

Module 10: Genome assembly

Eugene Gardner

Postdoctoral Fellow, Wellcome Sanger Institute

e : eg15@sanger.ac.uk

 : [@DrGeneUK](https://twitter.com/DrGeneUK)

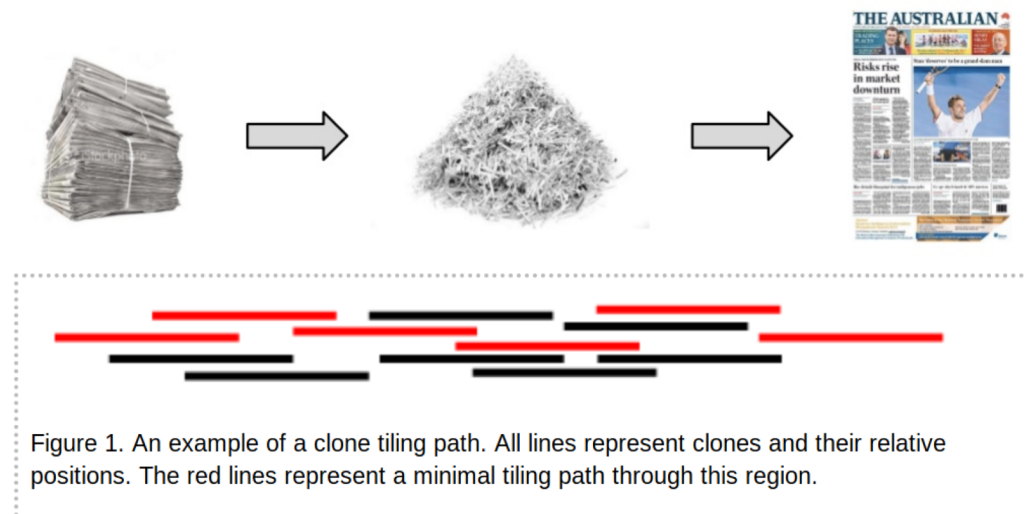
Presentation originally developed by:

Shane McCarthy

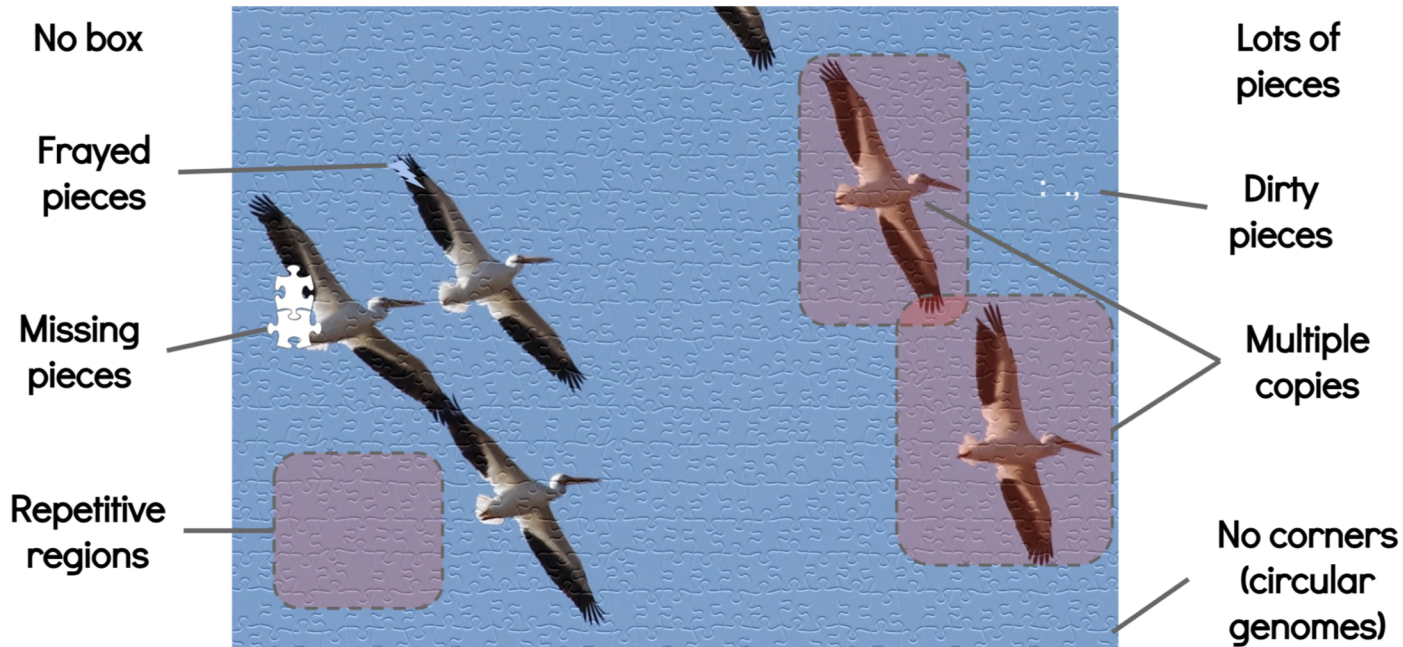
What is genome assembly?

