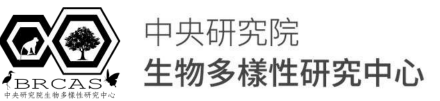
Genome assemblies

Isheng Jason Tsai





Lecture outline

- Introduction
- Assembly algorithm overview
- Long read technologies
- Scaffolding
 - Chromosome conformation capture
- Case studies

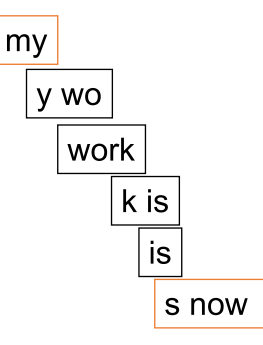
Problem

We accidently printed my five copies of "Origin of Species", and shredded into pieces



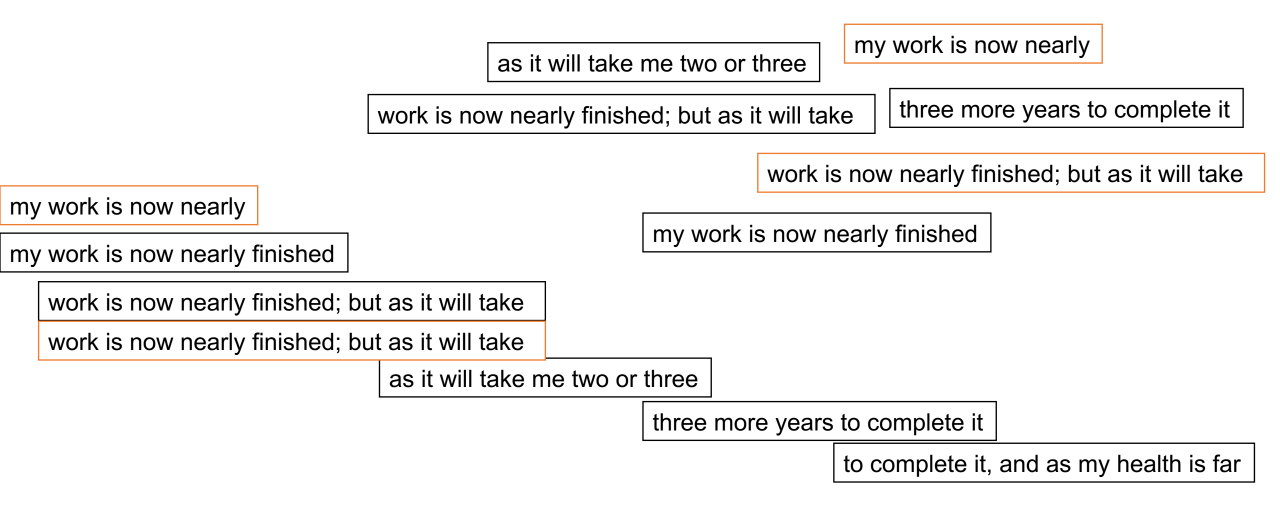
my work is now nearly finished; but as it will take me two or three more years to complete it, and as my health is far from strong, I have been urged to publish this Abstract. I have more especially been induced to do this, as Mr Wallace, who is now studying the natural history of the Malay archipelago, has arrived at almost exactly the same general conclusions that I have on the origin of species. Last year he sent to me a memoir on this subject, with a request that I would forward it to Sir Charles Lyell, who sent it to the Linnean Society, and it is published in the third volume of the journal of that Society. Sir C. Lyell and Dr Hooker, who both knew of my work -- the latter having read my sketch of 1844 -honoured me by thinking it advisable to publish, with Mr Wallace's excellent memoir, some brief extracts from my manuscripts.

Assemby = Piecing the pieces together



my work is now

Long shredded pieces (read) = easier assembly



my work is now nearly finished; but as it will take me two or three more years to complete it, and as my health is far

Why sequence a genome?

Genomics advance our understanding of organisms nature nature

2008

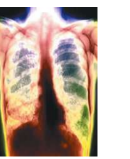
2007

2009

nature

genetics

across tree of life pathogens



TB

1996

997



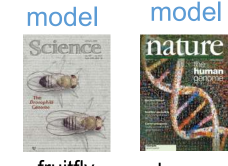
1998

666

2000

nature

Plasmodium genomic:





2002

nature

2001



2005



2010

2011

2012

2013









S. cerevisiae model





2006





nature THE TOMATO GENOME BLIGHTED

Blood fluke

2014





Phytophthora infestans Black death Smut fungi (potato blight) pathogens

Tapeworms

Assemble a genome is hard



Less complicated

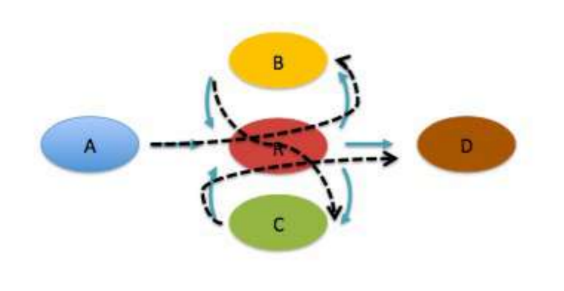


Complicated

Repeats

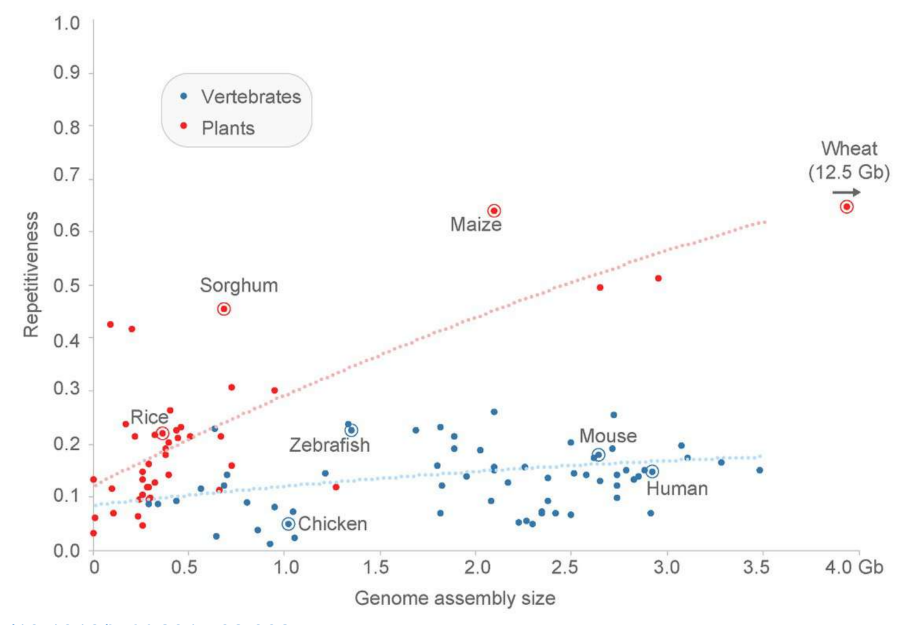
You can't 100% resolve repeats unless you have your sequence length > repeat





Credit: Michael Schatz

Repeat content of plant and vertebrate genomes



https://doi.org/10.1016/j.pbi.2017.02.002

Largest genome ever recorded: 149Gb



Organism Type	Organism Name	Approximate Genome size, in number of nucleotides ("letters")	Number of protein-coding genes
Bacterium	Nasuia deltocephalinicola, a tiny bacterium that lives inside an insect [3]	112,000 (0.112 million) * currently the smallest known bacterial genome	137
Bacterium	Escherichia coli [2]	4,600,000 (4.6 million)	5,000
Plant	Arabidopsis thaliana	135,000,000 (135 million)	27,416
Mammal	Homo sapiens, Humans	3,000,000,000 (3 billion)	20,000 [5]
Plant	Norway Spruce	19,000,000,000 (19 billion)	28,000
Plant	Paris japonica, a rare Japanese flower [4]	149,000,000,000 (149 billion) * currently the largest known genome	unknown

http://thatslifesci.com/2019-05-06-Biggest-genome-of-them-all-JBarnett/

Actually not too far off...

Article Giant lungfish genome elucidates the conquest of land by vertebrates

k-mers (Extended Data Fig. 2). We ascertained the high completeness of the 37-Gb assembly by observing that 88.2% of the DNA and 84% of the RNA sequencing (RNA-seq) reads aligned to the genome, which gives an estimated total genome size of 43 Gb (about 30% larger than the axolotl⁸). This matches the *k*-mer value but is smaller than that predicted by flow cytometry (52 Gb⁹) and Feulgen photometry (75 Gb¹⁰).

QC and understand your data before assembly

Conversation:

Jason, we don't know why our assembly is not as good as yours. We have about 90X of Mate pair data..

Email 2 weeks later:

Dear Jason,

I run the Trimmomatic analysis for the raw data of mate-pair libraries (10kL2_1, 10kL7_1, 15kL3_1 and 15kL8_4 as examples) with a custom adapter file containing mate-pair adapter sequences (junction and external adaptors, you may find them in the attached technote pdf file) and found **that over 80% reads of the library were dropped out.**

Always be careful of contamination

PNAS

Evidence for extensive horizontal gene transfer from the draft genome of a tardigrade

Thomas C. Boothby^{a,1}, Jennifer R. Tenlen^{a,2}, Frank W. Smith^a, Jeremy R. Wang^{a,b}, Kiera A. Patanella^a, Erin Osborne Nishimura^a, Sophia C. Tintori^a, Qing Li^c, Corbin D. Jones^a, Mark Yandell^c, David N. Messina^d, Jarret Glasscock^d, and Bob Goldstein^a

^aDepartment of Biology, University of North Carolina at Chapel Hill, Chapel Hill, NC 27599; ^bDepartment of Genetics, University of North Carolina at Chapel Hill, Chapel Hill, Chapel Hill, NC 27599; ^cEccles Institute of Human Genetics, University of Utah, Salt Lake City, UT 84112; and ^dCofactor Genomics, St. Louis, MO 63110

Edited by W. Ford Doolittle, Dalhousie University, Halifax, Canada, and approved September 28, 2015 (received for review May 28, 2015)

No evidence for extensive horizontal gene transfer in the genome of the tardigrade *Hypsibius dujardini*

Georgios Koutsovoulos^a, Sujai Kumar^a, Dominik R. Laetsch^{a,b}, Lewis Stevens^a, Jennifer Daub^a, Claire Conlon^a, Habib Maroon^a, Fran Thomas^a, Aziz A. Aboobaker^c, and Mark Blaxter^{a,1}

^aInstitute of Evolutionary Biology, University of Edinburgh, Edinburgh EH9 3FL, United Kingdom; ^bThe James Hutton Institute, Dundee DD2 5DA, United Kingdom; and ^cDepartment of Zoology, University of Oxford, Oxford OX1 3PS, United Kingdom

Edited by W. Ford Doolittle, Dalhousie University, Halifax, Canada, and approved March 1, 2016 (received for review January 8, 2016)

17.5 % HGT

0.4 % HGT

Always be careful of contamination

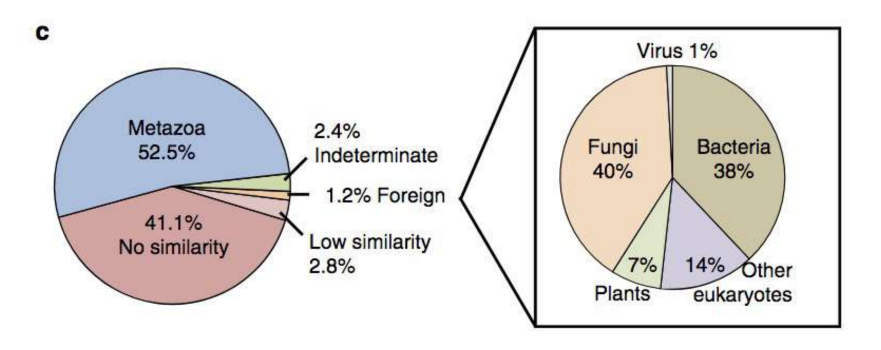
ARTICLE

Received 21 Jun 2015 | Accepted 3 Aug 2016 | Published 20 Sep 2016

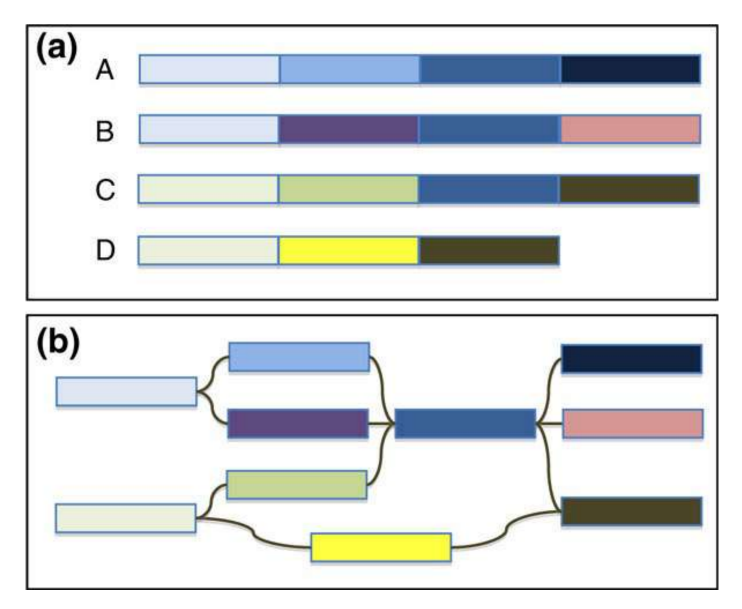
DOI: 10.1038/ncomms12808

OPEN

Extremotolerant tardigrade genome and improved radiotolerance of human cultured cells by tardigrade-unique protein



Ploidy, heterozygosity and the assembly graph



http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3446297/

Assembly process

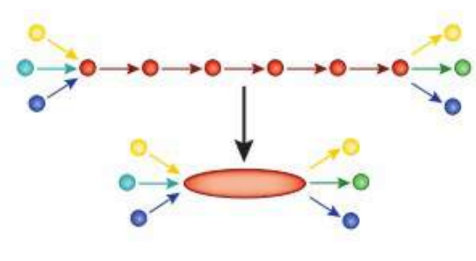
1. Fragment DNA and sequence



CGCATATCCGGT

2. Find overlaps between reads

3. Assemble overlaps into contigs



4. Assemble **contigs** into **scaffolds**

AGCCTAGACCTACA



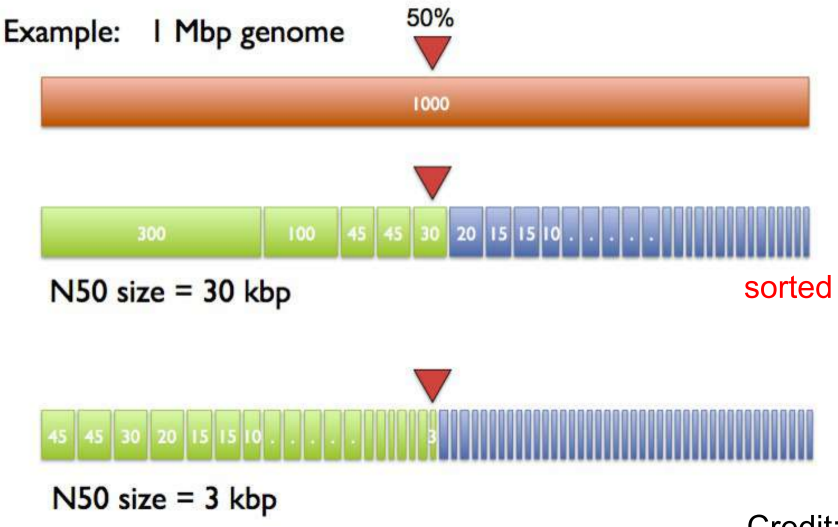
http://www.nature.com/nmeth/journal/v9/n4/pdf/nmeth.1935.pdf

Which program to choose?

SSPACE HaploMerger EULER-SR WhatsHap GARM SOAPdenovo mip GABenchToB Telescoper RAMPART fermi Contrail A5 QSRA Opera Arapan AutoAssemblyD Platanus SWAP-Assembler Newbler PCAP HapCompass Dazzler SSAKE Forge SHEAR Mapsembler 2 ALLPATHS-LG CLC VICUNA Edena TIGR PERGA KmerGenie CloudBrush Cortex REAPR TIGRA Amos gapfiller Ray Tedna MIRA dipSPAdes ATAC MetAMOS Nesoni Arachne Geneious SeqMan NGen Celera GAM Quast VCAKE PASHA MetaVelvet-SL SCARPA MHAP Hapsembler bambus2 **IDBA** GAML Sequencher BESST GGAKE Trinity GRIT MaSuRCA HiPGA PADENA Phrap SeqPrep Phusion PE-Assembler SGA KGBAssembler Metassembler Curtain SWiPS CGAL HGAP PRICE Pilon MSR-CA Taipan Pipeline Pilot SHRAP SILP3 IDBA-MTP Omega SUTTA ABySS HyDA-Vista SR-ASM OMACC Anchor Velvet Enly DNAnexus SOPRA Atlas Ragout SPAdes **iMetAMOS** FRCBam SAGE Monument Cerulean SAT-Assembler DNA Dragon CABOG SHORTY SHARCGS GAGM image ngsShoRT ABBA FALCON SuccinctAssembly DecGPU Lasergene PBJelly GenoMiner Khmer **ELOPER** GiaAssembler

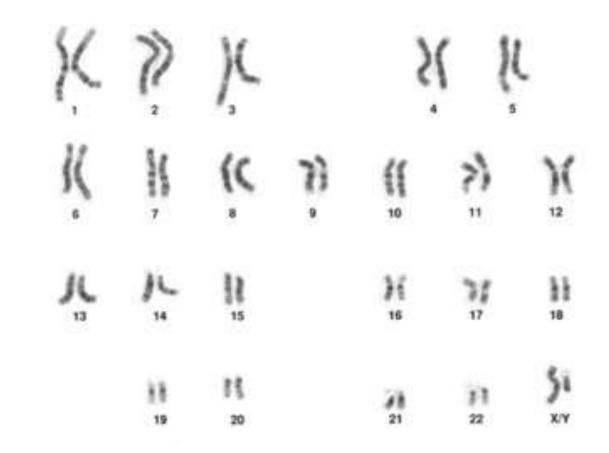
Contiguity (good) is a genome? **N50**

Definition: 50% of genome in contigs/scaffolds of length N50 bp or greater



Credit: Michael Schatz

Most assemblies are fragmented

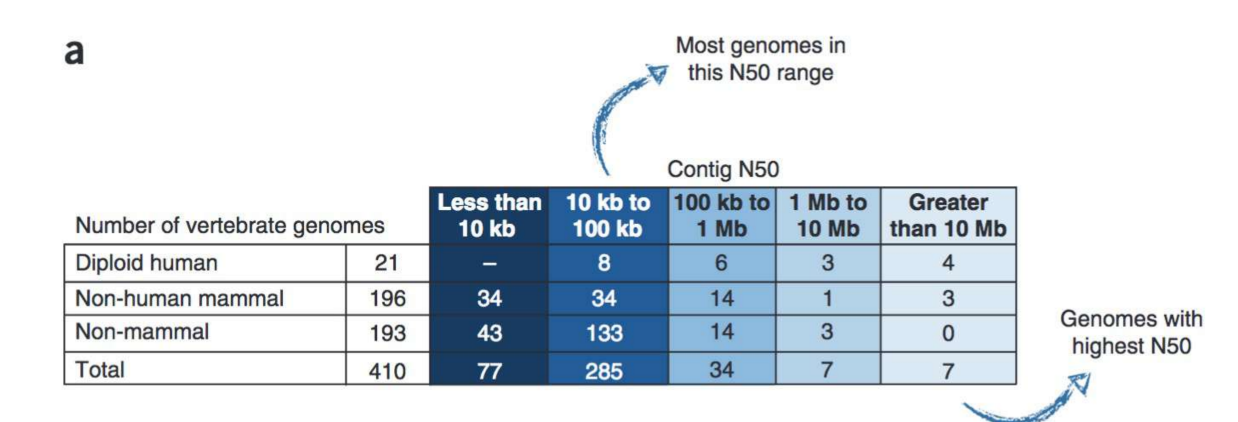


chromosomes



Contigs

Statistics for current GenBank assemblies.



doi:10.1038/ng.3824

Why do we need a good assembly?

It is easier to analyse 10000 pieces vs. 23 pieces (chromosomes)

Allows more accurate representation of genes locations on genome Is gene A close to gene B? On the same chromosome?

Transposon dynamics

Missing in most assemblies (located in NNNNNNNN gaps)

Responsibility to contribute to your community Do you want others to work on the same genome / species as well?

Bottom line:

It's really no point to do one if you can't produce an accurate and useful assembly

Assembly qualities

	Whole Genome Representation	Sequence Status	Genes	Usability	
1	Incomplete for non- repetitive regions	Small scaffolds and contigs	Incomplete genes	Markers development	
2	Complete for non- repetitive regions	Medium scaffolds and contigs	Complete but 1-2 genes/contig	Gene mining	
3	Complete for non- repetitive regions	Large scaffolds and contigs	Several dozens of genes/contig	Microsynteny	
4	Complete for almost the whole genome	Pseudomolecules	Hundreds of genes/ contig	Any (Synteny, Candidate gene by QTLs)	
5	Complete genome	Pseudomolecules	Thousands of genes/ contig	Candidate gene by QTES	

Credit: Aureliano Bombarely

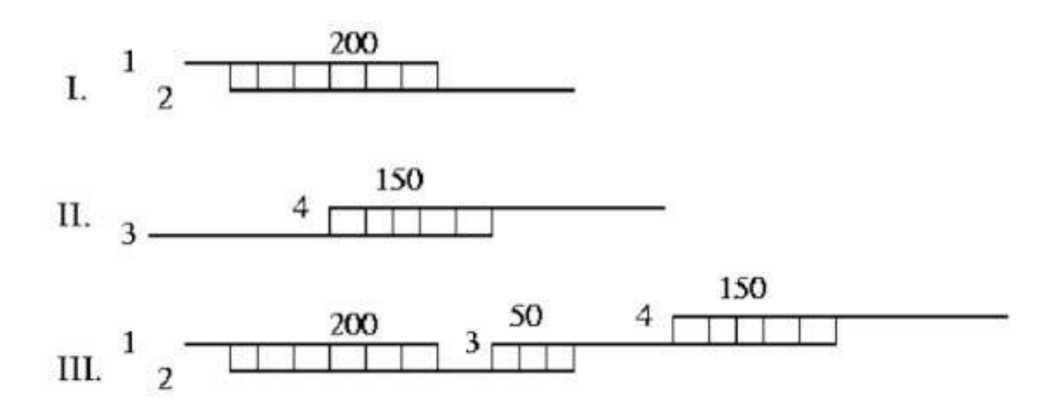
Assembly algorithms

Approaches

- Greedy extension
 - only mentioned for historical reasons
- Overlap Layout Consensus (OLC) assembly: 'traditional' and well established method, but challenging to implement at each stage
 - Most "old" and "newest" assemblies were produced using this approach
- de Bruijn graph (DBG) assembly

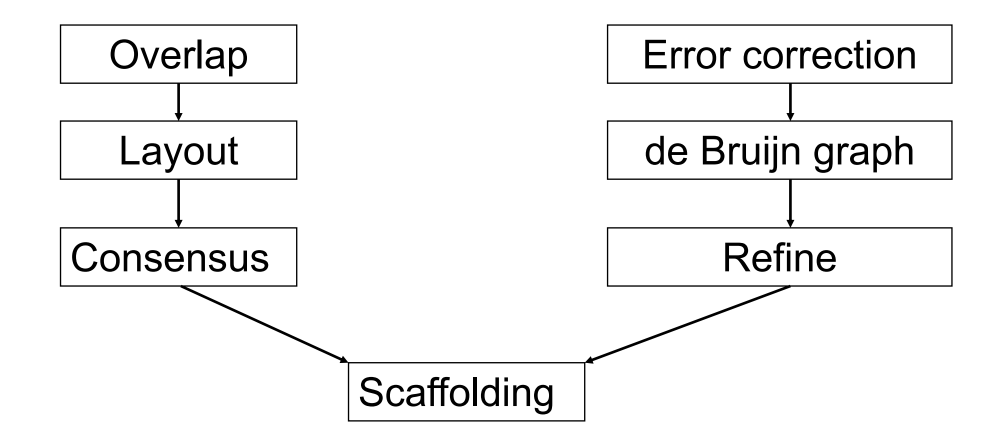
Greedy extension

Oldest and not really useful in most cases



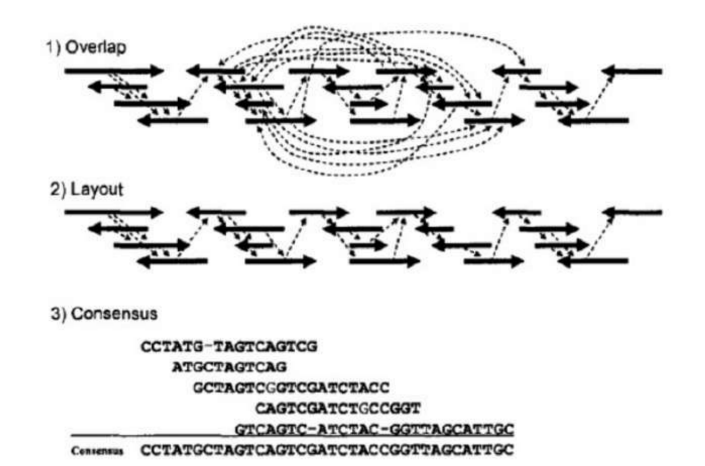
http://www.cbcb.umd.edu/research/assembly_primer.shtml

