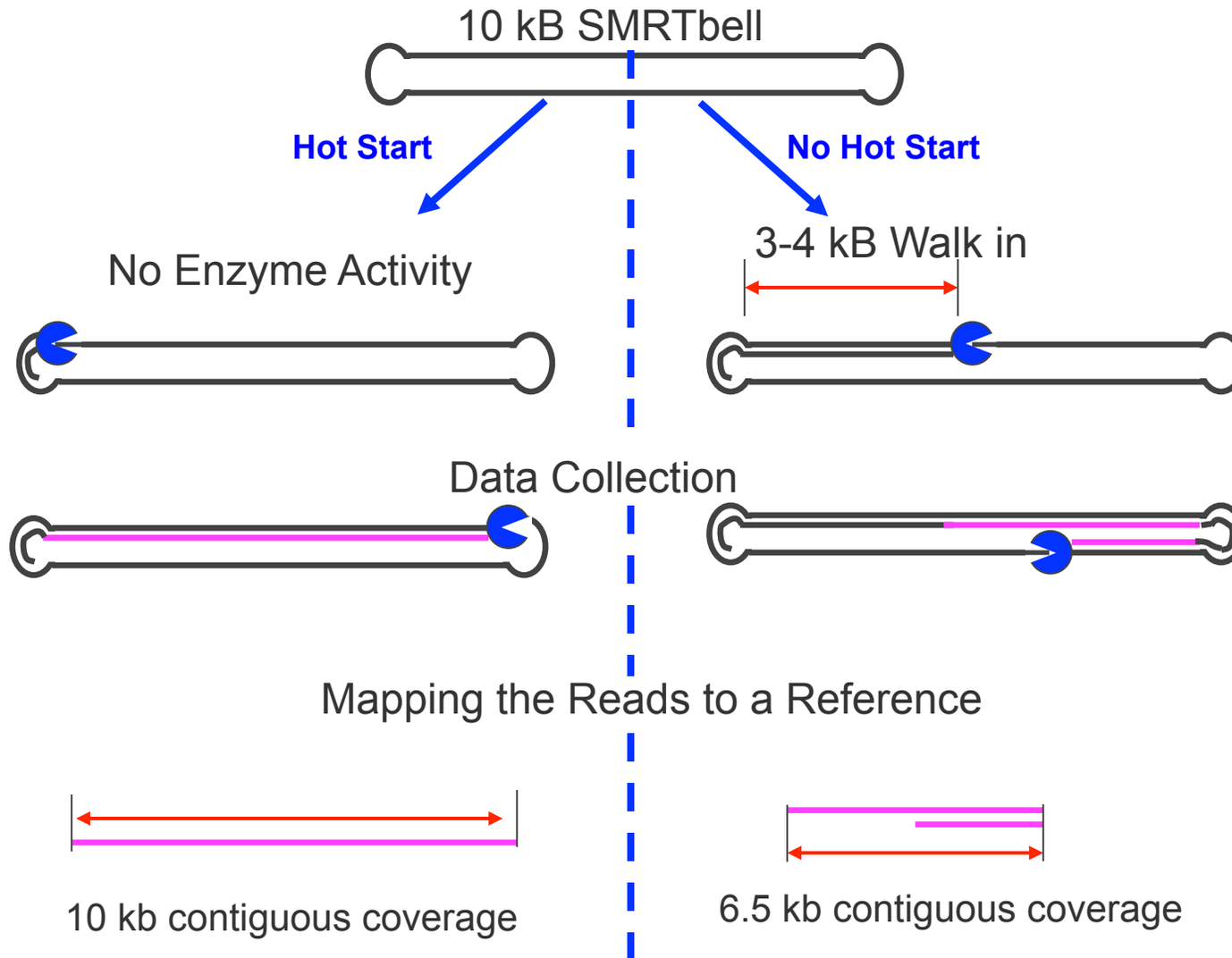


(a) Long reads close sequencing gaps

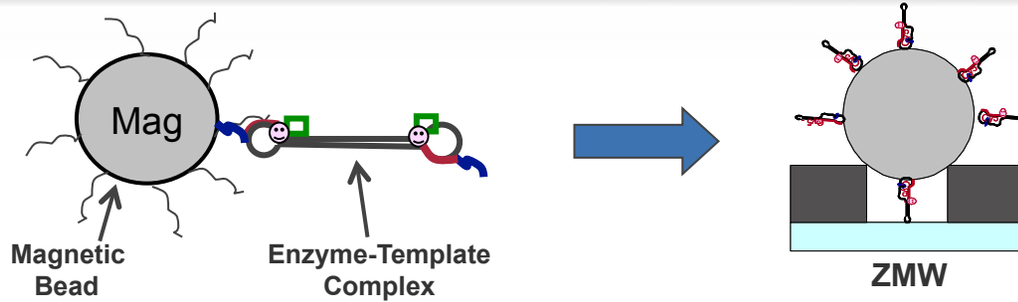
(b) Long reads assemble across long repeats