The process of reconstructing the genomic sequence from sequenced fragments

- Sequencing technologies can only sequence short stretches of DNA (100 bp-1M bp)
- Given many (millions or billions) of reads, produce a linear (or perhaps circular) genome
- Clone based
 - Select clones using markers
 - Sequence clones separately
- Whole-genome shotgun
 - Fragment whole-genome into tiny pieces and sequence



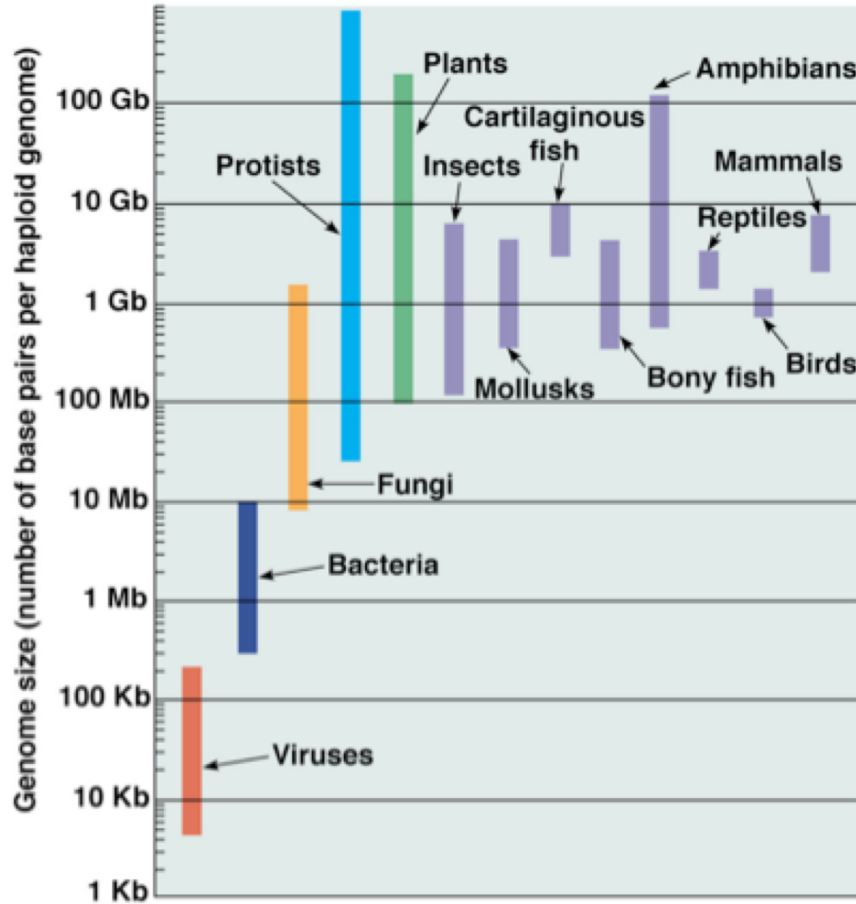
What makes assembling a jigsaw puzzle hard?



- **What helps?** Larger pieces (longer reads); fewer dirty or frayed pieces (more accurate reads); fewer repeats and copies (simpler genomes)

Key considerations in assembly

- genome size
- genome heterozygosity and ploidy
- sequencing coverage
- errors in reads
- read lengths
- non-uniqueness of the genome (repeat content)
- running time and memory of assembly strategy



Terminology

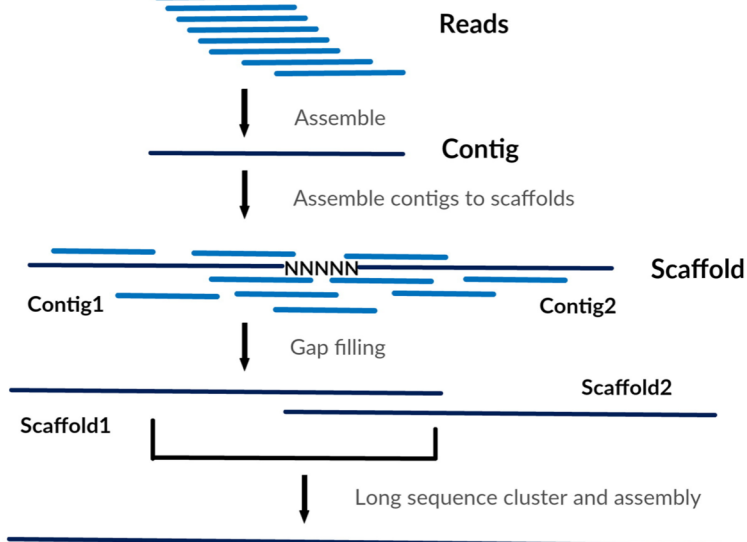
Puzzle pieces



Partial Puzzle



Full Puzzle



Chromosome Improvements and impacts of GRCh38 human reference on high throughput sequencing data analysis