OLC and DBG assemblers



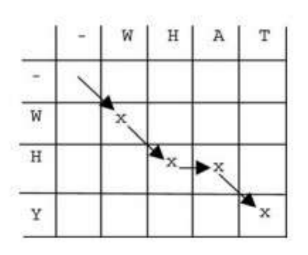
OLC approach

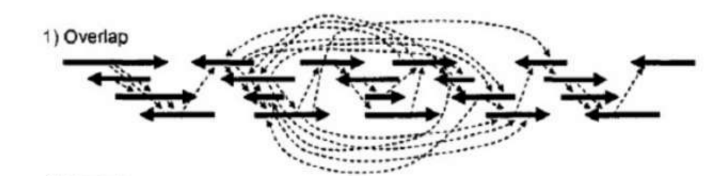
- Pairwise alignment of all reads to find **overlaps**
- Layout the reads to decide which read align to which
- Get consensus by joining join the read sequences, merging overlaps
- All three are challenging



Overlap

- All vs. all pairwise alignment
 - Smith-Waterman? Blast? Kmer based? Suffix tree? Dynamic programming?
- Computationally very intensive but can be parallelised
 - Need lots of CPUs!
 - Batch1 align Batch 1 in 1st CPU
 - Batch1 align Batch 2 in 2nd CPU





Computational Example 8.1: Pseudocode for overlap alignment

Input sequences A, B Set $O_{i,0} = O_{0,j} = 0$ for all i, jfor i = 1 to nfor j = 1 to m $O_{i,j} = \max\{O_{i-1,j} - \delta, O_{i-1,j-1} + s(a_i, b_j), O_{i,j-1} - \delta\}$ end end Best overlap = $\max\{O_{i,m}, O_{n,j}; 1 \le i \le n, 1 \le j \le m\}$

Deonier et al., Computational Genome Analysis

Build overlap graph

- It's common practice to represent them in graphs
- The actual overlaps are the edges
- Now we create the genome assembly graph

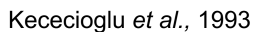
my work is now nearly finished

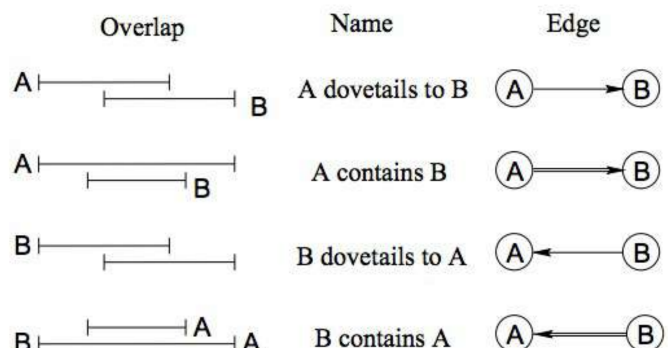
B

my work is now nearly finished

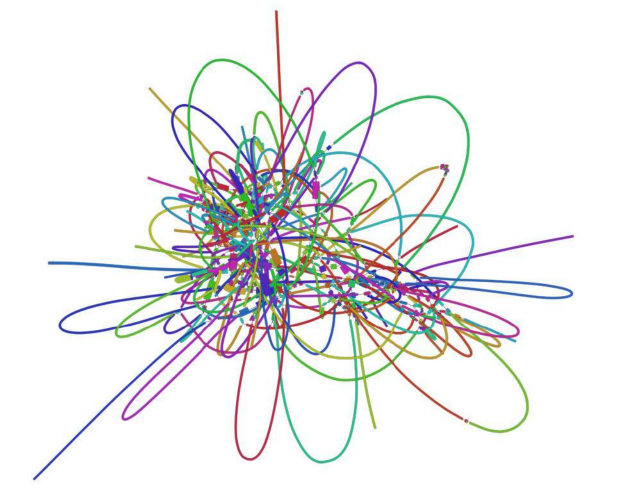
work is now nearly finished; but as it will take

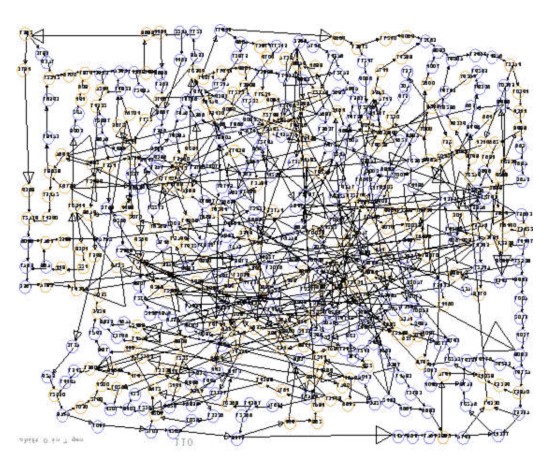
work is now nearly finished; but as it will take





Some assembly graph can be complicated...

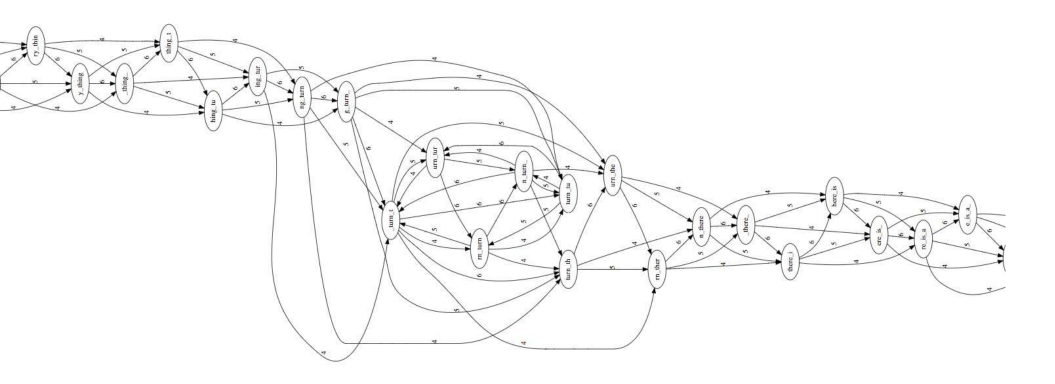




http://rrwick.github.io/Bandage/images/screenshots/screenshot02.png

Overlap graph is big and messy. Contigs don't "pop out" at us.

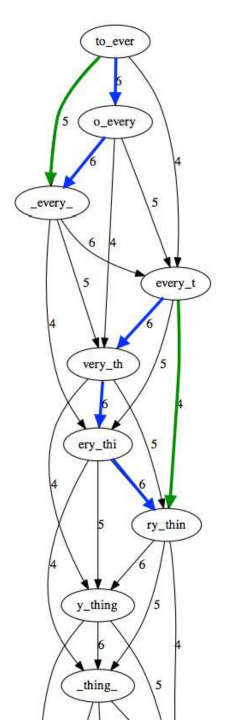
Below: part of the overlap graph for to_every_thing_turn_turn_there_is_a_season l = 4, k = 7



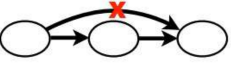
Order the reads into a consistent manner

Anything redundant about this part of overlap graph?

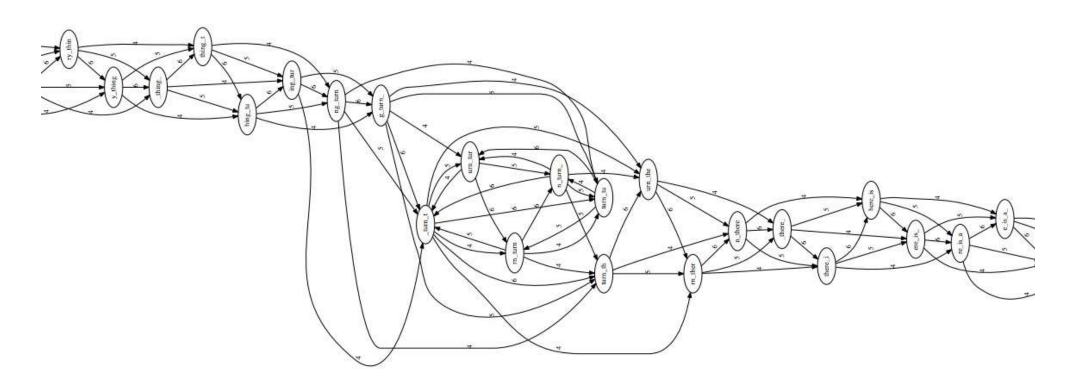
Some edges can be inferred from other edges E.g., green edge can be inferred from blue



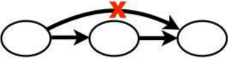
Remove transitively-inferrible edges, starting with edges that skip one node:



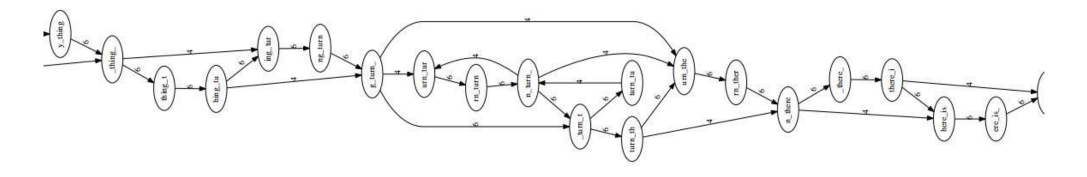
Before:



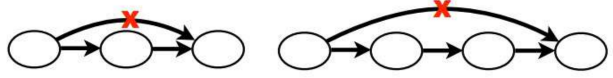
Remove transitively-inferrible edges, starting with edges that skip one node:



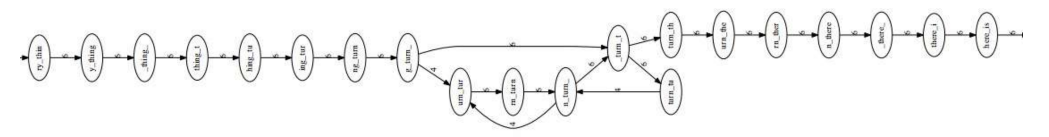
After:



Remove transitively-inferrible edges, starting with edges that skip one or two nodes:

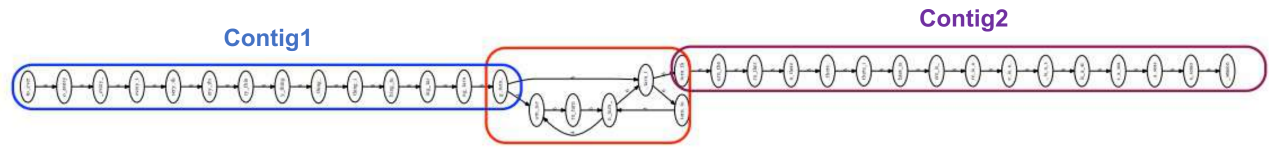


After:



Even simpler

Layout

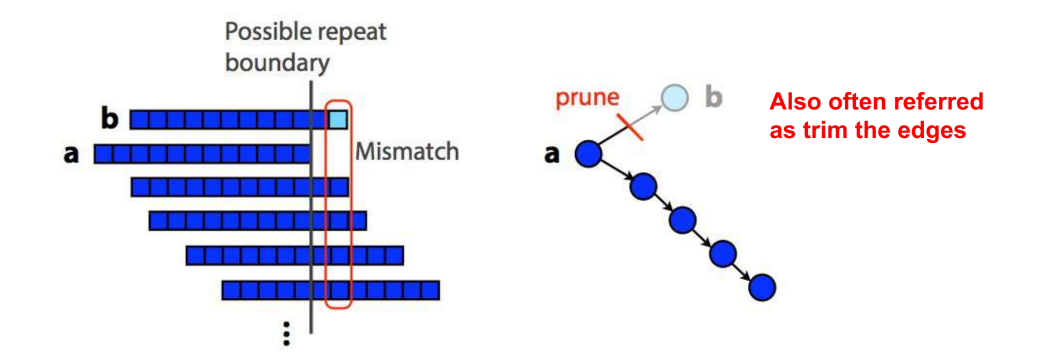


Unresolvable repeat

Depending on assemblers, they may result in 1 or 2 contigs

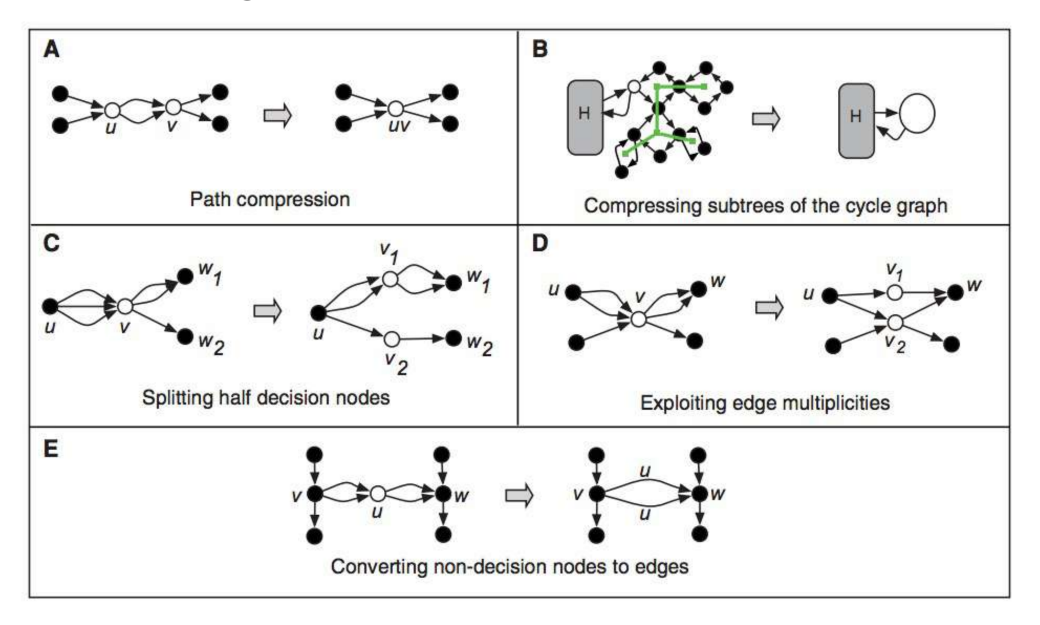
Layout – can we do more?

In practice, layout step also has to deal with spurious subgraphs, e.g. because of sequencing error



Mismatch could be due to sequencing error or repeat. Since the path through **b** ends abruptly we might conclude it's an error and prune **b**.

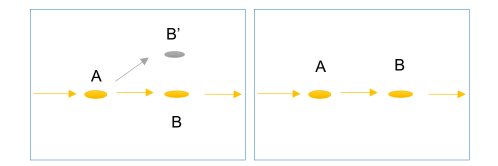
Usefulness of graph transformation



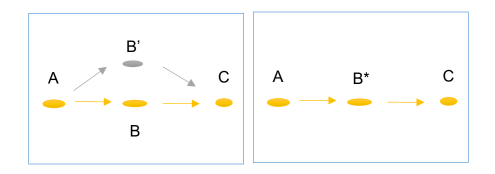
Kingsford *et al.,* (2010)

Error correction in graph

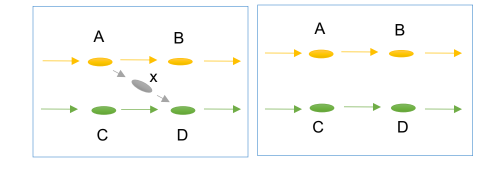
- Errors at end of read
 - Trim off 'dead-end' tips



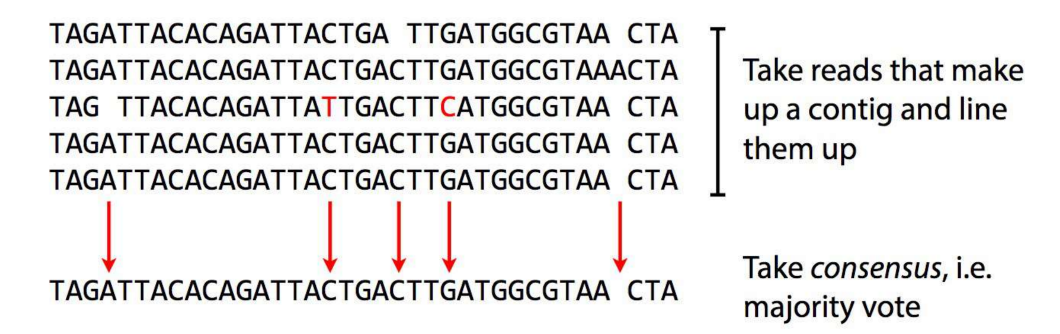
- Errors in middle of read
 - Pop Bubbles



- Chimeric Edges
 - Clip short, low coverage nodes



Consensus



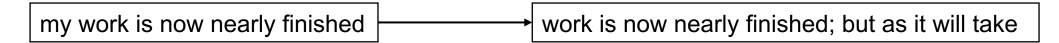
At each position, ask: what nucleotide (and/or gap) is here?

Complications: (a) sequencing error, (b) ploidy

Say the true genotype is AG, but we have a high sequencing error rate and only about 6 reads covering the position.

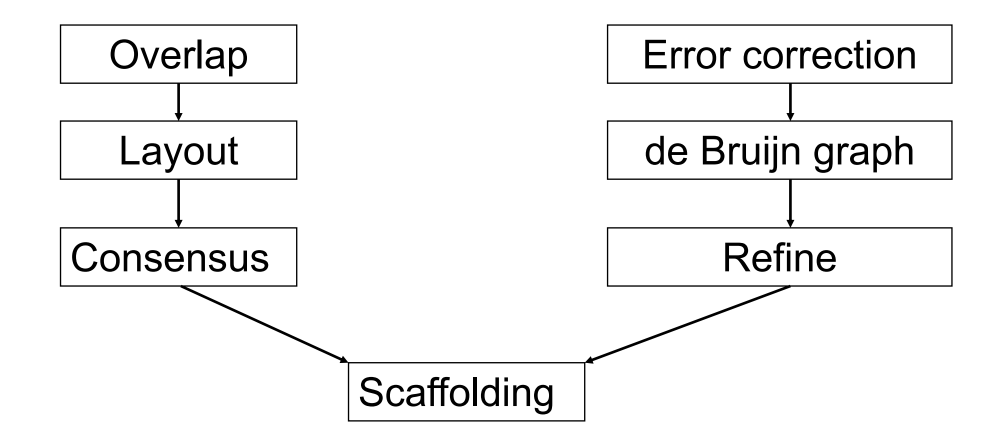
OLC Assemblers

- Mostly used in the Sanger sequencing era
 - Celera, Phusion, PCAP, Arachne
- Disadvantages of OLC
 - Computing overlaps is slow
 - 5 billion reads -> takes 400 years to compute overlaps if 1 million overlap per second
 - Overlap graph is big and complicated
 - One node per read
 - Number of edges grows superlinearly with number of reads



- When 2nd generation dataset first arrived
 - Millions and millions of reads
 - Short read length difficult to build sufficient overlap

OLC and DBG assemblers



k-mer

"k-mer" is a substring of length k

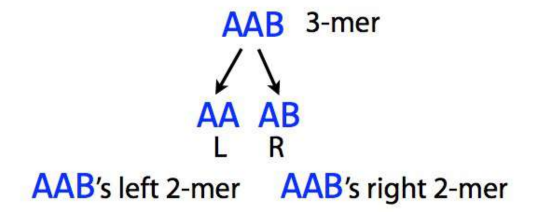
GGCGATTCATCG S: A 4-mer of S: ATTC All 3-mers of S: GGC GCG CGA GAT ATT TTC TCA CAT ATC TCG

mer: from Greek meaning "part"

We start with a collection of reads of **3bp** from the reference genome **AAABBBA**

AAA, AAB, ABB, BBB, BBA

AAB is a k-mer (k = 3). AA is its left k-1-mer, and AB is its right k-1-mer.



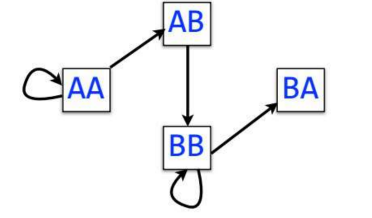
Take each length-3 input string and split it into two overlapping substrings of length 2. Call these the *left* and *right 2-mers*.

AAABBBA

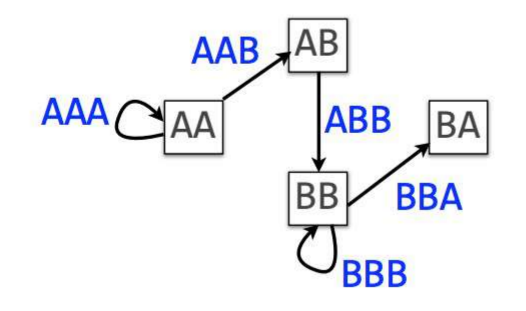
From these 2-mers, only AA, AB, BA, BB are present (they will be nodes)

Let 2-mers be nodes in a new graph. Draw a directed edge from each left 2-mer to corresponding right 2-mer:

So AAB will be $AA \rightarrow AB$



Each *edge* in this graph corresponds to a length-3 input string



How do we get contigs from the graph?

Intuitively we walk and visited all edges and node of the graph, but how?

An edge corresponds to an overlap (of length k-2) between two k-1 mers. More precisely, it corresponds to a k-mer from the input.

De Bruijn graph is a directed multigraph

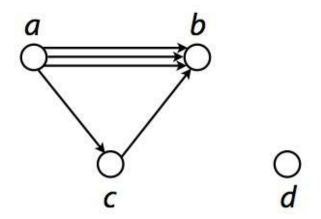
Directed **multigraph** G(V, E) consists of set of vertices, V and **multiset** of directed edges, E

Otherwise, like a directed graph

Node's *indegree* = # incoming edges

Node's *outdegree* = # outgoing edges

De Bruijn graph is a directed multigraph



$$V = \{ a, b, c, d \}$$

$$E = \{ (a, b), (a, b), (a, b), (a, c), (c, b) \}$$

$$\xrightarrow{\text{Repeated}}$$

https://en.wikipedia.org/wiki/Eulerian path

Eulerian walk definitions and statement

1.A **directed graph** is a *graph* in which each edge has a direction, usually represented as an arrow from a node *v* to a node *w*.

2.Graph is connected if each node can be reached by some other node.

3.Node is **balanced** if indegree equals outdegree. Node is **semi-balanced** if indegree differs from outdegree by 1

4.A directed, connected graph is Eulerian if and only if it has **at most 2 semi-balanced nodes** and all other nodes are balanced

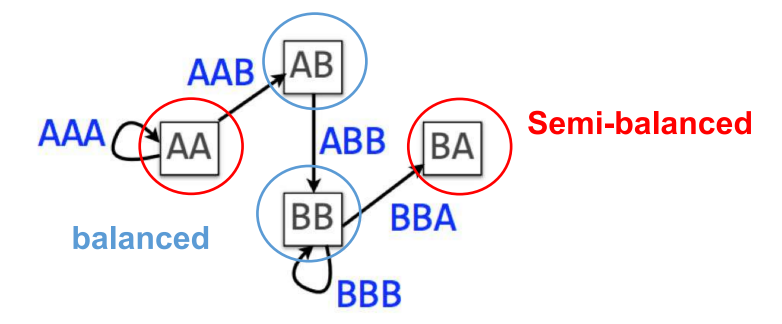
AAB AB AAA AA AB BA BB BBA OBBB

Yes

Yes

Eulerian walk definitions and statement

A directed, connected graph is Eulerian if and only if it has at most 2 semi-balanced nodes and all other nodes are balanced



Is it Eulerian? Yes

Then we can search for Eulerian path from the graph: Eulerian walk visits each edge exactly once ****

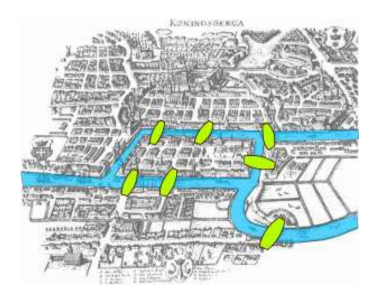
Argument 1: $AA \rightarrow AA \rightarrow AB \rightarrow BB \rightarrow BB \rightarrow BA$

Argument 2: AA and BA are semi-balanced, AB and BB are balanced

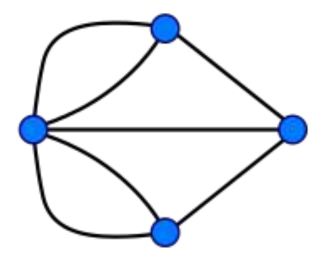
Eulerian walk

In graph theory, an Eulerian trail (or Eulerian path) is a trail in a graph which visits every edge exactly once. Similarly, an Eulerian circuit or Eulerian cycle is an Eulerian trail which starts and ends on the same vertex. They were first discussed by Leonhard Euler while solving the famous Seven Bridges of Königsberg problem in 1736. Mathematically the problem can be stated like this:

Given the graph in the image, is it possible to construct a path (or a <u>cycle</u>, i.e. a path starting and ending on the same vertex) which visits each edge exactly once?