(c) Long reads span complex microsatellites

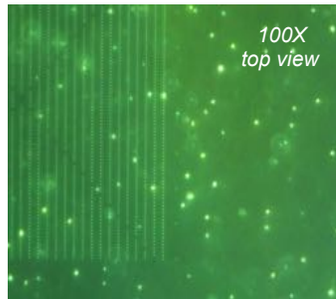
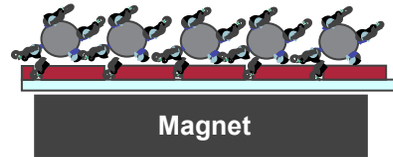
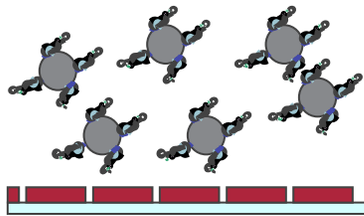
Theoretical Benefits of Hot Start Sequencing



Magnetic Bead Enzyme-Template Complex Loading



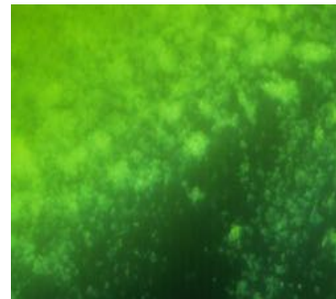
Multiple complexes attached to magnetic beads that are much larger than individual ZMWs



100X
top view

(I)

Pre-Deposition: Complex loaded beads in solution



(II)

Introduce magnet: Bead complexes pulled to chip surface



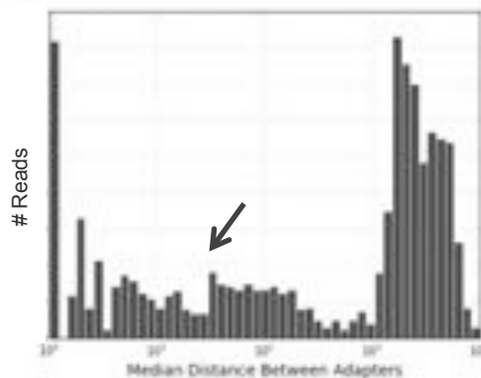
Rotate magnet to evenly disperse beads across entire chip surface

MBS (MagBead Station)

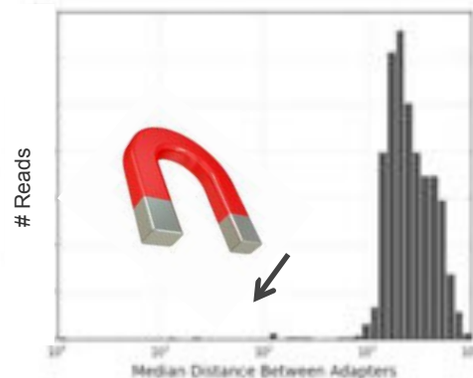
Listeria



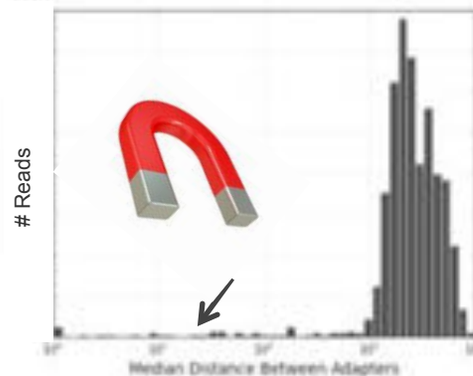
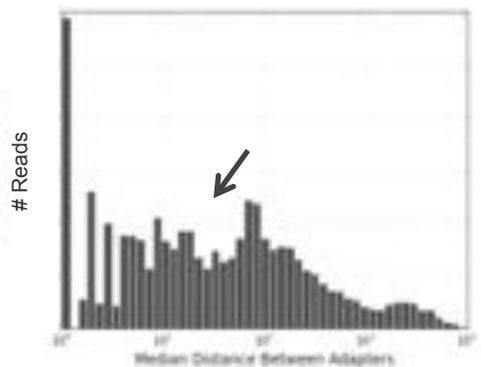
Diffusion



MBS



Cholera



Improvements to Sample Prep