PS. This graph is Not Eulerian

https://en.wikipedia.org/wiki/Eulerian_path

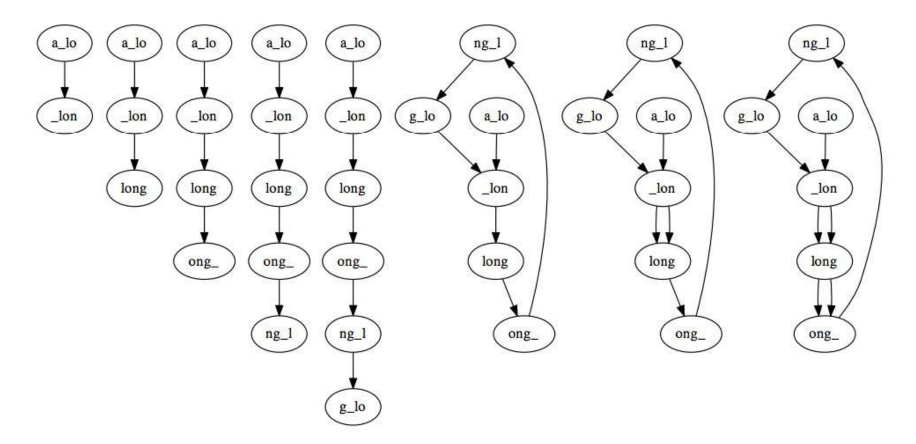
A procedure for making a De Bruijn graph for a genome

Assume *perfect sequencing* where each length-*k* substring is sequenced exactly once with no errors

Pick a substring length k: 5

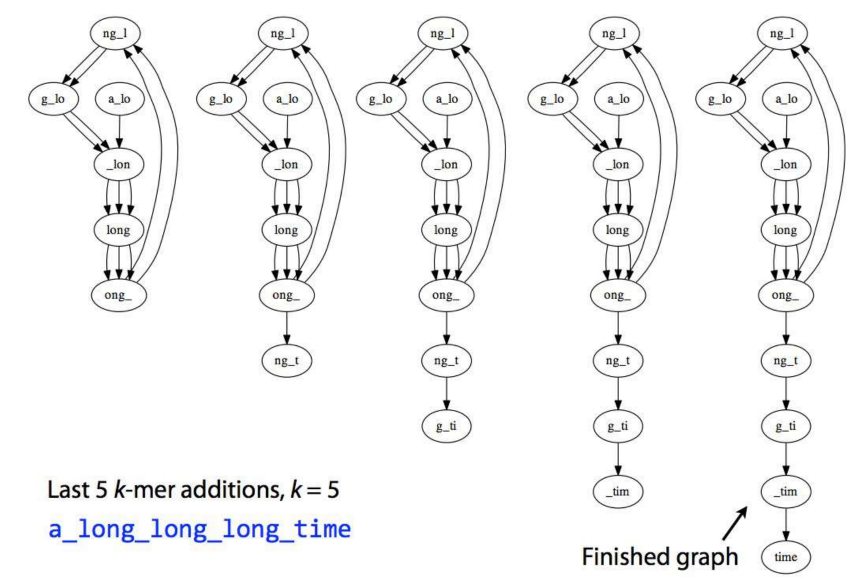
Start with an input string: $a_long_long_long_long_time$ Take each k mer and split
into left and right k-1 mers $long_$ Long ong_long ong_

Add k-1 mers as nodes to De Bruijn graph (if not already there), add edge from left k-1 mer to right k-1 mer



First 8 k-mer additions, k = 5

a_long_long_long_time



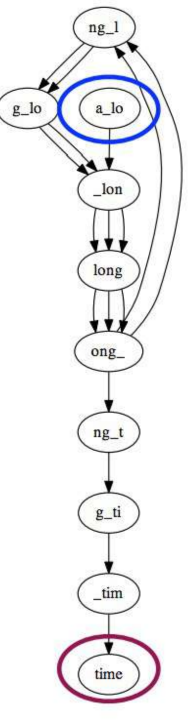
With perfect sequencing, this procedure always yields an Eulerian graph. Why?

Node for *k*-1-mer from left end is semi-balanced with one more outgoing edge than incoming *

Node for *k*-1-mer at right end is semi-balanced with one more incoming than outgoing *

Other nodes are balanced since # times k-1-mer occurs as a left k-1-mer = # times it occurs as a right k-1-mer

* Unless genome is circular



De Bruijn graph with actual data

Assuming perfect sequencing, procedure yields graph with Eulerian walk that can be found efficiently.

We saw cases where Eulerian walk corresponds to the original superstring. Is this always the case?

g_lo a lo lon long ong_ ng_t g_ti _tim time

When k-mer is repeat

No: graph can have multiple Eulerian walks, only one of which corresponds to original superstring

Right: graph for ZABCDABEFABY, k = 3

Alternative Eulerian walks:

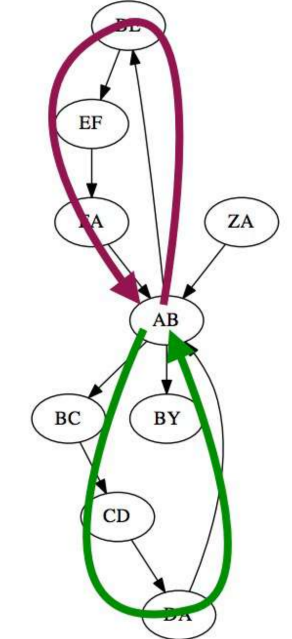
 $ZA \rightarrow AB \rightarrow BE \rightarrow EF \rightarrow FA \rightarrow AB \rightarrow BC \rightarrow CD \rightarrow DA \rightarrow AB \rightarrow BY$

 $\mathsf{ZA} \to \mathsf{AB} \to \mathsf{BC} \to \mathsf{CD} \to \mathsf{DA} \to \mathsf{AB} \to \mathsf{BE} \to \mathsf{EF} \to \mathsf{FA} \to \mathsf{AB} \to \mathsf{BY}$

These correspond to two edge-disjoint directed cycles joined by node AB

AB is a repeat: ZABCDABEFABY





When k-mer is repeat (in practice)

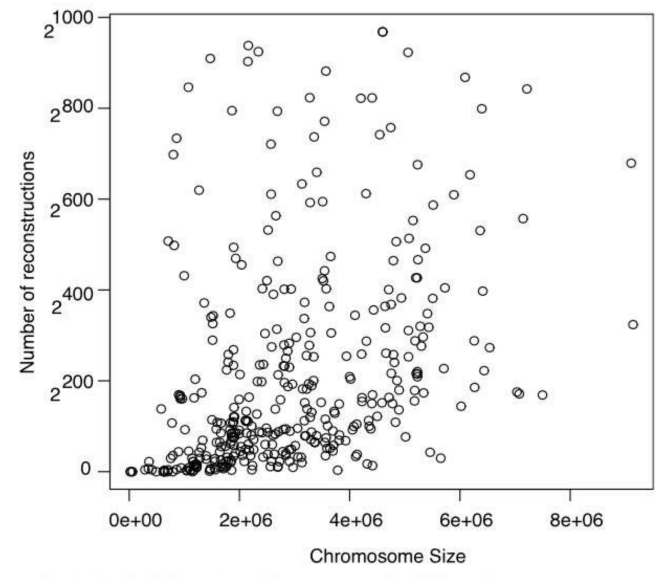
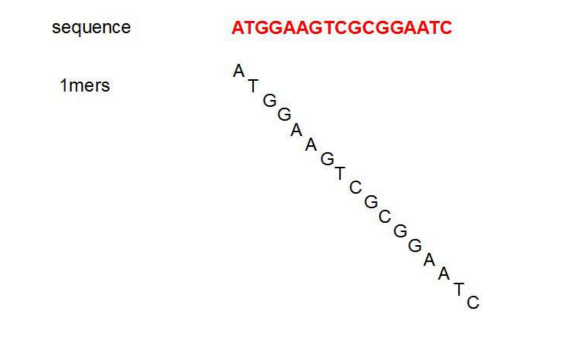


Figure 2 Number of words consistent with genome graphs. The

Kingsford et al., 2010

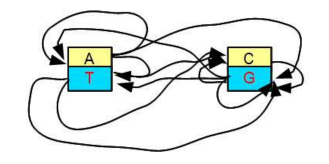
Impact of changing kmer size



Example of a kmer of 1 basically means A,C,T,G are all repeats..

Larger kmer will span more small repeat less than kmer size, but likely to have less overlap

de Bruijn graph

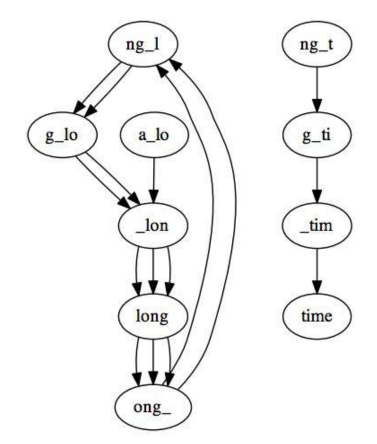


http://www.homolog.us/Tutorials/index.php?p=2.4&s=1

Low coverage = disconnected graph

Gaps in coverage can lead to disconnected graph

Graph for a_long_long_time, k = 5 but omitting ong_t:



Connected components are individually Eulerian, overall graph is not

Coverage difference = not Eulerian

Differences in coverage also lead to non-Eulerian graph

Graph for a_long_long_long_time, k = 5 but with extra copy of ong_t:

Graph has 4 semi-balanced nodes, isn't Eulerian

ng_l g_lo a lo lon long ong_ ng t g_ti _tim time

Gaining assembly as Eulerian walk is appealing, not many practical cases impede this:

- Uneven coverage, sequencing errors, make graph non-Eulerian
- Repeats produces many possible walks

But there is one major advantage of De Bruijn graph over OLC

Computationally efficient

Efficiency

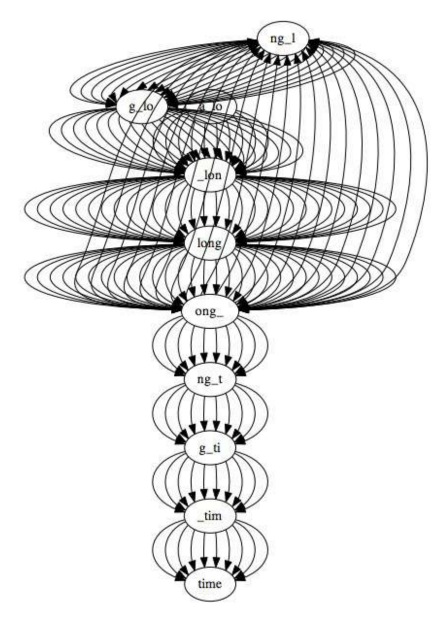
Assume you have 5 billion reads to assemble

- Not uncommon nowadays in some plant species
- OLC: 12.5 quadrillion overlaps to compute first Even 1 million overlap per sec will equate to **400 years**

DBG: depends on genome size

Size of De Bruijn graph depends on genome size

Usually ~50X paired end reads in coverage

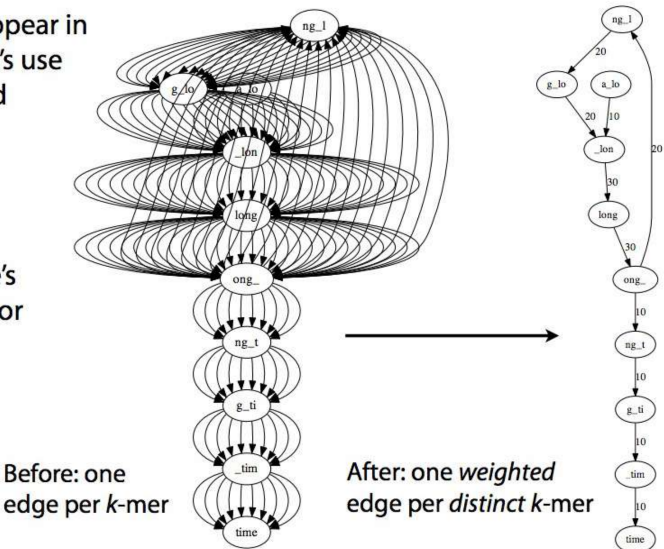


Size of De Bruijn graph depends on genome size

Same edge might appear in dozens of copies; let's use edge *weights* instead

Weight = # times k-mer occurs

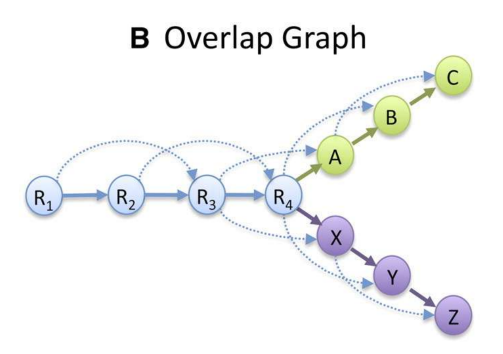
Using weights, there's one *weighted* edge for each *distinct* k-mer

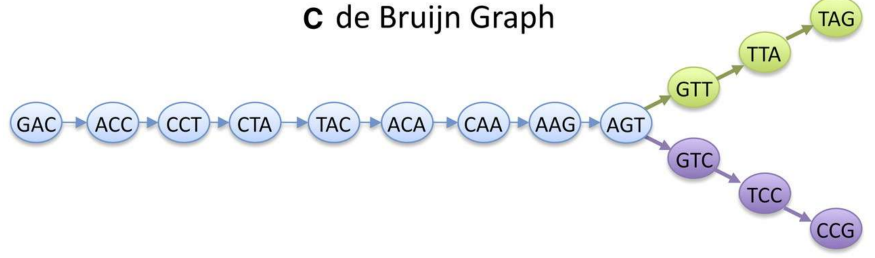


Summary

A Read Layout

- R₁: GACCTACA
- R₂: ACCTACAA
- R₃: CCTACAAG
- R₄: CTACAAGT
- A: TACAAGTT
- B: ACAAGTTA
- C: CAAGTTAG
- X: TACAAGTC
- Y: ACAAGTCC
- Z: CAAGTCCG





Schatz M C et al. Genome Res. 2010;20:1165-1173

Summary

Advantage of DBG:

Time to build based on Genome size (G) or total length of reads (N) For OLC: time to build overlap graph is based on number of reads

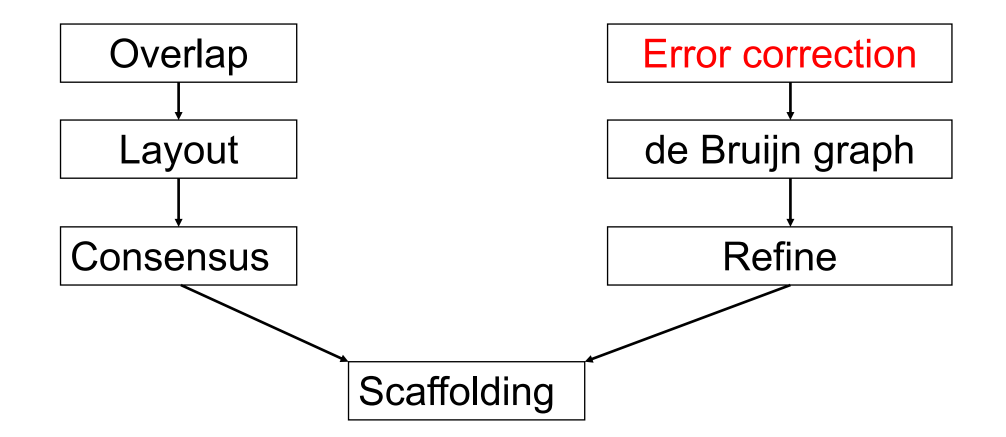
But:

DBG not flexible: only overlap of **fixed length** (=kmer) can't solve repeat with repeat > kmer

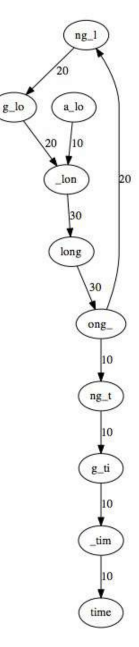
Read information is lost: All reads are split into kmers. (A lot of work on later DBG assemblers are put in this)

Tradeoff between DBG and OLC needed Some improve over existing approach: Spades Some combine both: Masurca

OLC and DBG assemblers



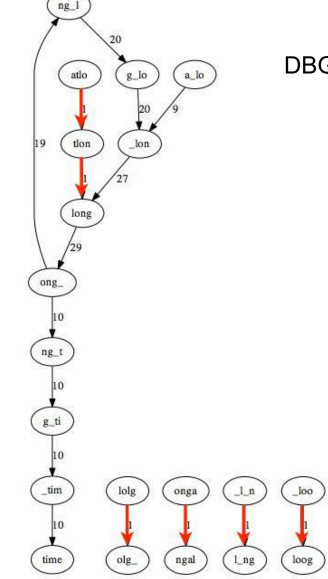
Error in graphs



DBG from perfect reads (10X)

As you can see all weighted **edges** have high coverage

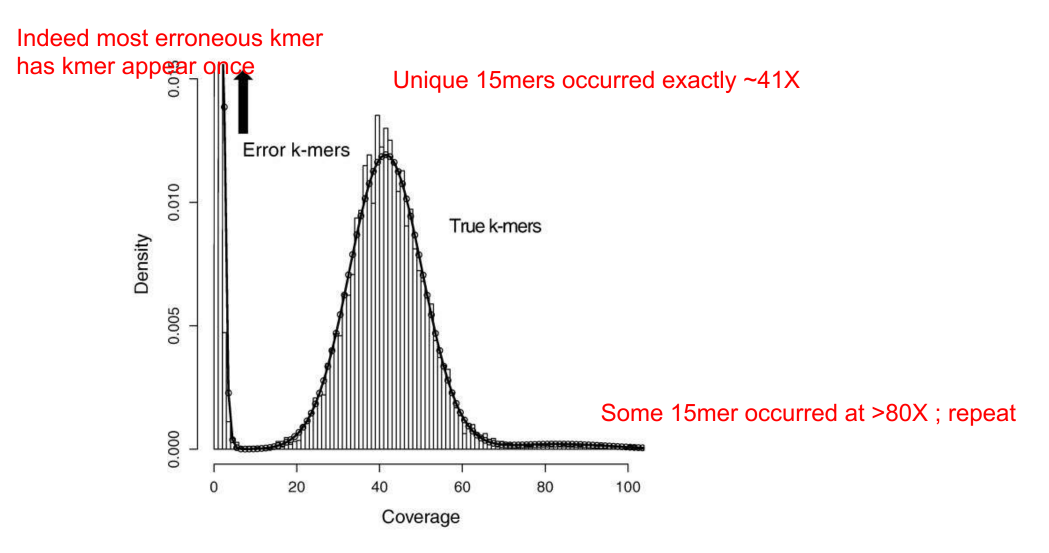
Some higher than the other; this is obviously repeat



DBG from reads with some errors

http://www.cs.jhu.edu/~langmea/resources/lecture_notes/error_correction.pdf

Kmer coverage



The mean and variance for true k-mers are 41 and 77 suggesting that a coverage bias exists as the variance is almost twice the theoretical 41 suggested by the Poisson distribution

Kelley et al., (2010)

Choosing the right kmer

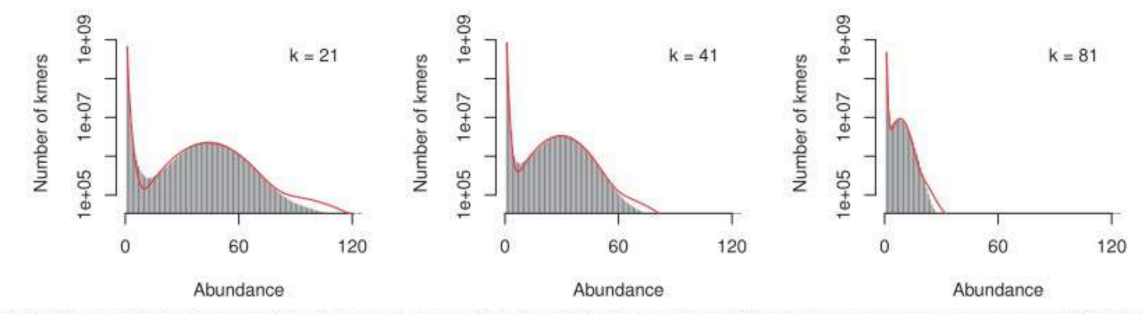


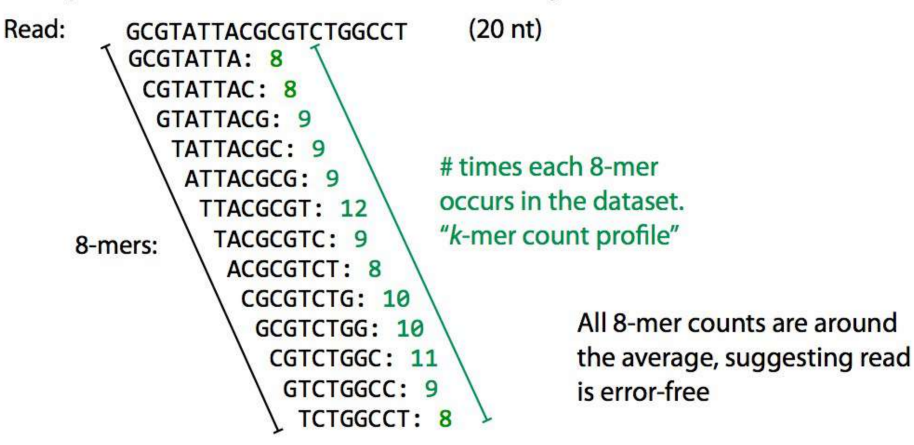
Fig. 2. The abundance histograms for chr14 with k values of 21, 41 and 81 (on a y log scale). Each plot also shows a curve corresponding to the optimized statistical model (haploid)

Chikhi and Medvedev et al., (2014)

Error correction: rationale

Idea: errors tend to turn frequent k-mers to infrequent k-mers, so corrections should do the reverse

Say we have a collection of reads where each distinct 8-mer occurs an average of ~10 times, and we have the following read:



http://www.cs.jhu.edu/~langmea/resources/lecture_notes/error_correction.pdf

Error correction: rationale

Suppose there's an error

```
Read:
        GCGTACTACGCGTCTGGCCT
         GCGTACTA: 1
                                            k-mer count profile has
          CGTACTAC: 3
                            Below average
                                            corresponding stretch of
           GTACTACG: 1
                                            below-average counts
            TACTACGC: 1
             ACTACGCG: 2
              CTACGCGT: 1
               TACGCGTC: 9
                ACGCGTCT: 8
                 CGCGTCTG: 10
                                     Around average
                  GCGTCTGG: 10
                   CGTCTGGC: 11
                     GTCTGGCC: 9
                      TCTGGCCT: 8
```

http://www.cs.jhu.edu/~langmea/resources/lecture_notes/error_correction.pdf

Error correction: rationale

k-mer count profiles when errors are in different parts of the read:

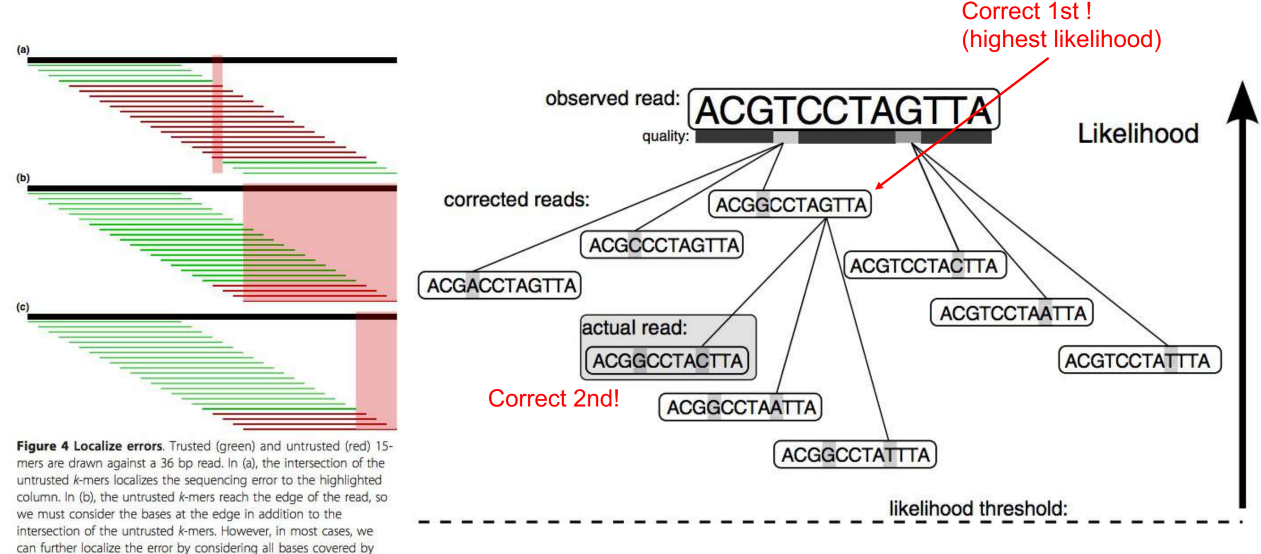
GCGTACTACGCGTCTGGCCT GCGTATTACACGTCTGGCCT GCGTATTACGCGTCTGGTCT GCGTACTA: 1 GCGTATTA: 8 GCGTATTA: 8 CGTATTAC: 8 CGTACTAC: 3 CGTATTAC: 8 GTACTACG: 1 GTATTACA: 1 GTATTACG: 9 TACTACGC: 1 TATTACAC: 1 TATTACGC: 9 ACTACGCG: 2 ATTACACG: 1 ATTACGCG: 9 CTACGCGT: 1 TTACACGT: 1 TTACGCGT: 12 TACGCGTC: 9 TACACGTC: 1 TACGCGTC: 9 ACGCGTCT: 8 ACACGTCT: 2 ACGCGTCT: 8 CGCGTCTG: 10 CACGTCTG: 1 CGCGTCTG: 10 GCGTCTGG: 10 GCGTCTGG: 10 GCGTCTGG: 10 CGTCTGGC: 11 CGTCTGGC: 11 CGTCTGGT: 1 GTCTGGCC: 9 GTCTGGCC: 9 GTCTGGTC: 2 TCTGGCCT: 8 TCTGGCCT: 8 TCTGGTCT: 1

http://www.cs.jhu.edu/~langmea/resources/lecture_notes/error_correction.pdf

Localize error and correct

the right-most trusted k-mer to be correct and removing them from

the error region as shown in (c).



Kelley et al., (2010)

Velvet: first de Bruijn graph assembler

- Cited 8812 times
- Still being used in some metagenomics dataset

1

Resource

Velvet: Algorithms for de novo short read assembly using de Bruijn graphs

Daniel R. Zerbino and Ewan Birney¹

EMBL-European Bioinformatics Institute, Wellcome Trust Genome Campus, Hinxton, Cambridge CB10 1SD, United Kingdom

We have developed a new set of algorithms, collectively called "Velvet," to manipulate de Bruijn graphs for genomic sequence assembly. A de Bruijn graph is a compact representation based on short words (k-mers) that is ideal for high coverage, very short read (25–50 bp) data sets. Applying Velvet to very short reads and paired-ends information only, one can produce contigs of significant length, up to 50-kb N50 length in simulations of prokaryotic data and 3-kb N50 on simulated mammalian BACs. When applied to real Solexa data sets without read pairs, Velvet generated contigs of ~8 kb in a prokaryote and 2 kb in a mammalian BAC, in close agreement with our simulated results without read-pair information. Velvet represents a new approach to assembly that can leverage very short reads in combination with read pairs to produce useful assemblies.

[Supplemental material is available online at www.genome.org. The code for Velvet is freely available, under the GNU Public License, at http://www.ebi.ac.uk/~zerbino/velvet.]

Summary

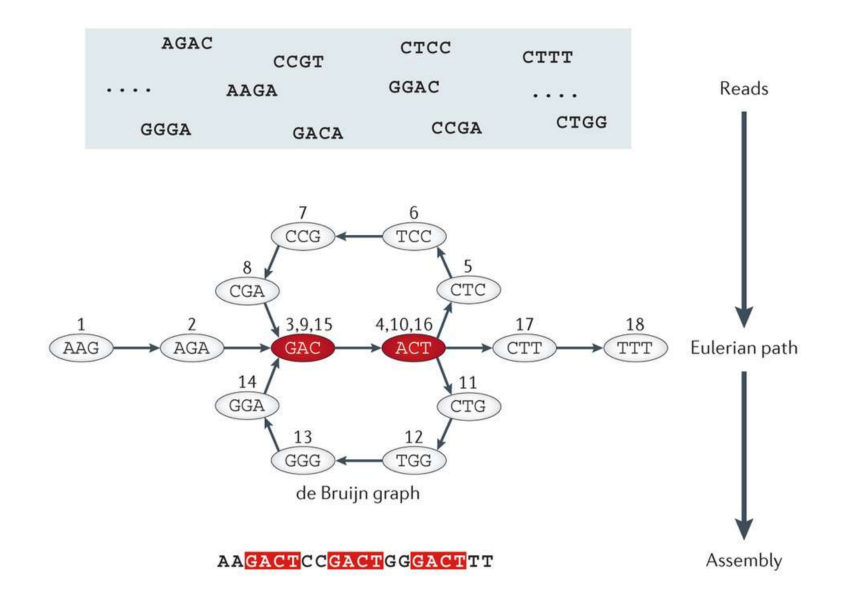
Error correction will **definitely** improve assembly

But, for it to work well:

Sequenced coverage should be high enough Choose kmer wisely otherwise we can't distinguish erroneous kmer from frequent kmers

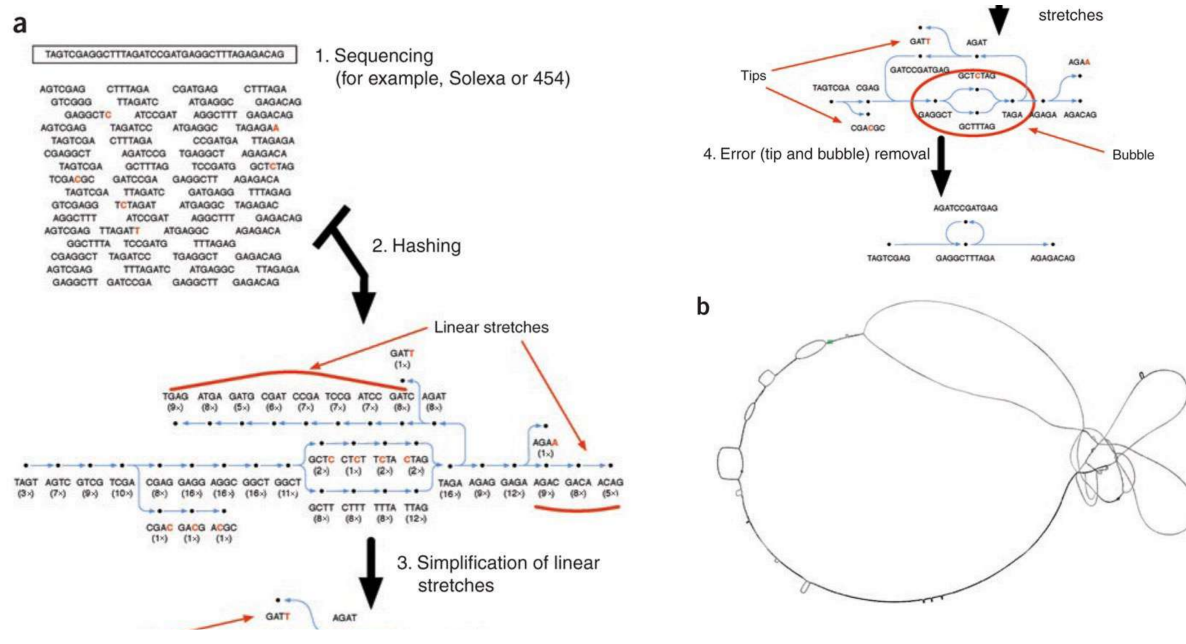
http://www.cs.jhu.edu/~langmea/resources/lecture_notes/assembly_dbg.pdf

Summary I

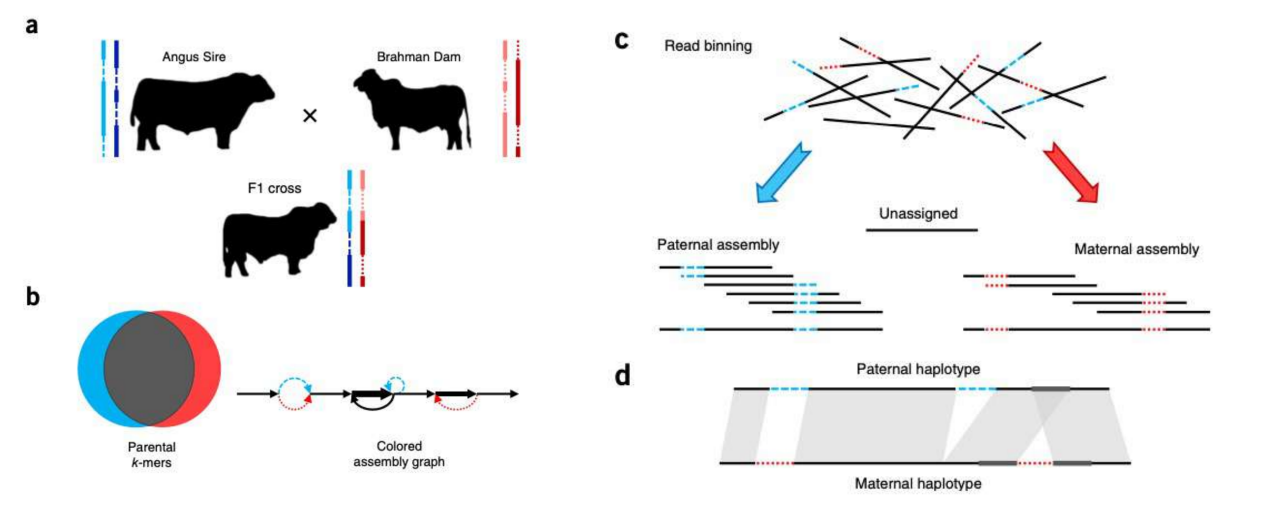


http://www.nature.com/nrg/journal/v14/n5/full/nrg3433.html

Summary II



Trio binning



Koren et al (2018) Nature Biotechnology

Binning (in metagenomics ; trios)

Keyword: MAG (metagenome-assembled genomes)

Advantage of metagenomics approach

Better classification with Increasing number of complete genomes Focus on whole genome based phylogeny (whole genome phylotyping)

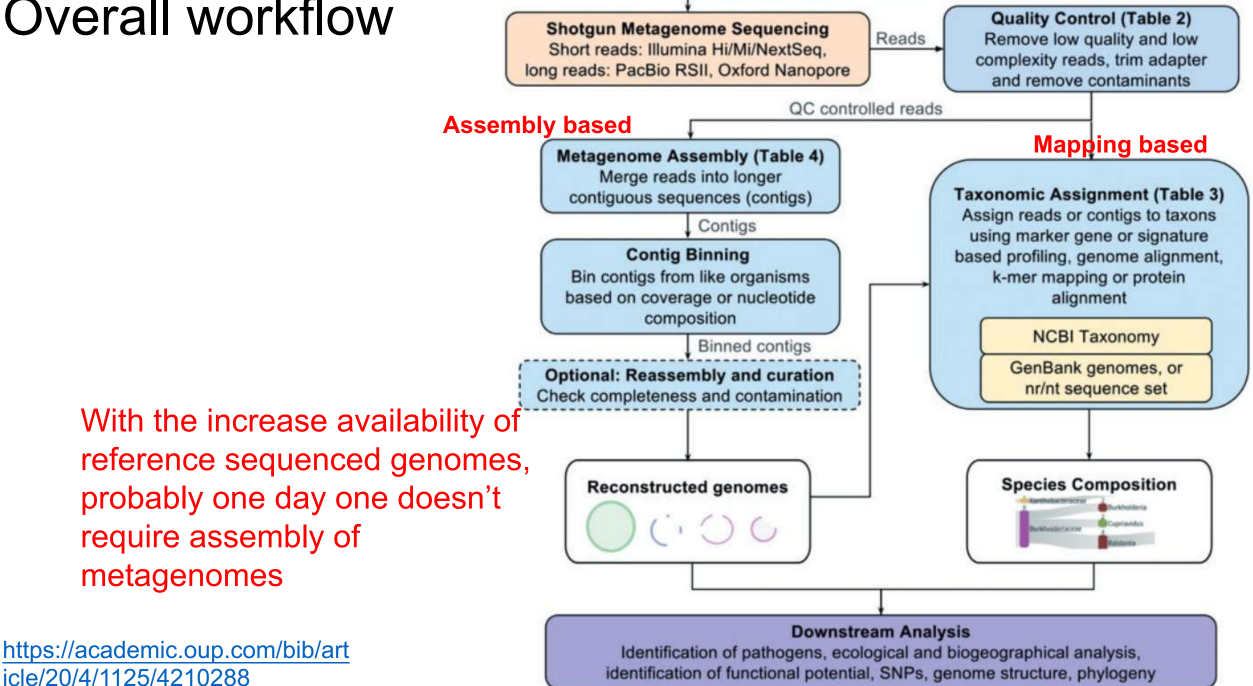
•Advantages No amplification bias like in 16S/ITS

Issues

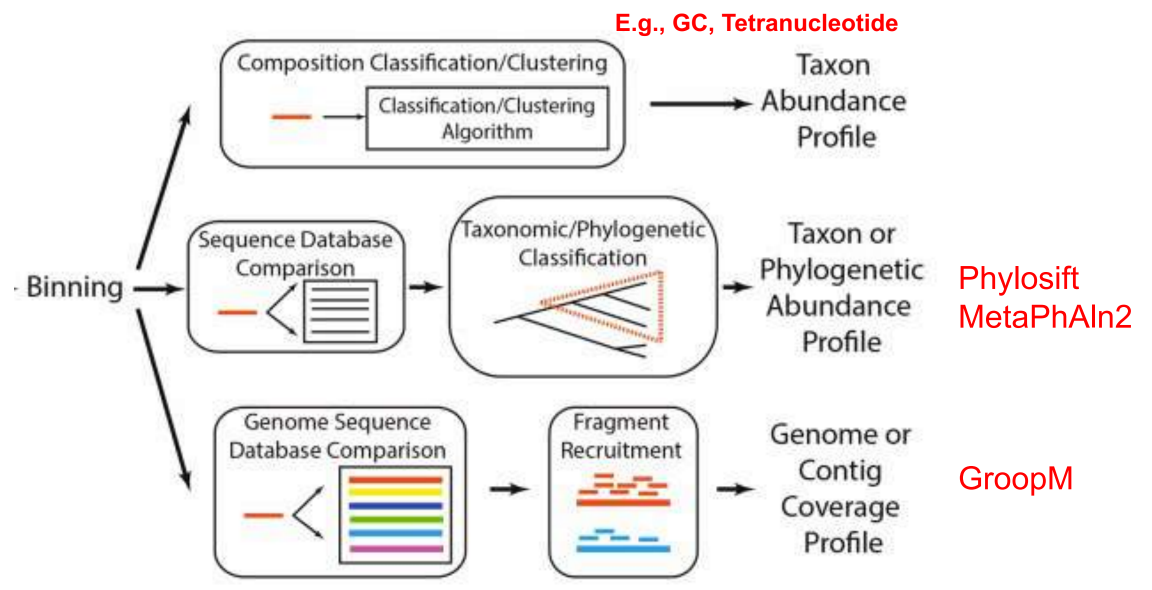
Poor sampling beyond eukaryotic diversity

Assembly of metagenomes is **challenging** due to uneven coverage Requires **high** depth of coverage

Overall workflow

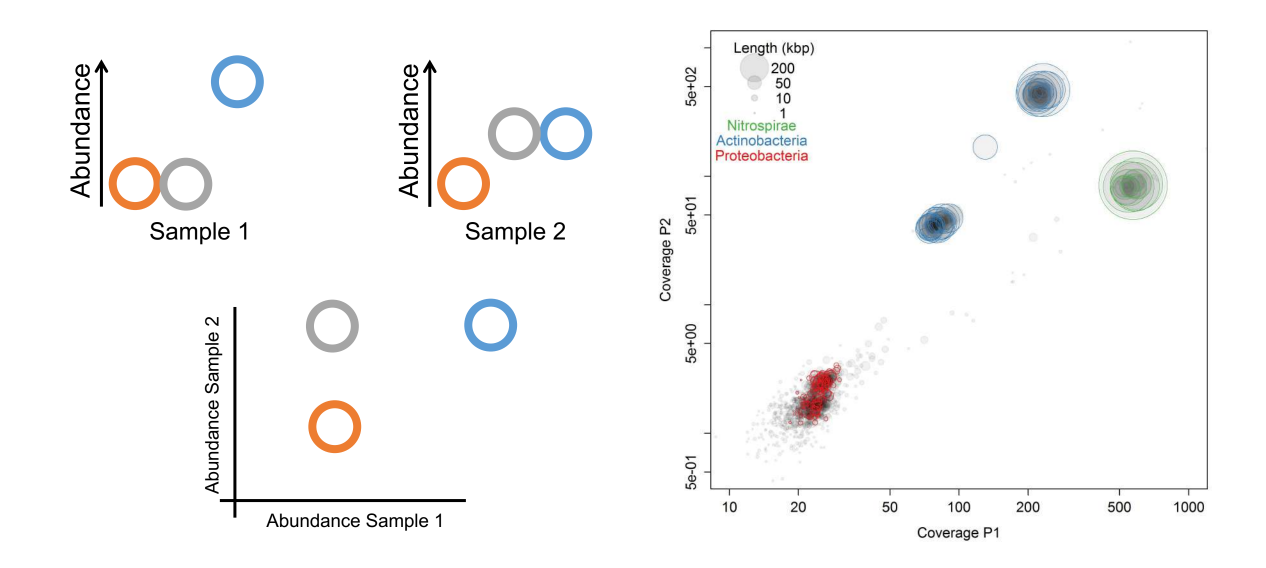


Binning methods



Sharpton (2014)

Example of binning based on differential coverage



H. Daims & C. Dorninger, DOME, University of Vienna

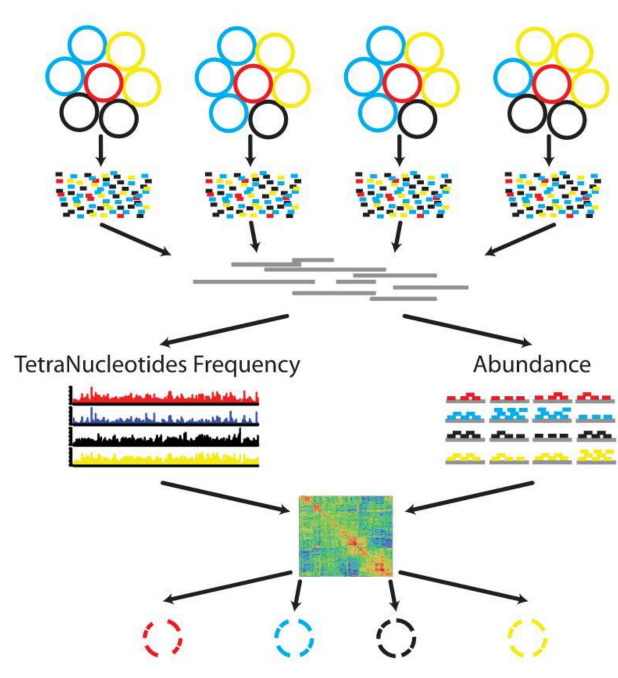
Binning methods: A combination of

Classification based on **sequence composition**:

- Advantage : all reads can be categorised into bins
- **Disadvantage**: no taxonomy / function of the bins.

Classification based on **sequence similarity (of known genes) Advantage**: One can determine taxonomy and function of reads. **Disadvantage**: reads with similarity can not be classified .

Metabat



Preprocessing

Samples from multiple sites or times

2

Metagenome libraries

3

Initial de-novo assembly using the combined library

MetaBAT

4

Calculate TNF for each contig

5 Calc

Calculate Abundance per library for each contig

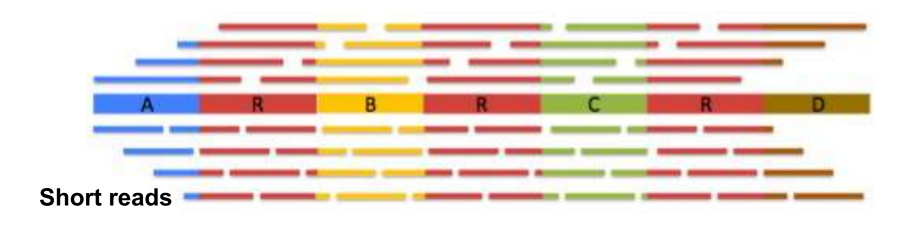
6

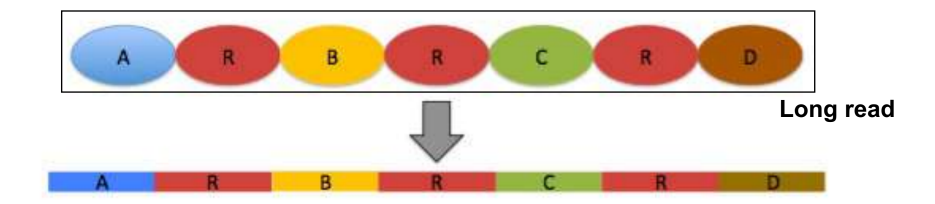
Calculate the pairwise distance matrix using pre-trained probabilistic models

Forming genome bins iteratively

https://peerj.com/articles/7359/

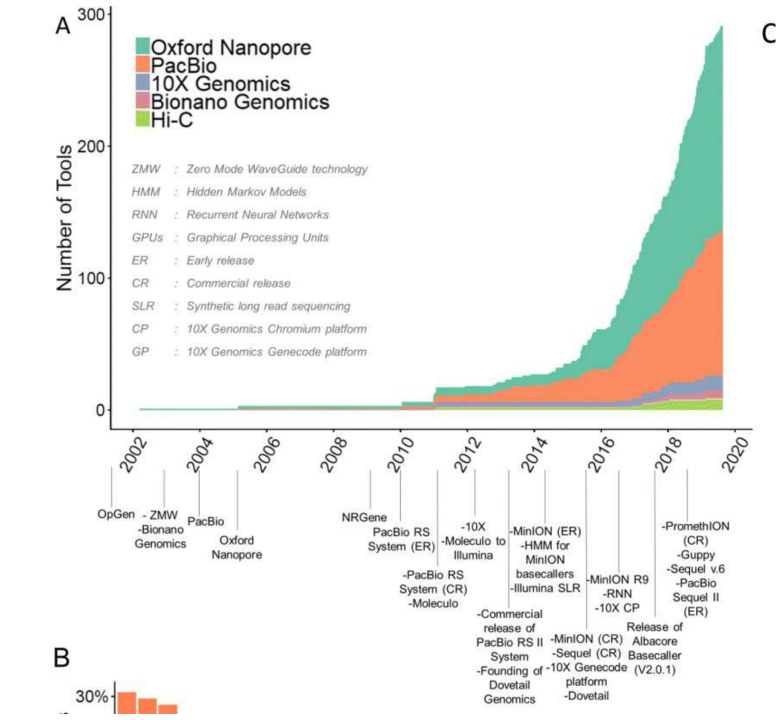
Long read technologies





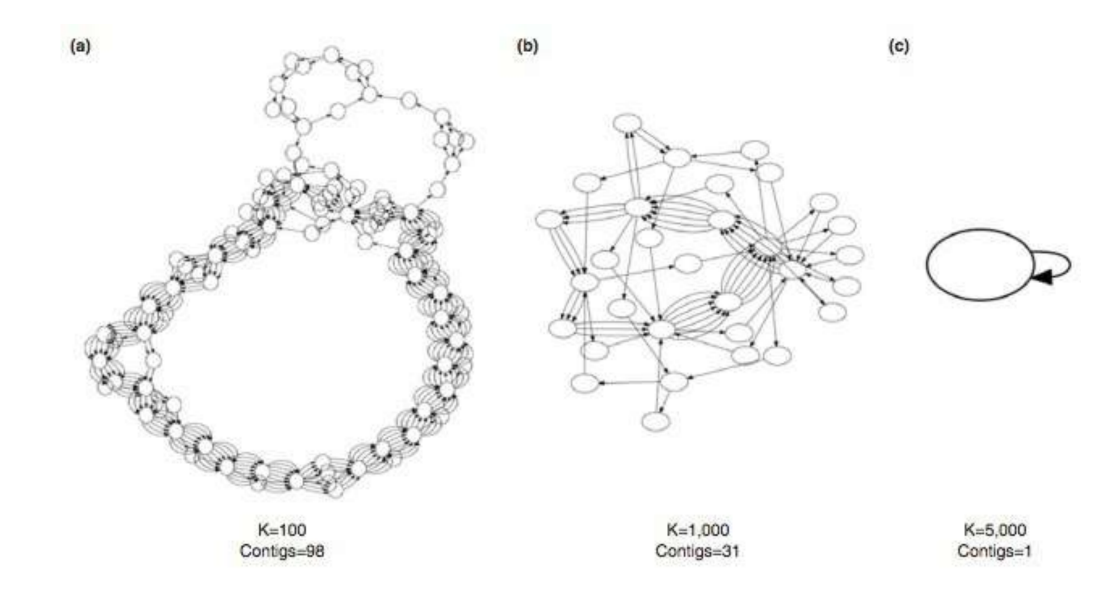
Credit: Michael Schatz

New tools continue to be developed



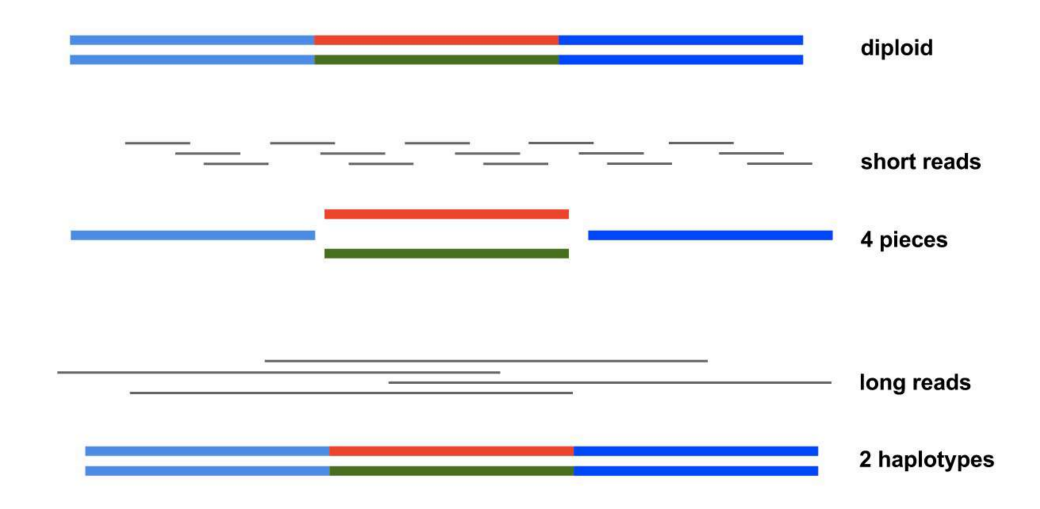
Amarasinghe et al (2020) Genome Biology

Simpler graphs

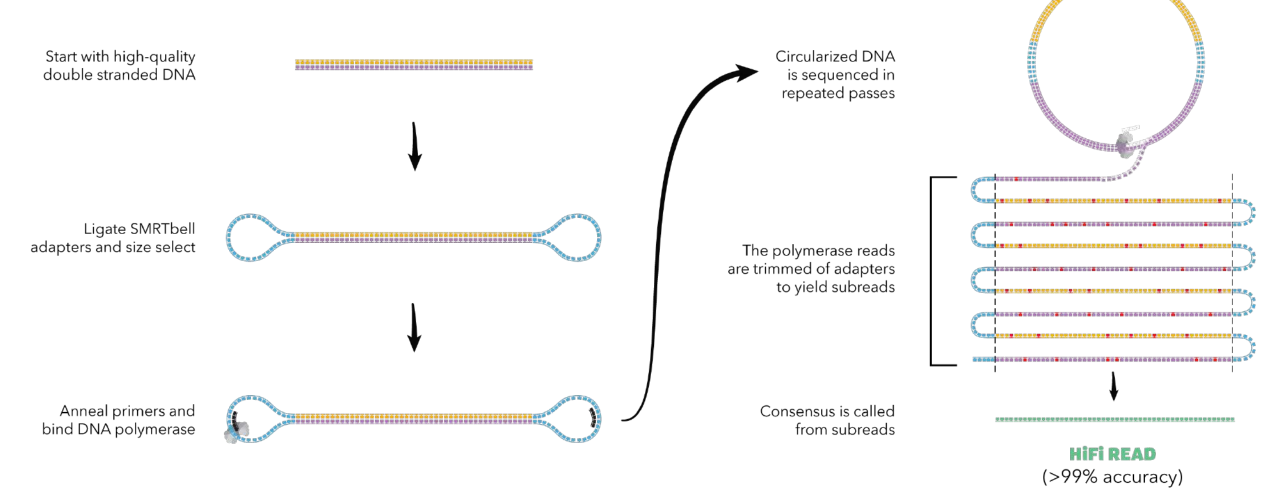


Roberts et al., 2013

Long reads can also resolve haplotypes (with sufficient coverage)



HiFi reads



https://www.pacb.com/smrt-science/smrt-sequencing/hifireads-for-highly-accurate-long-read-sequencing/

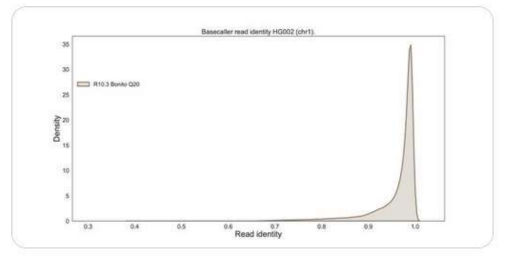


Miten Jain @mitenjain · Mar 11 Impressive numbers on human genome @nanopore data using Q20 early-access chemistry.

PEPPER-Margin-DeepVariant achieves amazing, precisionFDA level, performance (F1 scores >0.996) with modest coverage

@kishwarshafin @BenedictPaten @pichuan @acarroll_ATG

biorxiv.org/content/10.110...

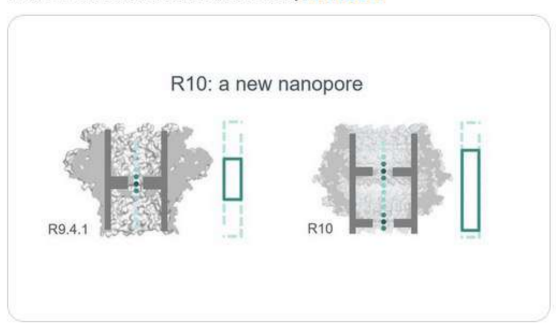




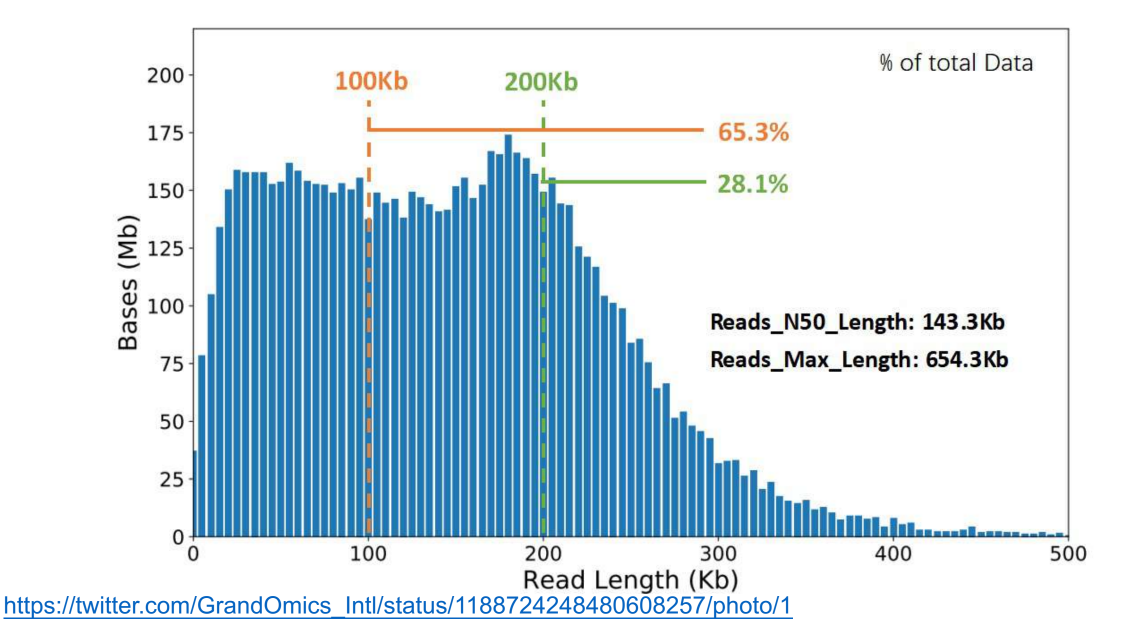
...

Alexander Wittenberg @AW_NGS · Mar 12

Just obtained amazing results on Fusarium spp genome using R10.3 @nanopore PromethION data, Bonito basecalling and Medaka consensus calling. Achieved chromosome-level assembly with QV52. That is >99.999% consensus accuracy! #RNGS21

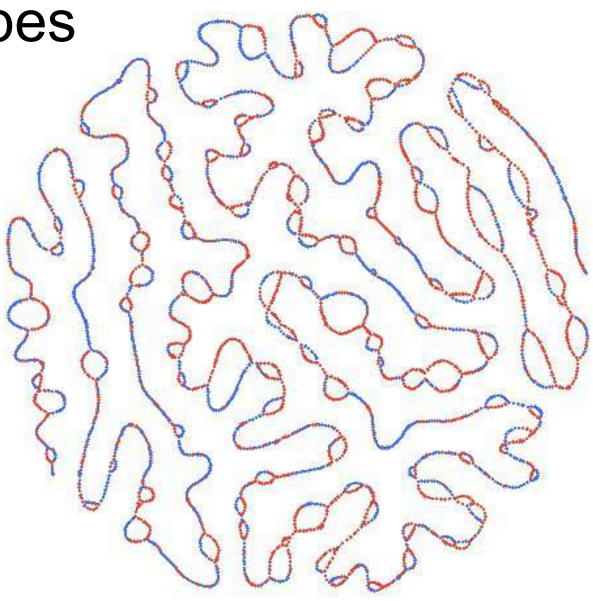


Read length and capacity go beyond



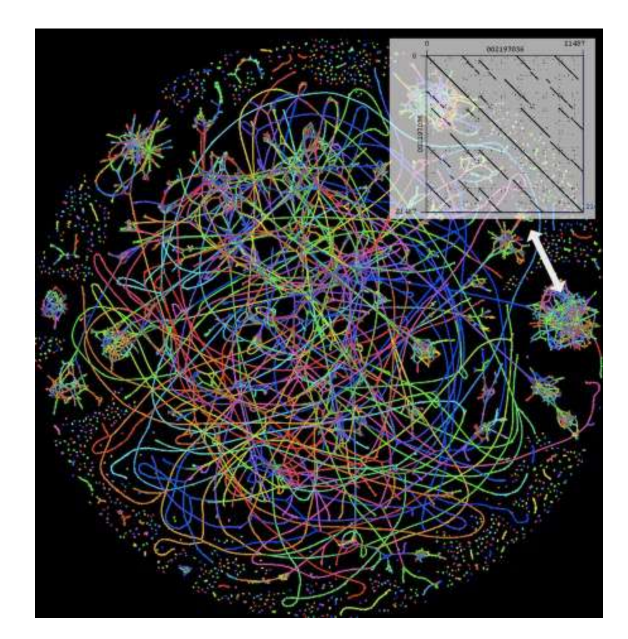
What you can do with long reads: Resolving haplotypes

Credit: Jason Chin Two genomes. One assembly. Two colors. Many bubbles. Game against entropy. http://t.co/uCPmxCRiZ6



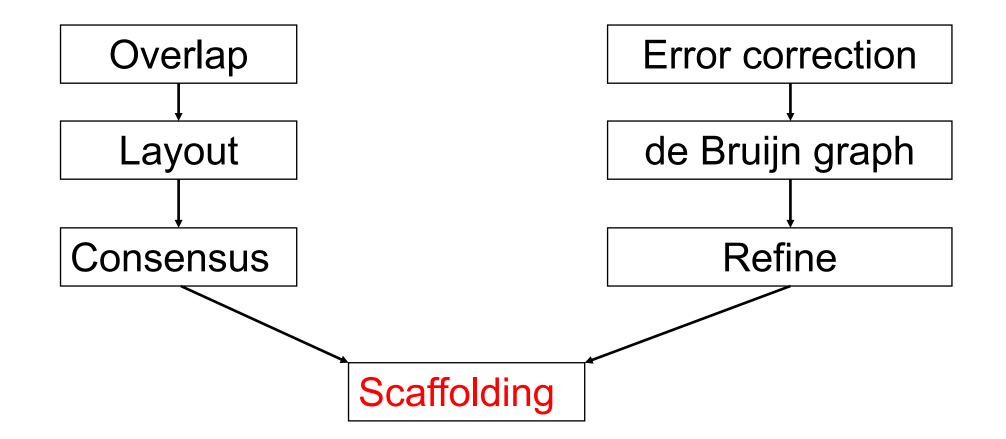
Current limitation of long reads..

Credit: Jason Chin What are those blobs in the genome assembly graph? Intriguing repeats that have no NCBI blast hit. http://t.co/2y7stBGs4W



Scaffolding

OLC and DBG assemblers

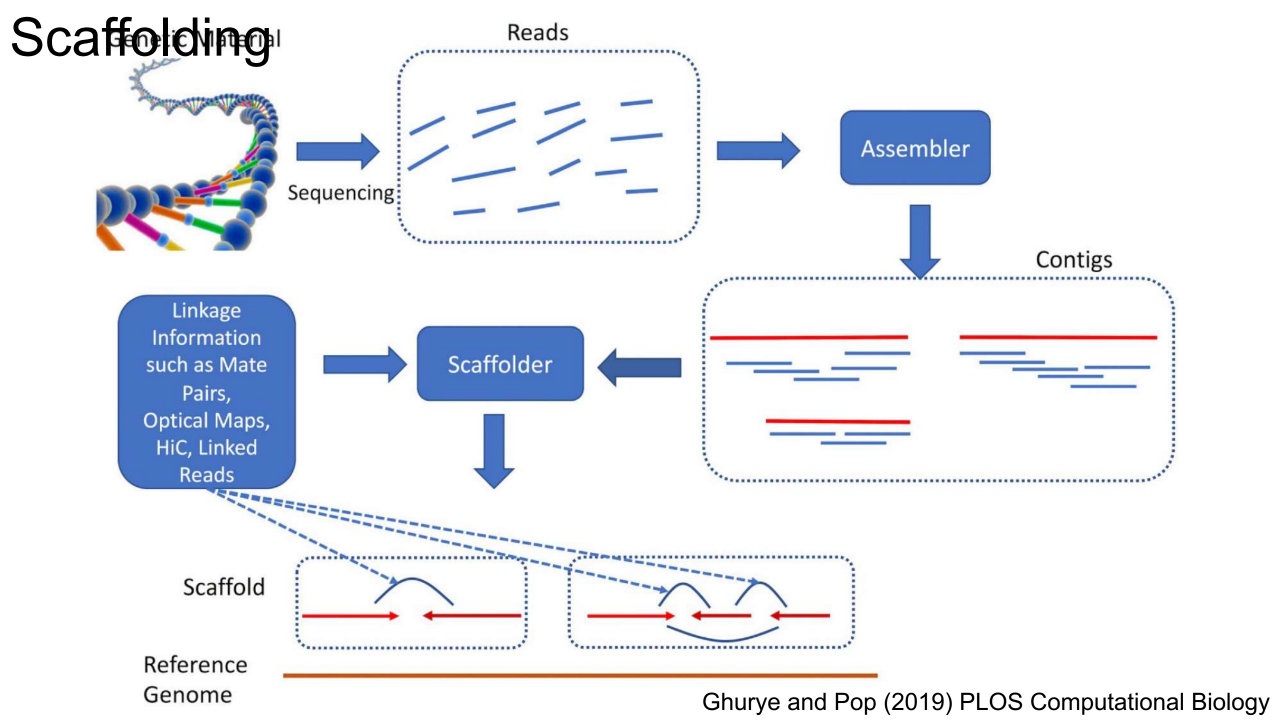


Scaffolding

OLC and DBG attempt to construct longest and most accurate **contigs** (**contig**uous stretch of assembled bases)

Scaffolding is to order and orient contigs with respect to each other

Various data types: Paired ends / Mate pairs Genetic map Additional long range information



Scaffolding: Paired end sequencing

Fragment GCATCATTGCCAATATATGGCTCTAGCATAAAACC GCATCATTG Mate 1

Because of technology limitation (usually ~150bp at each end), whole fragment is not sequenced. But the distance between two mates equals to length of fragment (insert size)

Scaffolding: Paired end sequencing

• DNA fragment (200-800 bp)

• Single end

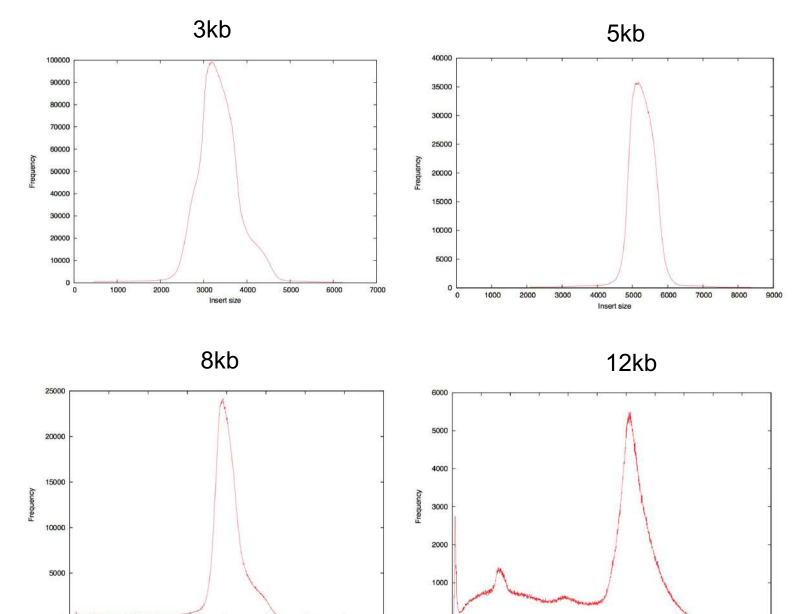
- Paired end (up to 800 bp span)
- Mate pair (up to 40 kbp span)

Examples

How to check insert size?

Remap the data back to the assembly

Problem can arise in larger insert sizes



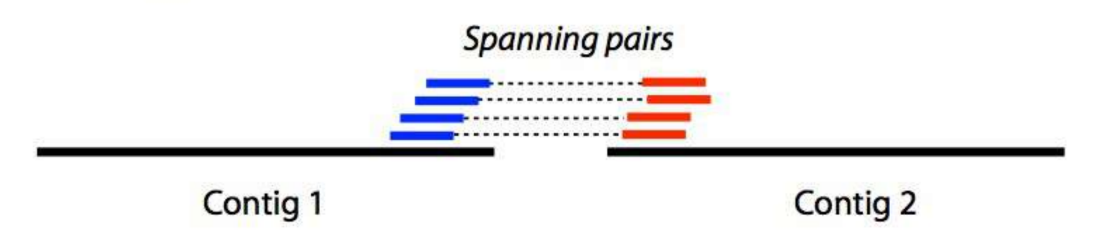
Insert size

Insert size

14000 16000 18000 20000 22000

Scaffolding (Illumina; obsoleted)

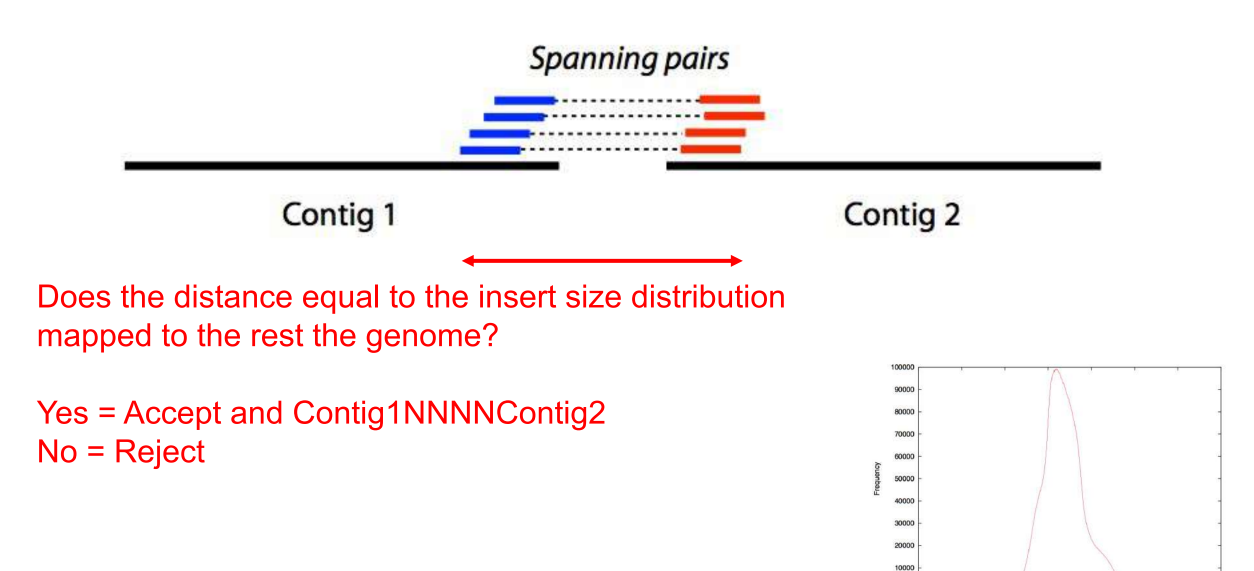
Say we have a collection of pairs and we assemble them as usual Assembly yields two contigs:



...and we discover that some of the mates at one edge of contig 1 are paired with mates in contig 2

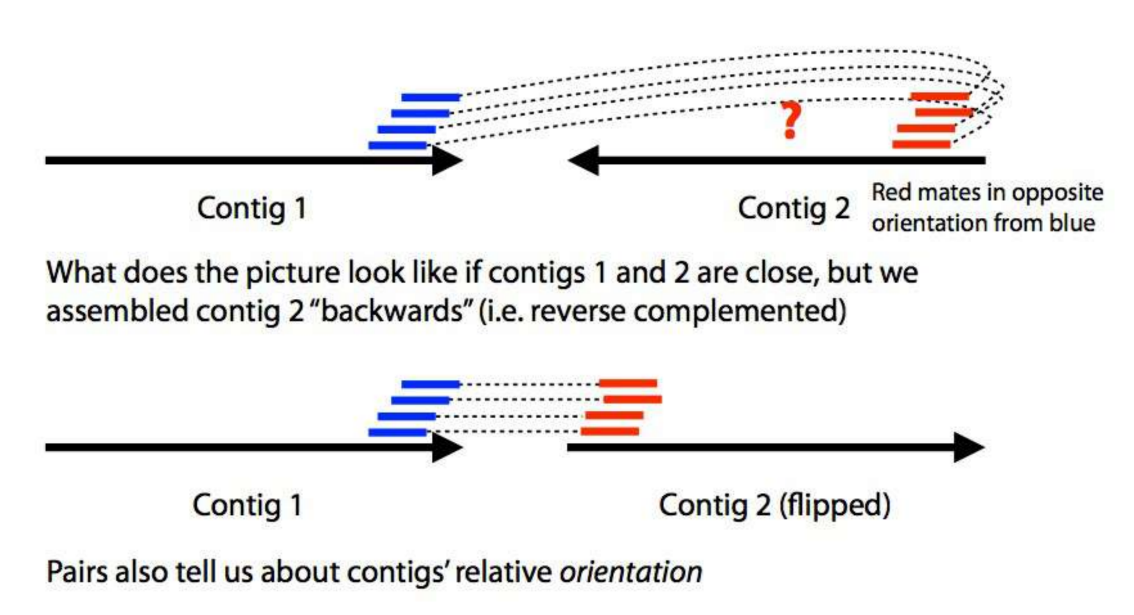
Call these spanning pairs

Scaffolding (Illumina; obsoleted)

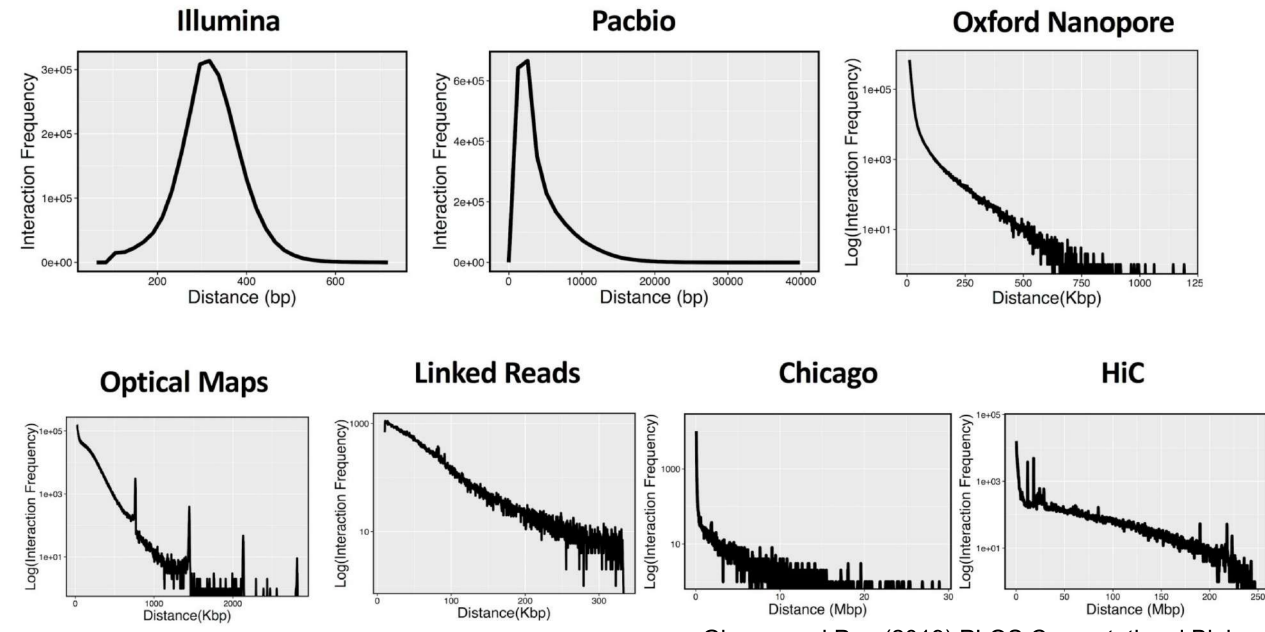


Insert size

Scaffolding (Illumina; obsoleted)



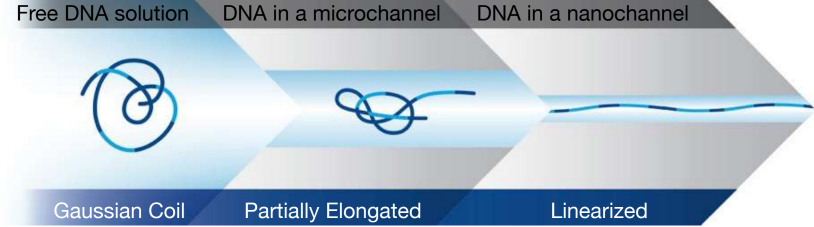
The genomic span covered by different technologies



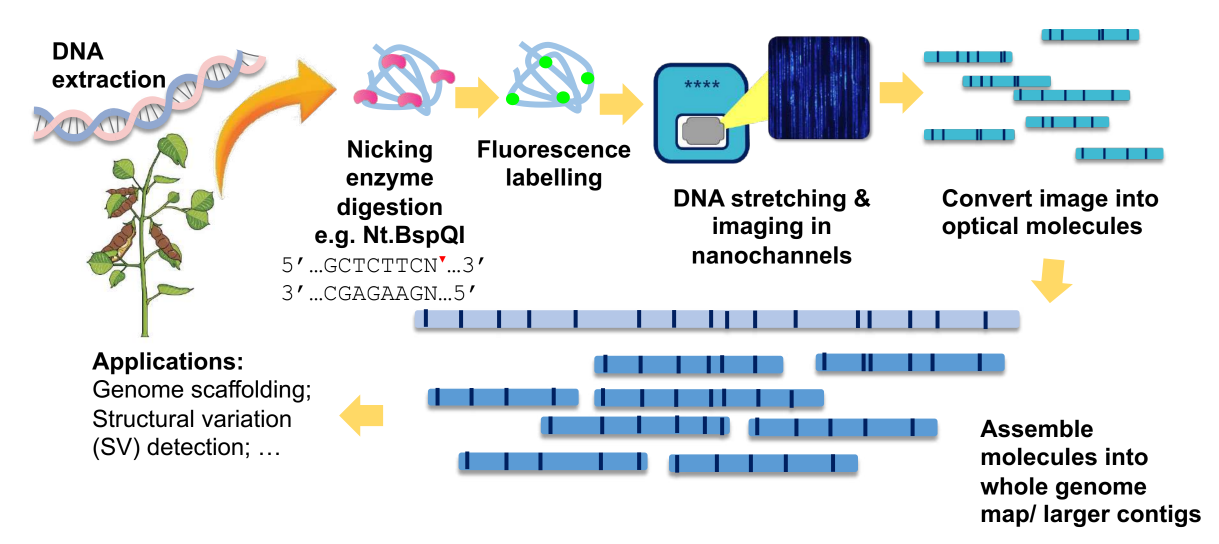
Ghurye and Pop (2019) PLOS Computational Biology

Optical map: DNA linearized in nanochannel array

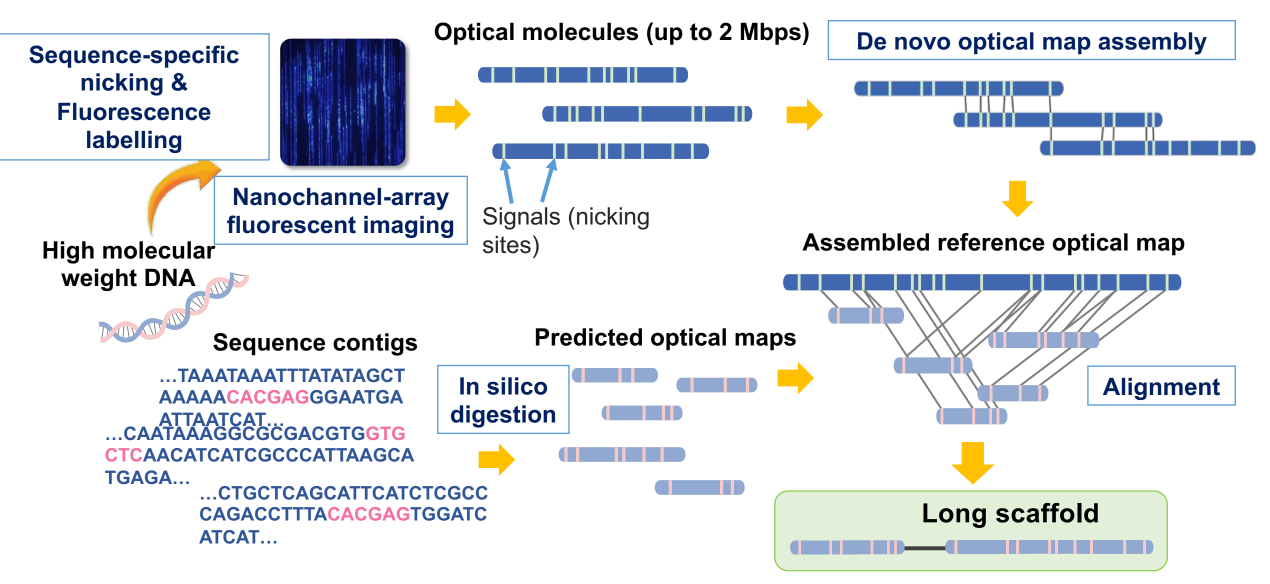


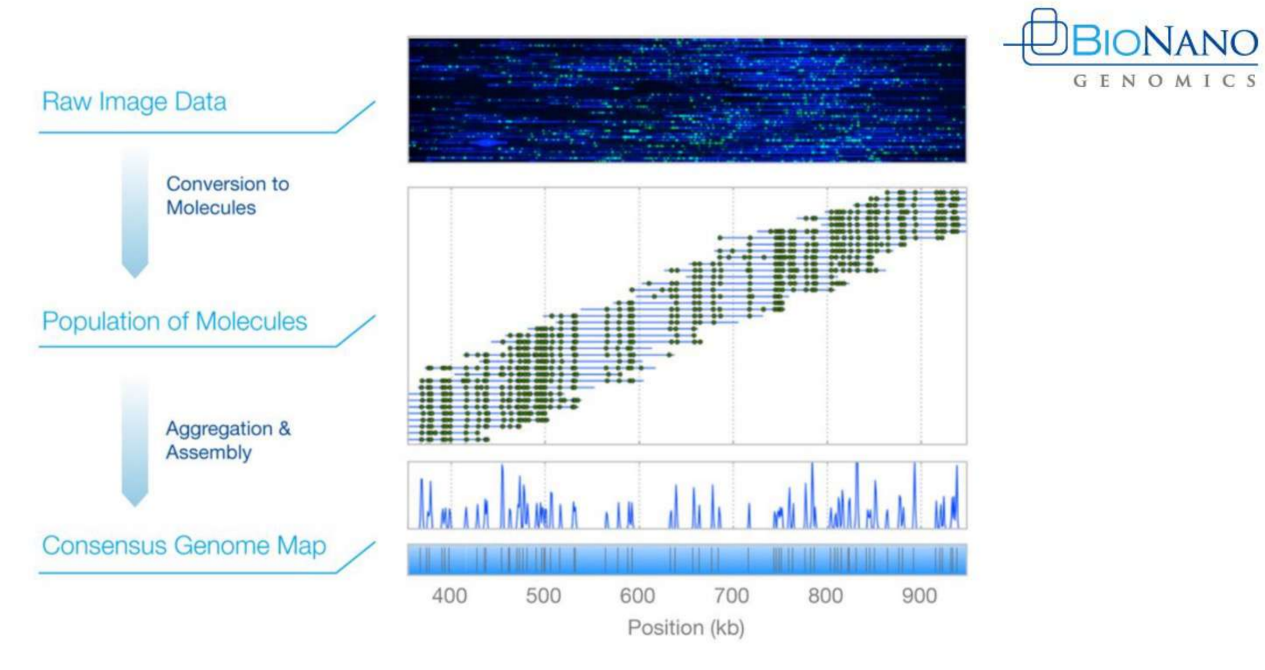


Workflow of an optical mapping procedure



Optical mapping-assisted scaffolding principles

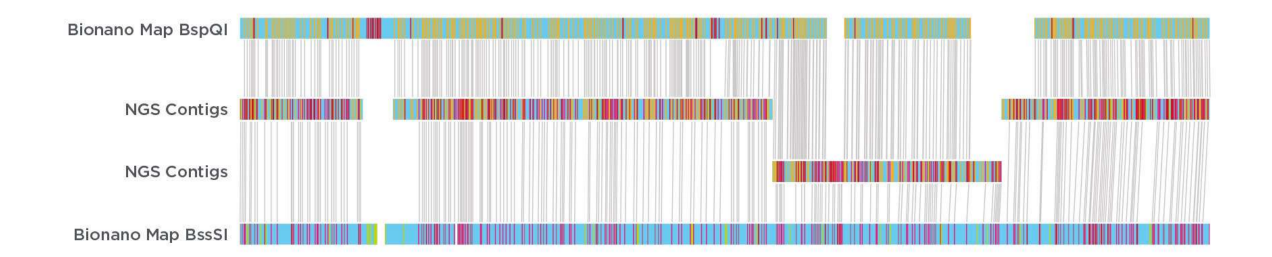




https://bionanogenomics.com/technology/platform-technology/



TWO ENZYME HYBRID SCAFFOLDING



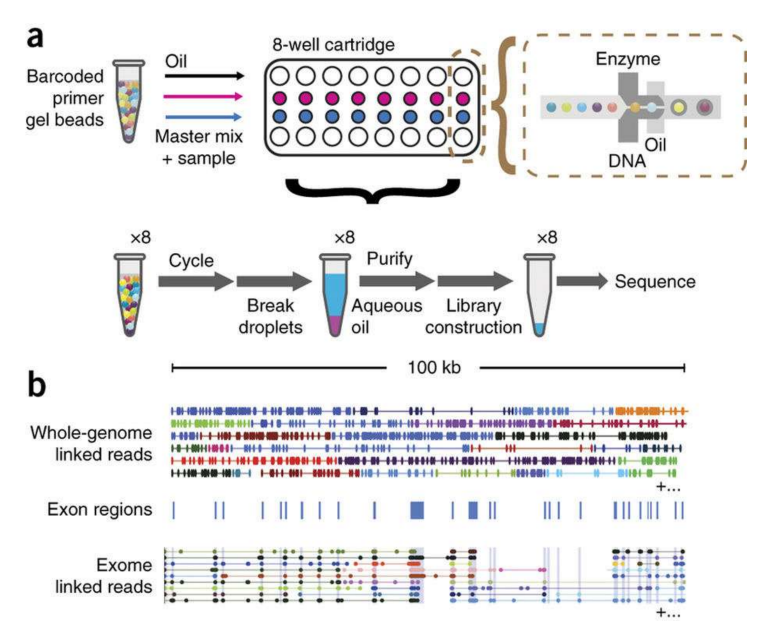
https://bionanogenomics.com/technology/platform-technology/



- Long range information from short reads using 14bp barcodes
- Very low input DNA (ng) and 20 mins preparation time
- 1ng of DNA is split across 100,000 Gel coated beads
- Single-cell available

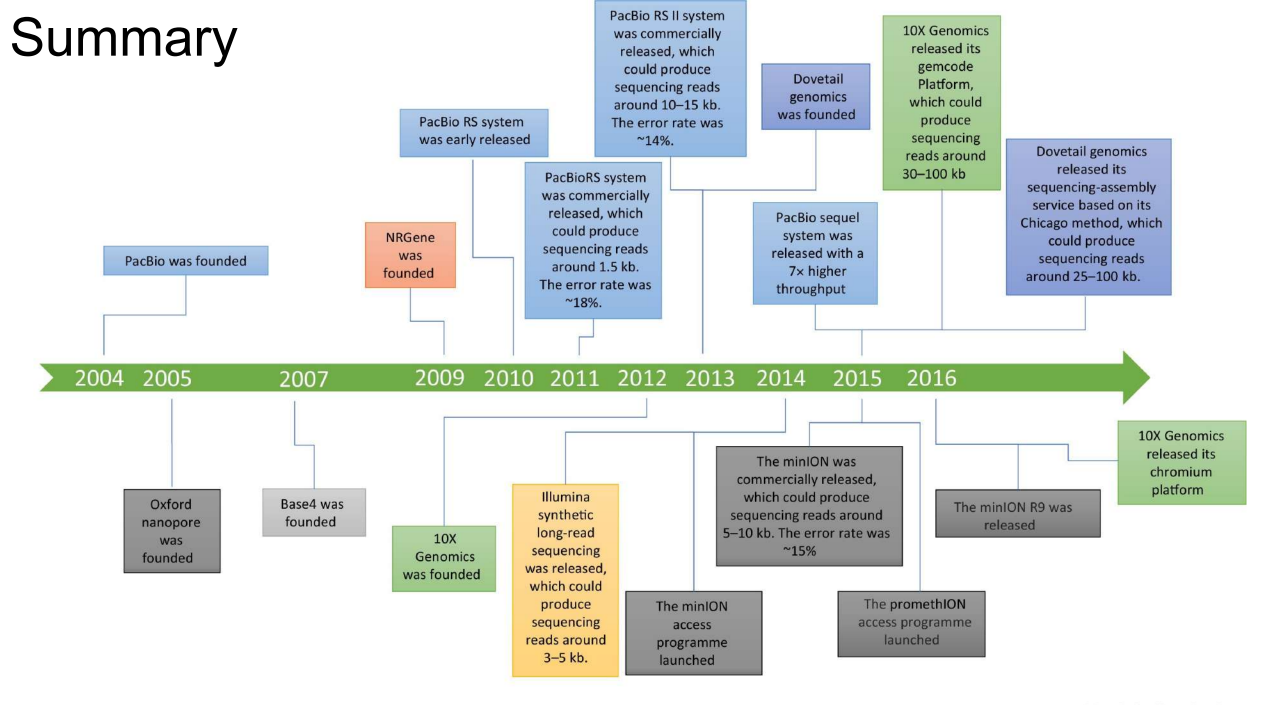
https://www.nature.com/articles/nbt.3432

Haplotyping germline and cancer genomes with high-throughput linked-read sequencing

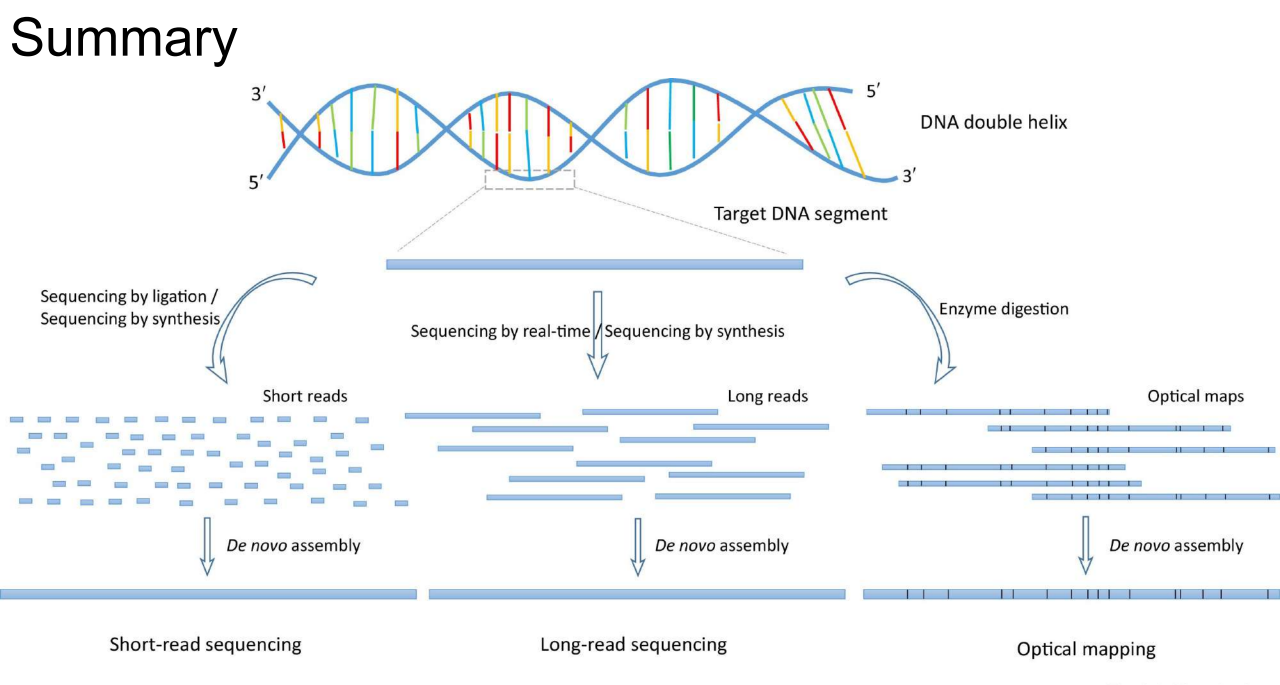


10X Genomics

https://www.youtube.com/watch?v=aUyFzwRFWJQ



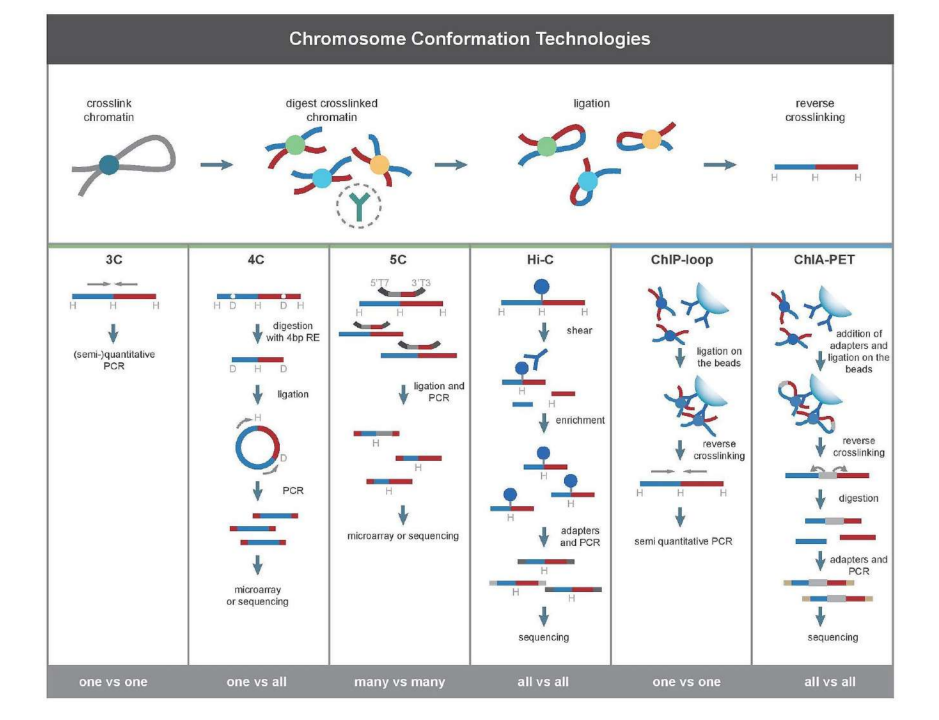
Trends in Biotechnology



Scaffolding using Chromosome conformation capture

Chromosome conformation capture techniques (often abbreviated to 3C technologies or 3C-based methods) are a set of molecular biology methods used to analyze the spatial organization of chromatin in a cell. These methods quantify the **number of interactions between genomic loci that are nearby in 3-D space**, but may be **separated by many nucleotides** in the linear genome

Flemmi the terr chroma		Kossel discove histone	ers	Waldem coins the chromos	ne term	Waddii postula epigen landsca	ates the netic	Structu DNA	ire of	Discoveri the chror fiber	les of	Chromo terrutor		Dekker innovat 3C tech	tes the	Simonis Dostie –	,	Lieberman Aiden - Hi Fullwood ChIA-PET	i-C, −
1879	1883	1884	1888	1889	1928	1942	1948	1953	1961	1973	1975	1982	1984	2002	2003	2006	2007	2009	2012
	Weisma connect chromat heredity	ts itin with		propose tinuity of tin during	heteroc		Hotchkis discovers ⁿ methylat	rs DNA	Lyon pos the princ XCI		Chambon the term nucleosoi		Lis innov the ChIP techniqu			Genome finished	Pugh inno the ChIP-s technique	seq	Ren discovers TADs in mammals
				, cic															



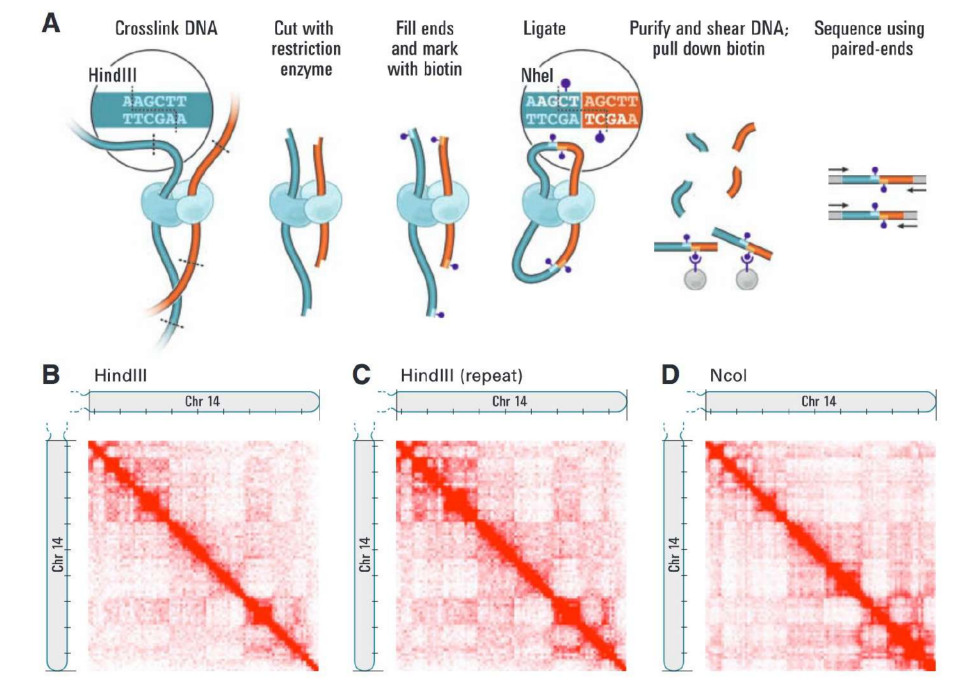
Comprehensive Mapping of Long-Range Interactions Reveals Folding Principles of the Human Genome

Erez Lieberman-Aiden,^{1,2,3,4}* Nynke L. van Berkum,⁵* Louise Williams,¹ Maxim Imakaev,² Tobias Ragoczy,^{6,7} Agnes Telling,^{6,7} Ido Amit,¹ Bryan R. Lajoie,⁵ Peter J. Sabo,⁸ Michael O. Dorschner,⁸ Richard Sandstrom,⁸ Bradley Bernstein,^{1,9} M. A. Bender,¹⁰ Mark Groudine,^{6,7} Andreas Gnirke,¹ John Stamatoyannopoulos,⁸ Leonid A. Mirny,^{2,11} Eric S. Lander,^{1,12,13}† Job Dekker⁵†

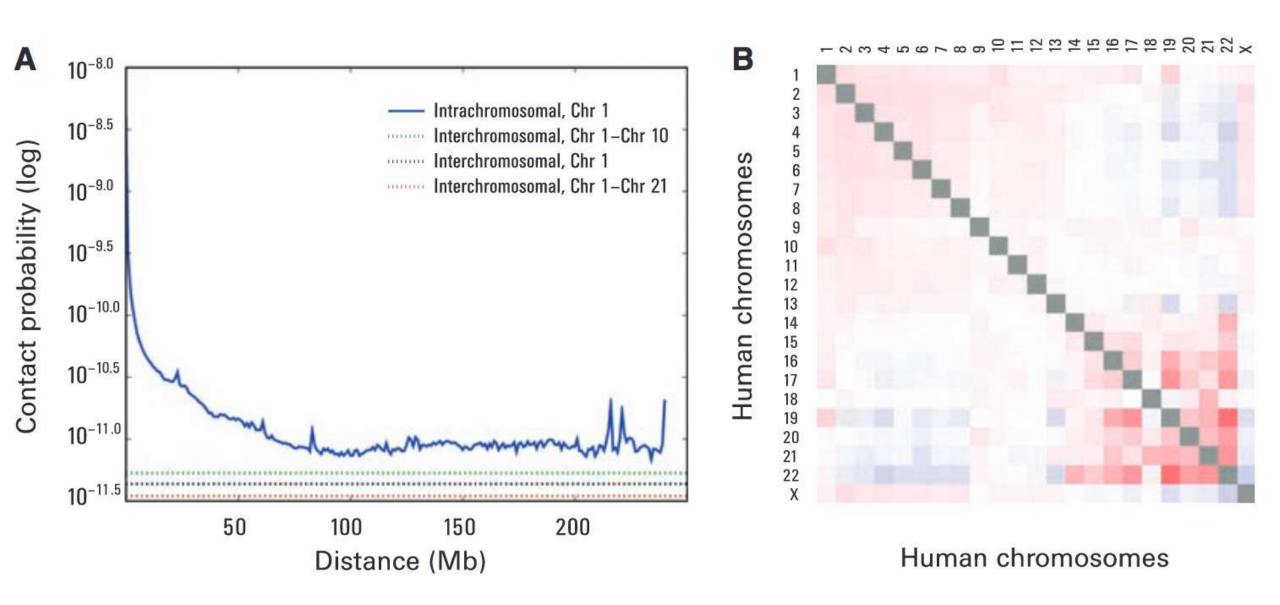
We describe Hi-C, a method that probes the three-dimensional architecture of whole genomes by coupling proximity-based ligation with massively parallel sequencing. We constructed spatial proximity maps of the human genome with Hi-C at a resolution of 1 megabase. These maps confirm the presence of chromosome territories and the spatial proximity of small, gene-rich chromosomes. We identified an additional level of genome organization that is characterized by the spatial segregation of open and closed chromatin to form two genome-wide compartments.

Lieberman-Aiden (2009) Science 10.1126/science.1181369

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Lieberman-Aiden (2009) Science 10.1126/science.1181369



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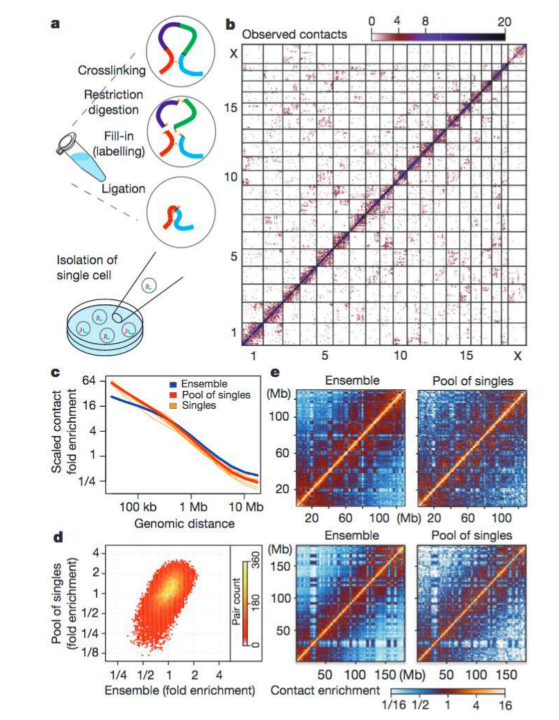
Hi-C at single cell level

ARTICLE

doi:10.1038/nature12593

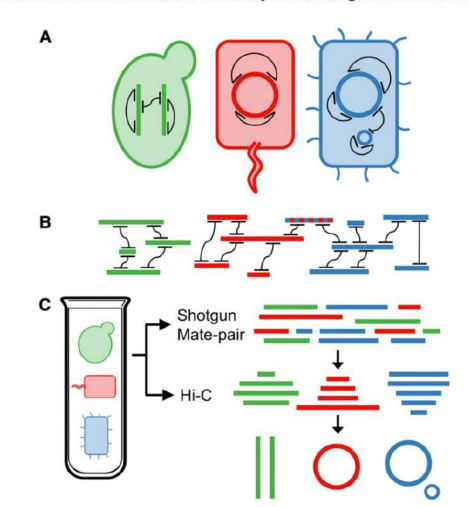
Single-cell Hi-C reveals cell-to-cell variability in chromosome structure

Takashi Nagano¹*, Yaniv Lubling²*, Tim J. Stevens³*, Stefan Schoenfelder¹, Eitan Yaffe², Wendy Dean⁴, Ernest D. Laue³, Amos Tanay² & Peter Fraser¹



Species-Level Deconvolution of Metagenome Assemblies with Hi-C–Based Contact Probability Maps

Joshua N. Burton,¹ Ivan Liachko,¹ Maitreya J. Dunham,² and Jay Shendure² Department of Genome Sciences, University of Washington, Seattle, Washington 98195-5065



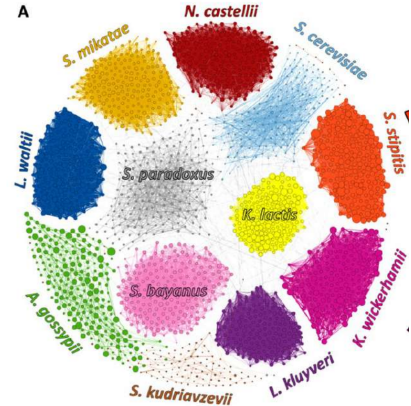
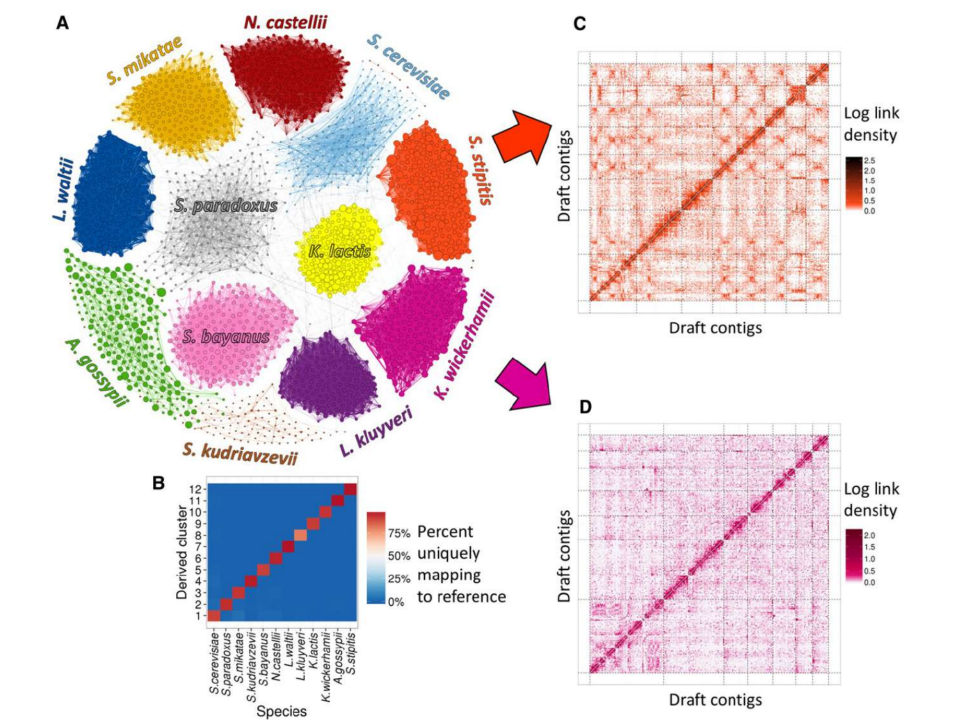


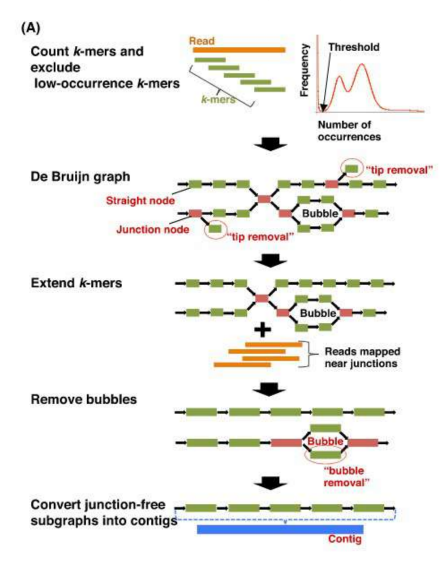
Table 2	Sequencing	libraries	used in	MetaPhase	analyses
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Sample	Library Type	Read Length, bp	Read Pairs, millions
M-Y	Shotgun	101	85.7
	Mate-pair	100	9.2
	Hi-C	100	81.0



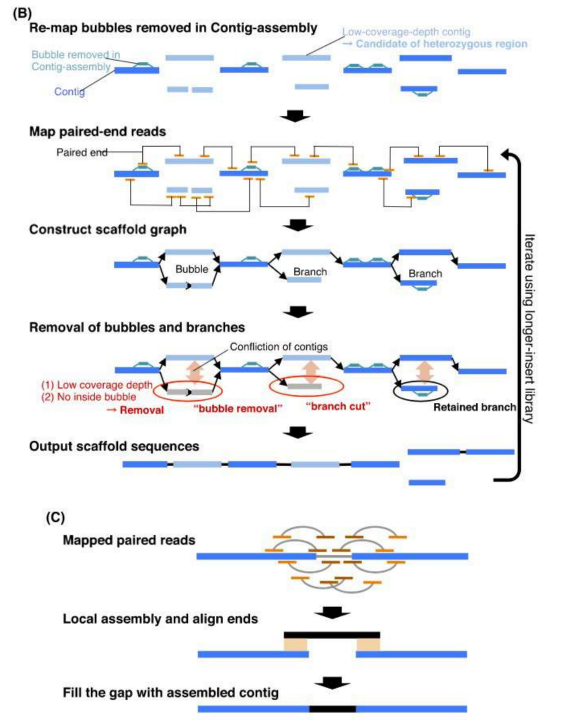
Case studies

Would you understand everything in this paper?



Resource-

Efficient de novo assembly of highly heterozygous genomes from whole-genome shotgun short reads



Why sequence a genome? (2021 version)

- Genomics advance our understanding of organisms across tree of life
 - All previous published genomes that were fragmented are being redone again
 - To reveal greater insights and ease of use for community
 - New genomes are expected to be of good quality
- Reveal more variations within species
 - Population genomics is not just **remapping** anymore
 - More accurate inference of structure variation
 - Gene level
- Better analysis power in a genomics world

Lower quality genomes are getting improved

LETTER

OPEN doi:10.1038/nature22971

Improved maize reference genome with single-molecule technologies

Yinping Jiao¹, Paul Peluso², Jinghua Shi³, Tiffany Liang³, Michelle C. Stitzer⁴, Bo Wang¹, Michael S. Campbell¹, Joshua C. Stein¹, Xuehong Wei¹, Chen–Shan Chin², Katherine Guill⁵, Michael Regulski¹, Sunita Kumari¹, Andrew Olson¹, Jonathan Gent⁶, Kevin L. Schneider⁷, Thomas K. Wolfgruber⁷, Michael R. May⁸, Nathan M. Springer⁹, Eric Antoniou¹, W. Richard McCombie¹, Gernot G. Presting⁷, Michael McMullen⁵, Jeffrey Ross–Ibarra¹⁰, R. Kelly Dawe⁶, Alex Hastie³, David R. Rank² & Doreen Ware^{1,11}

RESEARCH ARTICLE

Open Access

CrossMark

An improved genome assembly uncovers prolific tandem repeats in Atlantic cod

Ole K. Tørresen^{1*}, Bastiaan Star¹, Sissel Jentoft^{1,2}, William B. Reinar¹, Harald Grove³, Jason R. Miller⁴, Brian P. Walenz⁵, James Knight⁶, Jenny M. Ekholm⁷, Paul Peluso⁷, Rolf B. Edvardsen⁸, Ave Tooming-Klunderud¹, Morten Skage¹, Sigbjørn Lien³, Kjetill S. Jakobsen¹ and Alexander J. Nederbragt^{1,9*}

An improved assembly of the loblolly pine

mega-genome using long-read single-molecule sequencing

Aleksey V. Zimin^{1,2}, Kristian A. Stevens³, Marc W. Crepeau³, Daniela Puiu², Jill L. Wegrzyn⁴, James A. Yorke¹, Charles H. Langley³, David B. Neale⁵ and Steven L. Salzberg^{2,6,*}

Improved genome assembly of American alligator genome reveals conserved architecture of estrogen signaling

Edward S. Rice,¹ Satomi Kohno,² John St. John,³ Son Pham,⁴ Jonathan Howard,⁵ Liana F. Lareau,⁶ Brendan L. O'Connell,^{1,7} Glenn Hickey,¹ Joel Armstrong,¹ Alden Deran,¹ Ian Fiddes,¹ Roy N. Platt II,⁸ Cathy Gresham,⁹ Fiona McCarthy,¹⁰ Colin Kern,¹¹ David Haan,¹ Tan Phan,¹² Carl Schmidt,¹³ Jeremy R. Sanford,¹⁴ David A. Ray,⁸ Benedict Paten,¹⁵ Louis J. Guillette Jr.,^{16,†} and Richard E. Green^{1,6,7}

Single-molecule sequencing and optical mapping yields an improved genome of woodland strawberry (Fragaria vesca) with chromosome-scale contiguity

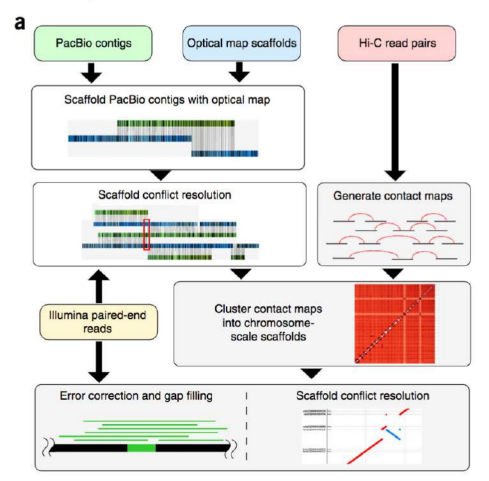
Patrick P. Edger^{1,2,*,†}, Robert VanBuren^{1,†}, Marivi Colle¹, Thomas J. Poorten³, Ching Man Wai¹, Chad E. Niederhuth⁴, Elizabeth I. Alger¹, Shujun Ou^{1,2}, Charlotte B. Acharya³, Jie Wang⁵, Pete Callow¹, Michael R. McKain⁶, Jinghua Shi⁷, Chad Collier⁷, Zhiyong Xiong⁸, Jeffrey P. Mower⁹, Janet P. Slovin¹⁰, Timo Hytönen¹¹, Ning Jiang^{1,2}, Kevin L. Childs^{5,12} and Steven J. Knapp^{3,*}

An improved assembly and annotation of the allohexaploid wheat genome identifies complete families of agronomic genes and provides genomic evidence for chromosomal translocations

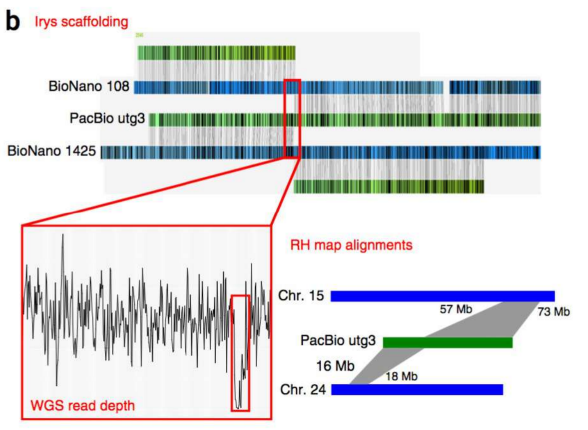
Bernardo J. Clavijo,^{1,9} Luca Venturini,^{1,9} Christian Schudoma,¹ Gonzalo Garcia Accinelli,¹ Gemy Kaithakottil,¹ Jonathan Wright,¹ Philippa Borrill,² George Kettleborough,¹ Darren Heavens,¹ Helen Chapman,¹ James Lipscombe,¹ Tom Barker,¹ Fu-Hao Lu,² Neil McKenzie,² Dina Raats,¹ Ricardo H. Ramirez-Gonzalez,^{1,2} Aurore Coince,¹ Ned Peel,¹ Lawrence Percival-Alwyn,¹ Owen Duncan,³ Josua Trösch,³ Guotai Yu,² Dan M. Bolser,⁴ Guy Namaati,⁴ Arnaud Kerhornou,⁴ Manuel Spannagl,⁵ Heidrun Gundlach,⁵ Georg Haberer,⁵ Robert P. Davey,^{1,6} Christine Fosker,¹ Federica Di Palma,^{1,6} Andrew L. Phillips,⁷ A. Harvey Millar,³ Paul J. Kersey,⁴ Cristobal Uauy,² Ksenia V. Krasileva,^{1,6,8} David Swarbreck,^{1,6} Michael W. Bevan,² and Matthew D. Clark^{1,6}

Single-molecule sequencing and chromatin conformation capture enable *de novo* reference assembly of the domestic goat genome

Derek M Bickhart^{1,18}, Benjamin D Rosen^{2,18}, Sergey Koren^{3,18}, Brian L Sayre⁴, Alex R Hastie⁵, Saki Chan⁵, Joyce Lee⁵, Ernest T Lam⁵, Ivan Liachko⁶, Shawn T Sullivan⁷, Joshua N Burton⁶, Heather J Huson⁸, John C Nystrom⁸, Christy M Kelley⁹, Jana L Hutchison², Yang Zhou^{2,10}, Jiajie Sun¹¹, Alessandra Crisà¹², F Abel Ponce de León¹³, John C Schwartz¹⁴, John A Hammond¹⁴, Geoffrey C Waldbieser¹⁵, Steven G Schroeder², George E Liu², Maitreya J Dunham⁶, Jay Shendure^{6,16}, Tad S Sonstegard¹⁷, Adam M Phillippy³, Curtis P Van Tassell² & Timothy P L Smith⁹



New genomes are expected to have high quality



Correcting misassemblies

... ARS1 comprises just 31 scaffolds and 649 gaps covering 30 of the 31 haploid, acrocentric goat chromosomes (excluding only the Y chromosome),

Table 1 Assembly statistics

4	o i b	0 (())	Unplaced	Degenerate	Contig	Scaffold	Assembly	Assembly in
Assembly ^a	Contigs ^b	Scaffolds	contigsc	contigs ^d	NG50 (Mb) ^e	NG50 (Mb) ^{e,f}	size (Gb)	scaffolds (%)
PacBio	3,074	-		30,693	3.795		2.914	N/A
Optical Map	-	2,944	-	-	_	1.487	2.748	N/A
PacBio + Optical Map	1,109	333	1,242	30,693	10.197	20.623	2.910	90.89
PacBio + Hi-C	2,115	31	959	30,693	3.795	88.799	2.910	87.97
PacBio + Optical Map + Hi-C	1,780	31	571	30,693	10.197	87.347	2.910	89.05
ARS1	680	31	654	29,315	18.702	87.277	2.924	88.32

^aAssemblies are listed in order of inclusion of scaffolding technologies toward the final assembly (ARS1), with the original contigs (PacBio) scaffolded using different technologies (Optical Map and Hi-C). Because the optical map program (Irys Scaffold) generates an assembly from the consensus of labeled DNA molecules, we have included scaffold statistics from these data (optical map) for comparison. ^bThe number of continuous stretches of sequence within the scaffold without gaps >3 bases in length of at least 10 bases. ^cUnplaced contigs are defined as input contigs or scaffolds that were not placed by the optical map or Hi-C in a scaffold were excluded from the scaffold counts. ^dDegenerate contigs were assembled unitigs that had less than 50 PacBio reads supporting their assembly (**Supplementary Note**). Differences in degenerate contig counts in the final ARS1 assembly are due to PBJelly merging of degenerate contigs (538 contigs) or removal due to no supporting PacBio read alignments (840). ^eAll NG50 values are based on the ARS1 assembly size (2.924 Gb). ^fNo scaffolds were generated for the PacBio entry.

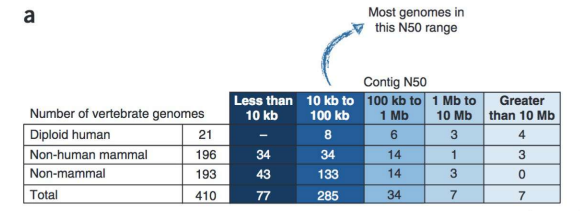
A golden goat genome

Kim C Worley

The newly described *de novo* goat genome sequence is the most contiguous diploid vertebrate assembly generated thus far using whole-genome assembly and scaffolding methods. The contiguity of this assembly is approaching that of the finished human and mouse genomes and suggests an affordable roadmap to high-quality references for thousands of species.

.... This report generates sighs of relief from researchers frustrated with the highly fragmented genome sequences available for most species....

... The lower costs and greater accessibility of these methods bring potential for wider impact....



b

	Goat CHIR_1.0	Goat ARS1	Human GRCh38
Total sequence length	2.6 Gb	2.9 Gb	3.2 Gb
Total assembly gap length	140 Mb	38 Mb	160 Mb
Gaps between scaffolds	411	0	349
Number of scaffolds	77,431	29,907	735
Scaffold N50	14 Mb	87 Mb	67 Mb
Number of contigs	337,494	30,399	1,385
Contig N50	18.9 kb	26.2 Mb	56.4 Mb
Number of chromosomes and plasmids	30	31	25



С

Genomes with

highest N50

Difficult regions are being targeted

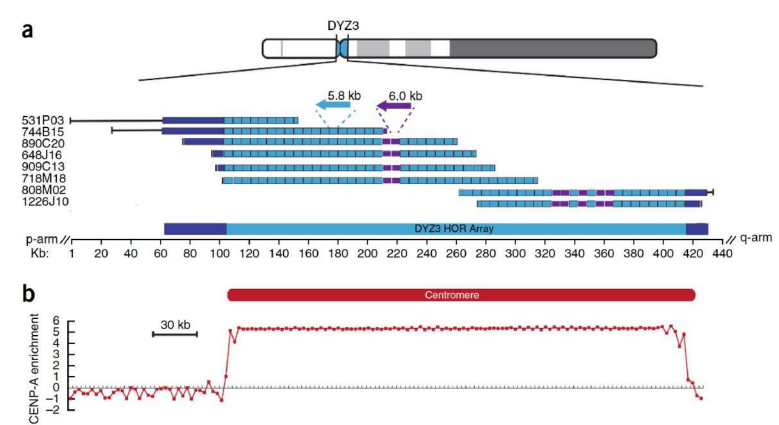
nature biotechnology

OPEN

Linear assembly of a human centromere on the Y chromosome

Miten Jain^{1,5}, Hugh E Olsen^{1,5}, Daniel J Turner², David Stoddart², Kira V Bulazel³, Benedict Paten¹, David Haussler¹, Huntington F Willard^{3,4}, Mark Akeson¹ & Karen H Miga^{1,3}

The human genome reference sequence remains incomplete owing to the challenge of assembling long tracts of nearidentical tandem repeats in centromeres. We implemented a nanopore sequencing strategy to generate high-quality reads that span hundreds of kilobases of highly repetitive DNA in a human Y chromosome centromere. Combining these data with short-read variant validation, we assembled and characterized the centromeric region of a human Y chromosome.



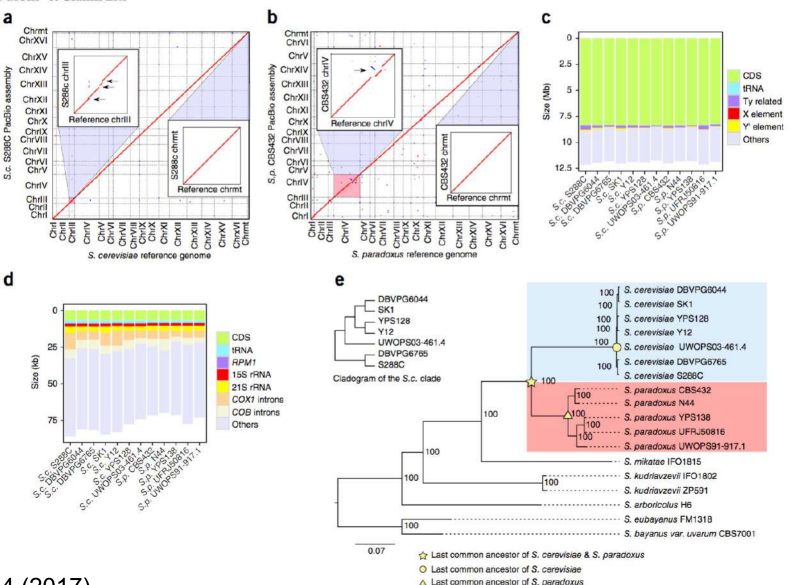
Jain *et al* (2018) Nature Biotechnology

Contrasting evolutionary genome dynamics between domesticated and wild yeasts

Population genomics

Jia-Xing Yue¹, Jing Li¹, Louise Aigrain², Johan Hallin¹, Karl Persson³, Karen Oliver², Anders Bergström², Paul Coupland^{2,5}, Jonas Warringer³, Marco Cosentino Lagomarsino⁴, Gilles Fischer⁴, Richard Durbin² & Gianni Liti¹

 long-read sequencing to generate end-to-end genome assemblies for
 12 strains representing major subpopulations of the partially domesticated yeast Saccharomyces cerevisiae and its wild relative S. paradoxus.



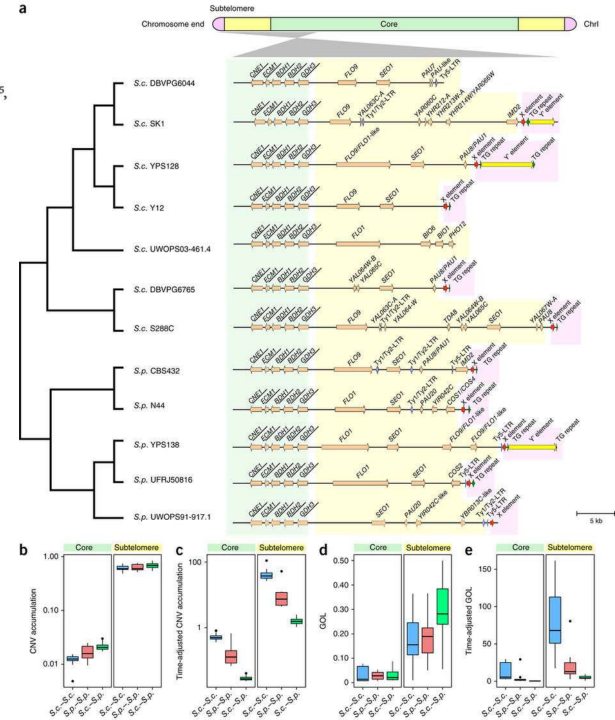
Nature Genetics volume 49, pages 913–924 (2017)

Contrasting evolutionary genome dynamics between domesticated and wild yeasts

Jia-Xing Yue¹, Jing Li¹, Louise Aigrain², Johan Hallin¹, Karl Persson³, Karen Oliver², Anders Bergström², Paul Coupland^{2,5}, Jonas Warringer³, Marco Cosentino Lagomarsino⁴, Gilles Fischer⁴, Richard Durbin² & Gianni Liti¹

- enable precise definition of chromosomal boundaries between cores and subtelomeres
- S. paradoxus shows faster accumulation of balanced rearrangements (inversions, reciprocal translocations and transpositions), S. cerevisiae accumulates unbalanced rearrangements (novel insertions, deletions and duplications) more rapidly.
- Such striking contrasts between wild and domesticated yeasts are likely to reflect the influence of human activities on structural genome evolution.

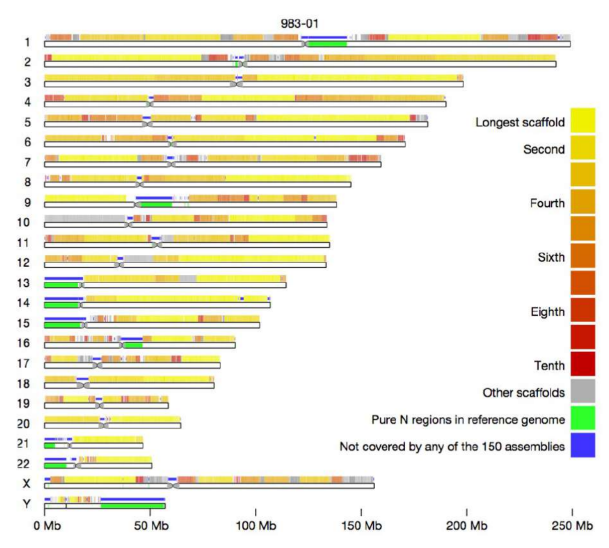
Nature Genetics volume 49, pages 913–924 (2017)



Sequencing and *de novo* assembly of 150 genomes from Denmark as a population reference

Lasse Maretty^{1*}, Jacob Malte Jensen^{2,3*}, Bent Petersen^{4*}, Jonas Andreas Sibbesen^{1*}, Siyang Liu^{1,5*}, Palle Villesen^{2,3,6*}, Laurits Skov^{2,3*}, Kirstine Belling^{4*}, Christian Theil Have⁷, Jose M. G. Izarzugaza⁴, Marie Grosjean⁴, Jette Bork-Jensen⁷, Jakob Grove^{3,8,9}, Thomas D. Als^{3,8,9}, Shujia Huang^{10,11}, Yuqi Chang¹⁰, Ruiqi Xu⁵, Weijian Ye⁵, Junhua Rao⁵, Xiaosen Guo^{10,12}, Jihua Sun^{5,7}, Hongzhi Cao¹⁰, Chen Ye¹⁰, Johan van Beusekom⁴, Thomas Espeseth^{13,14}, Esben Flindt¹², Rune M. Friborg^{2,3}, Anders E. Halager^{2,3}, Stephanie Le Hellard^{14,15}, Christina M. Hultman¹⁶, Francesco Lescai^{3,8,9}, Shengting Li^{3,8,9}, Ole Lund⁴, Peter Løngren⁴, Thomas Mailund^{2,3}, Maria Luisa Matey–Hernandez⁴, Ole Mors^{3,6,9}, Christian N. S. Pedersen^{2,3}, Thomas Sicheritz–Pontén⁴, Patrick Sullivan^{16,17}, Ali Syed⁴, David Westergaard⁴, Rachita Yadav⁴, Ning Li⁵, Xun Xu¹⁰, Torben Hansen⁷, Anders Krogh¹, Lars Bolund^{8,10}, Thorkild I. A. Sørensen^{7,18,19}, Oluf Pedersen⁷, Ramneek Gupta⁴, Simon Rasmussen⁴§, Søren Besenbacher^{2,6}§, Anders D. Børglum^{3,8,9}§, Jun Wang^{3,10,12}§, Hans Eiberg²⁰§, Karsten Kristiansen^{10,12}§, Søren Brunak^{4,21}§ & Mikkel Heide Schierup^{2,3,22}§

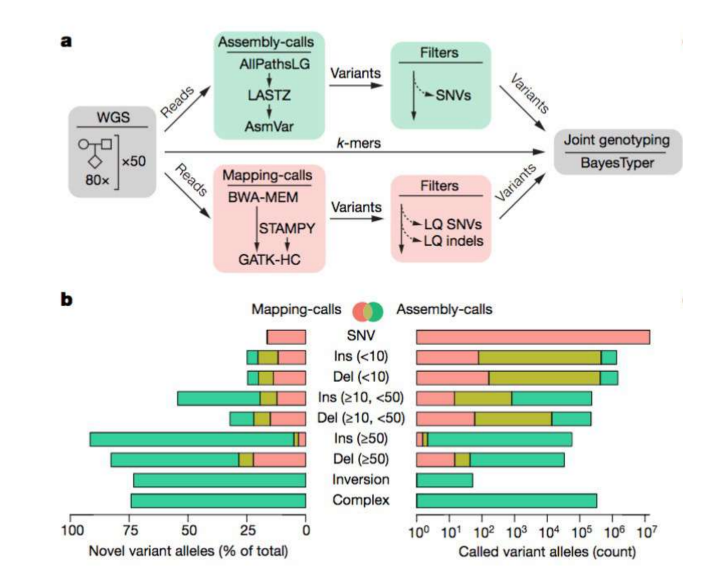
- Hundreds of thousands of human genomes are now being sequenced to characterize genetic variation and use this information to augment association mapping studies of complex disorders and other phenotypic traits.
- Genetic variation is identified mainly by mapping short reads to the reference genome. However, these approaches are biased against discovery of structural variants and variation in the more complex parts of the genome.
- report *de novo* assemblies of 150 individuals (50 trios) from the GenomeDenmark project.



We found that 16.4% of the called SNVs were novel (not in the Single Nucleotide Polymorphism database 142 (dbSNP142) or 1000 Genomes Project phase 3 structural variant call-set), whereas as many as 91.6% of insertions ≥50 bp were novel (Fig. 2b).

The fraction of novel variants increased rapidly with variant length, especially for insertions (Fig. 2d), with most longer variants contributed by the assembly-based approach...

For instance, …we called 33,653 deletions ≥50 bp, whereas the 1000 Genomes Project identified 42,279 such variants in 25 times more individuals who were more diverse than our study population.



ARTICLE

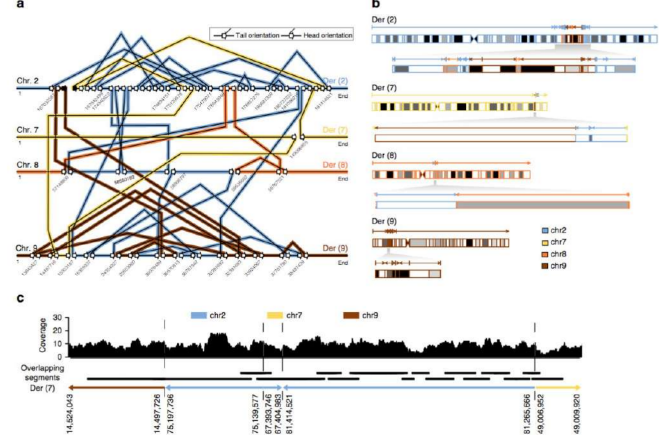
DOI: 10.1038/s41467-017-01343-4

OPEN

Mapping and phasing of structural variation in patient genomes using nanopore sequencing

Mircea Cretu Stancu¹, Markus J. van Roosmalen¹, Ivo Renkens¹, Marleen M. Nieboer¹, Sjors Middelkamp¹, Joep de Ligt ¹, Giulia Pregno², Daniela Giachino ², Giorgia Mandrile², Jose Espejo Valle-Inclan¹, Jerome Korzelius¹, Ewart de Bruijn¹, Edwin Cuppen³, Michael E. Talkowski^{4,5,6}, Tobias Marschall ^{7,8}, Jeroen de Ridder¹ & Wigard P. Kloosterman¹

- long reads are superior to short reads with regard to detection of de novo chromothripsis rearrangements.
- long reads also enable efficient phasing of genetic variations, which we leveraged to determine the parental origin of all de novo chromothripsis breakpoints and to resolve the structure of these complex rearrangements.



Long reads lead to better SV

Example of metagenomics

Metagenomic Discovery of Biomass-Degrading Genes and Genomes from Cow Rumen

Matthias Hess,^{1,2}* Alexander Sczyrba,^{1,2}* Rob Egan,^{1,2} Tae-Wan Kim,³ Harshal Chokhawala,³ Gary Schroth,⁴ Shujun Luo,⁴ Douglas S. Clark,^{3,5} Feng Chen,^{1,2} Tao Zhang,^{1,2} Roderick I. Mackie,⁶ Len A. Pennacchio,^{1,2} Susannah G. Tringe,^{1,2} Axel Visel,^{1,2} Tanja Woyke,^{1,2} Zhong Wang,^{1,2} Edward M. Rubin^{1,2}†

- 268Gb of metagenomics data
- Identified 27,755 putative carbohydrateactive genes from a cow rumen metagenome
- Expressed 90 candidates of which 57% had enzymatic activity against cellulosic substrates
- Assembled 15 uncultured microbial genomes

Hess et al., 2011 Science

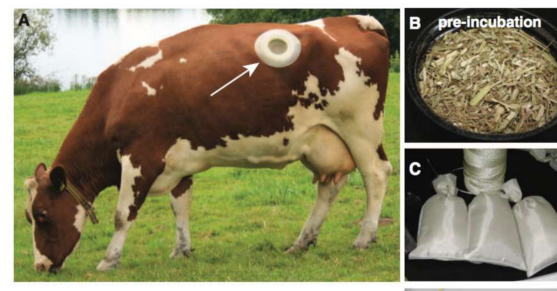


Fig. 1. (**A**) A surgically created fistula (arrow) sealed with a flexible cannula was used to study the degradation of switchgrass within the rumen. (**B**) Switchgrass before rumen incubation. (**C**) Nylon bags filled with switchgrass before insertion into the rumen. (**D**) Switchgrass after 72 hours of rumen incubation.



Genome Bin	Genome Size (Mb)	Phylogenetic Order	Estimated Complete- ness
AFa	2.87	Spirochaetales	92.98%
AMa	2.21	Spirochaetales	91.23%
Ala	2.53	Clostridiales	90.10%
AGa	3.08	Bacteroidales	89.77%
AN	2.02	Clostridiales	78.50%
AJ	2.24	Bacteroidales	75.96%
AC2a	2.07	Bacteroidales	75.96%
AWa	2.02	Clostridiales	75.77%
AH	2.52	Bacteroidales	75.45%
AQ	1.91	Bacteroidales	71.36%
ASta	1.75	Clostridiales	70.99%
APb	2.41	Clostridiales	64.85%
BOa	1.67	Clostridiales	64.16%
ADa	2.99	Myxococcales	62.13%
ATa	1.87	Clostridiales	60.41%

enome; innermost circle (green tick marks), location of glycoside hydrolase genes on draft genome.

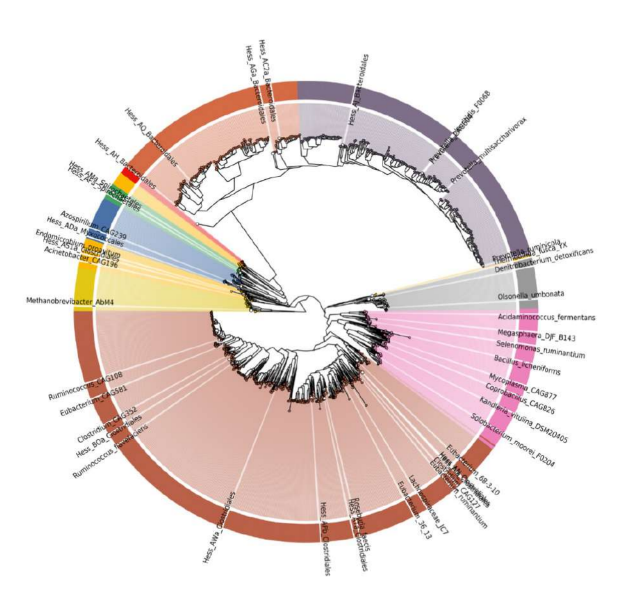
ARTICLE

DOI: 10.1038/s41467-018-03317-6

Assembly of 913 microbial genomes from metagenomic sequencing of the cow rumen

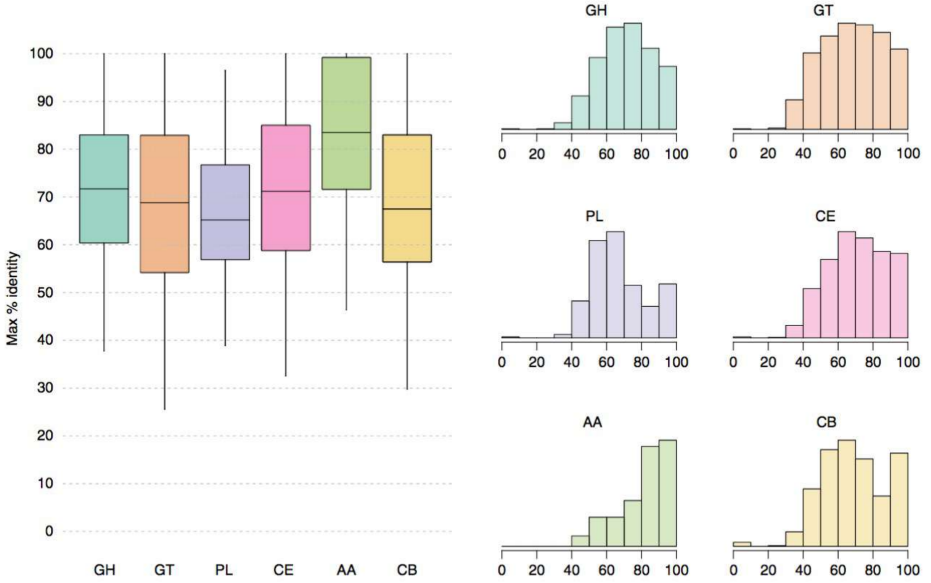
Robert D. Stewart¹, Marc D. Auffret², Amanda Warr¹, Andrew H. Wiser³, Maximilian O. Press³, Kyle W. Langford³, Ivan Liachko³, Timothy J. Snelling⁴, Richard J. Dewhurst¹, Alan W. Walker⁴, Rainer Roehe² & Mick Watson¹

- 800 Gb of sequence data derived from 43 Scottish cattle
- Using metagenomic binning and Hi-C techniques
- Assembly of 913 draft bacterial and archaeal genomes
- Most of these genomes represent previously unsequenced strains and species.
- The draft genomes contain over 69,000 proteins predicted to be involved in carbohydrate metabolism, over 90% of which do not have a good match in public databases.



Stewart et al (2018) Nature Communications

 The draft genomes contain over 69,000 proteins predicted to be involved in carbohydrate metabolism, over 90% of which do not have a good match in public databases.



- Inclusion of the 913 genomes presented here improves metagenomic read classification by sevenfold against the study's own data, and by fivefold against other publicly available rumen datasets.
- dataset substantially improves the coverage of rumen microbial genomes in the public databases and represents a valuable resource for biomassdegrading enzyme discovery and studies of the rumen microbiome

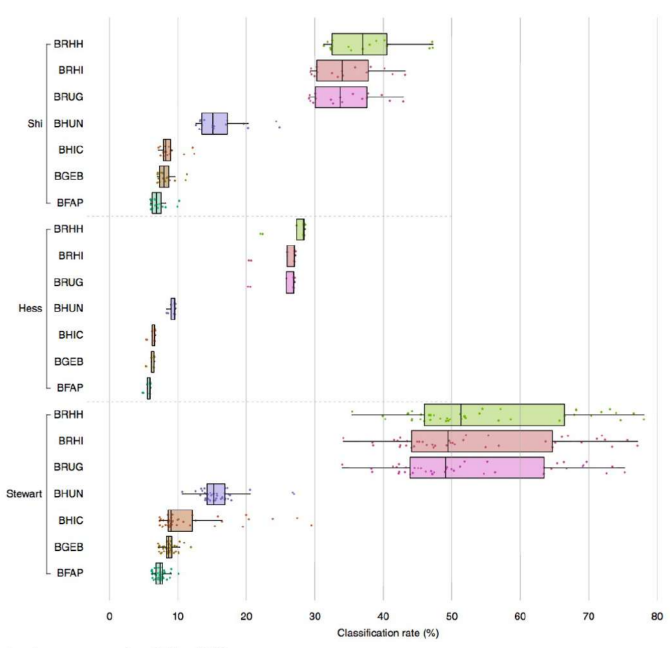


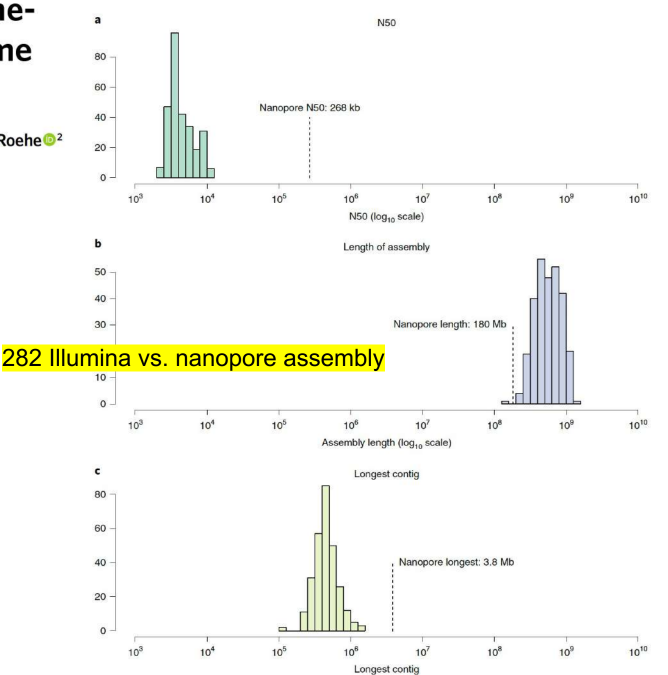
Fig. 4 Classification rate for three datasets against various Kraken databases. BFAP bacterial, archaeal, fungal and protozoan genomes from RefSeq, BGEB BFAP + 1003 GEBA genomes, BHIC BFAP + 63 hRUG genomes, BHUN BFAP + 410 genomes from the Hungate 1000 project, BRUG BFAP + 850 RUG MAGs, BRHI BFAP + all 913 genomes from this study, BRHH BFAP + 913 RUGs + 410 Hungate 1000 genomes. Addition of rumen-specific RUGs or Hungate 1000 genomes has the most dramatic effect

Compendium of 4,941 rumen metagenomeassembled genomes for rumen microbiome biology and enzyme discovery

Robert D. Stewart¹, Marc D. Auffret¹, Amanda Warr¹, Alan W. Walker¹, Rainer Roehe¹ and Mick Watson¹

- **6.5 Tb** of sequence data derived from **283** ruminant cattles
- Using metagenomic binning and Hi-C techniques
- Assembly of 4,941 draft bacterial and archaeal genomes
- Long read is being used: "We also present a metagenomic assembly of nanopore (MinION) sequencing data (from one rumen sample) that contains at least three whole bacterial chromosomes as single contigs"

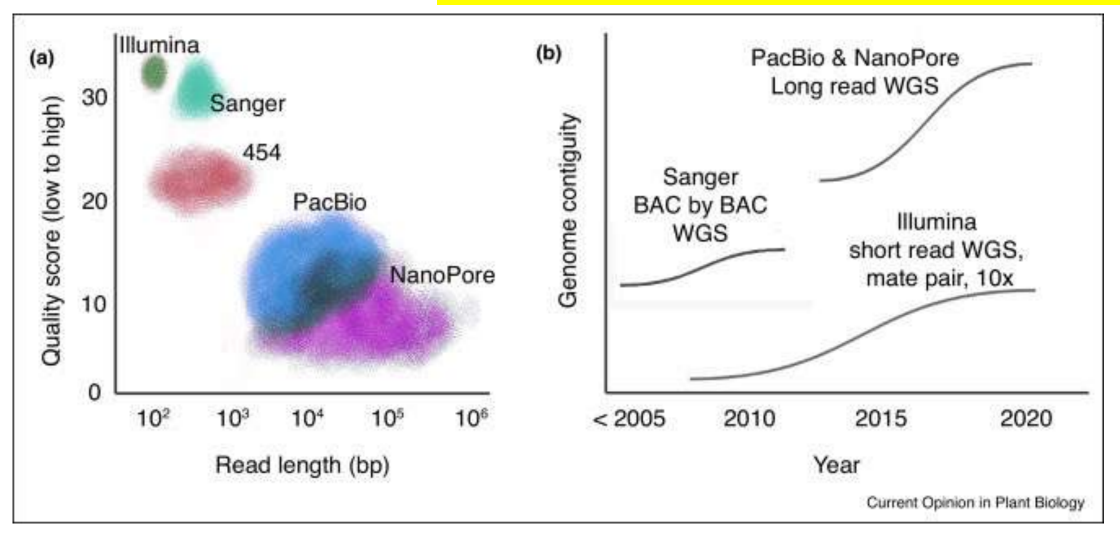
Stewart et al (2019) Nature Biotechnology



Summary and opinions

Summary and opinions

Advances in sequencing technology have dramatically improved genome contiguity over the last two decades

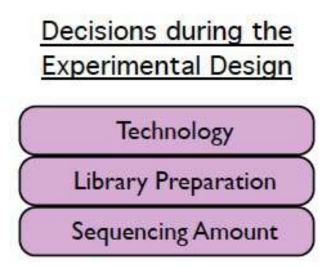


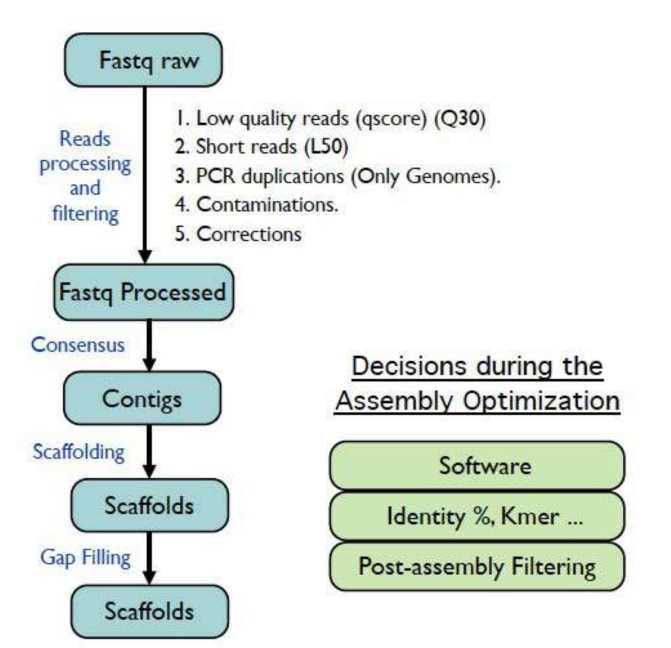
Todd and VanBuren (2020) Current Opinion in Plant Biology

Summary

- Expectation is higher in assembly of a genome
- Population genomics is moving from a mapping (resequencing) to assembly approach
- Long read technologies are improving rapidly fast so every standard lab can generate a high quality assembly
- Assembly processes need to scale up to accommodate the advancing technologies and changing biological questions

Overview of a sequencing project





Credit: Aureliano Bombarely

Things to consider

- \$\$\$\$\$\$\$
- Project type

virus, bacteria, eukaryote, meta-genome

• Goals

Just an assembly to showcase the world?

Sequence pandemic species = conservation? (No right or wrong answer) Any biological question?

Why *de novo* sequence a species?

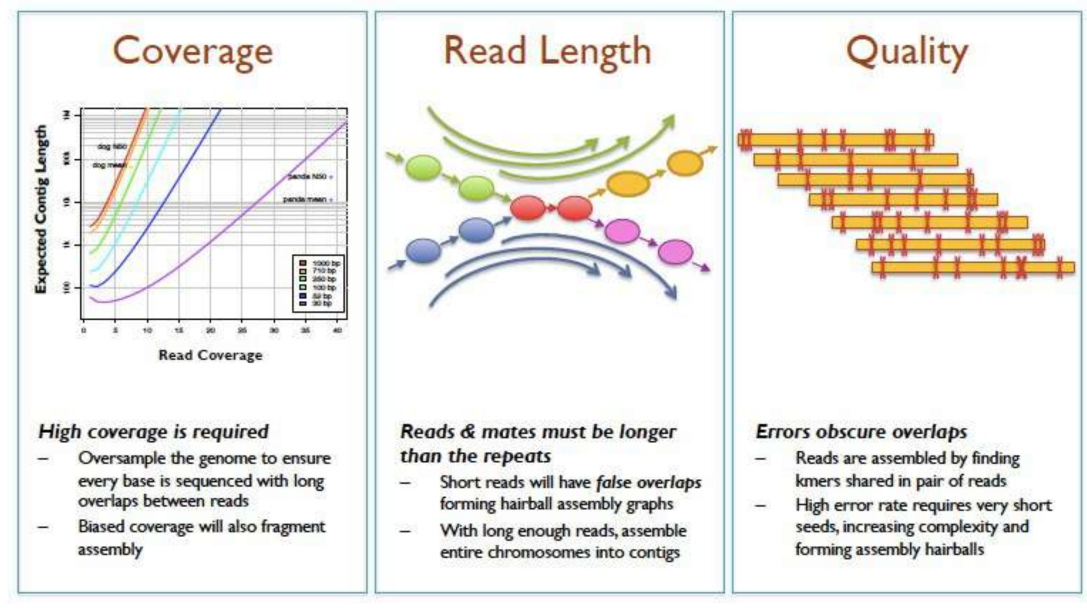
• Hardware

- You need CPUs, but RAM is more important
 - Imagine storing all the hashes or kmers
- This may change depending on nature of data (all long reads within n years?)

Consider technologies and experiments

- Use multiple techniques to answer your questions
 - Long read only
 - Long read + Hi-C
 - Long read + optical maps + Hi-C
 - 10X linked reads
 - 10X + Hi-C
 - Single cell?
 - Experimental advancement?
- Sometimes limitations becomes experiment preparation rather than the technological one

Ingredients for a good assembly



Credit: Michael Schatz

FUTURE ISSUES

- 1. High-error reads: Third-generation DNA sequencing technologies (e.g., from Pacific Biosciences and Oxford Nanopore) generate much longer reads than previously possible (tens of thousands of base pairs), but with the cost of much higher error rates.
- 2. Metagenomics/mixtures of organisms: Increasingly, scientists are sequencing the genomes within mixtures of organisms, whether in the context of metagenomics or within clinical samples (e.g., mixtures of tumor cells). Most of the theoretical framework for sequence assembly was developed for isolated genomes. New methods will need to be developed that can both cope with and characterize heterogeneity within closely related genomes.
- 3. Dealing with big data: As DNA sequencing costs drop, scientists are increasingly able to focus on larger genomes (such as those of plants) and mixtures (such as soil metagenomes). New approaches will be needed that allow genome sequence assemblers to scale with the amount of data being generated.

Not covered but should be

Alignment method in overlap graph String graph (reduced overlap graph) Shortest Superstring Problem (SSP) Hamiltonian path

Choice of kmers in DBG Bloom filter

References

shorturl.at/xBC06

Assignment

- 1. Choose a group of species, or a species.
- 2. Please write a short review (~10 references) on how analyses of comparative/population genomics have been transformed by recent advances (algorithm and experimental approaches) in sequencing.