Yan Guo ^{a,*}, Yulin Dai ^a, Hui Yu ^a, Shilin Zhao ^a, David C. Samuels ^b, Yu Shyr ^{c,*}

- **contig:** contiguous stretch of assembled sequence
- **scaffold:** ordered set of contigs with gaps (Ns) placed between

a) Multiple copies of genome



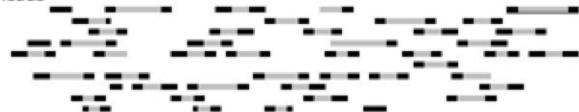
b) Sheared random fragments



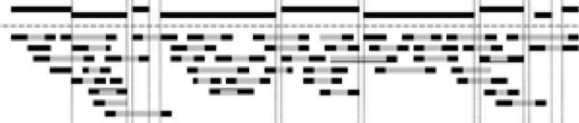
c) Size fractionated fragments



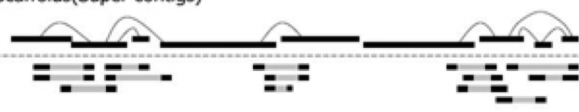
d) Reads



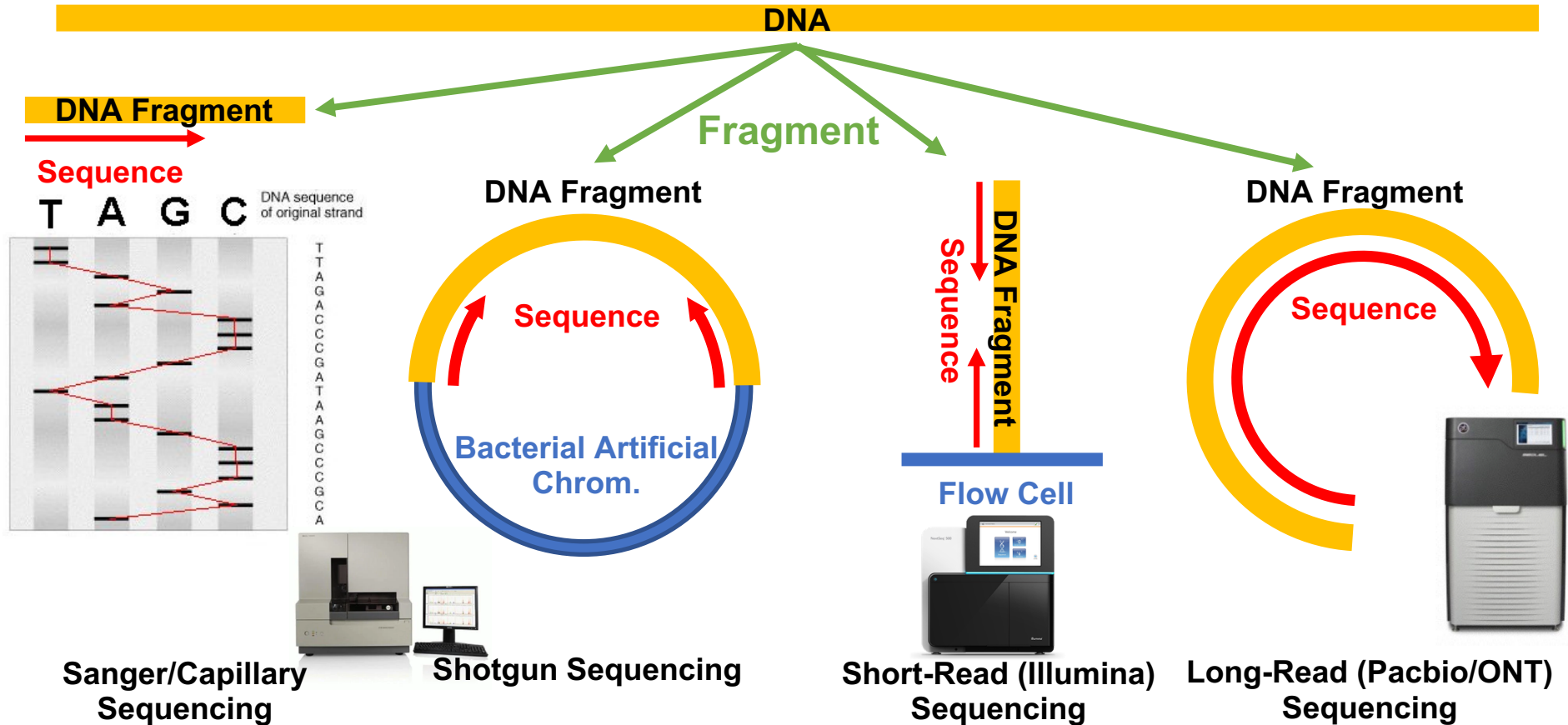
e) Contigs



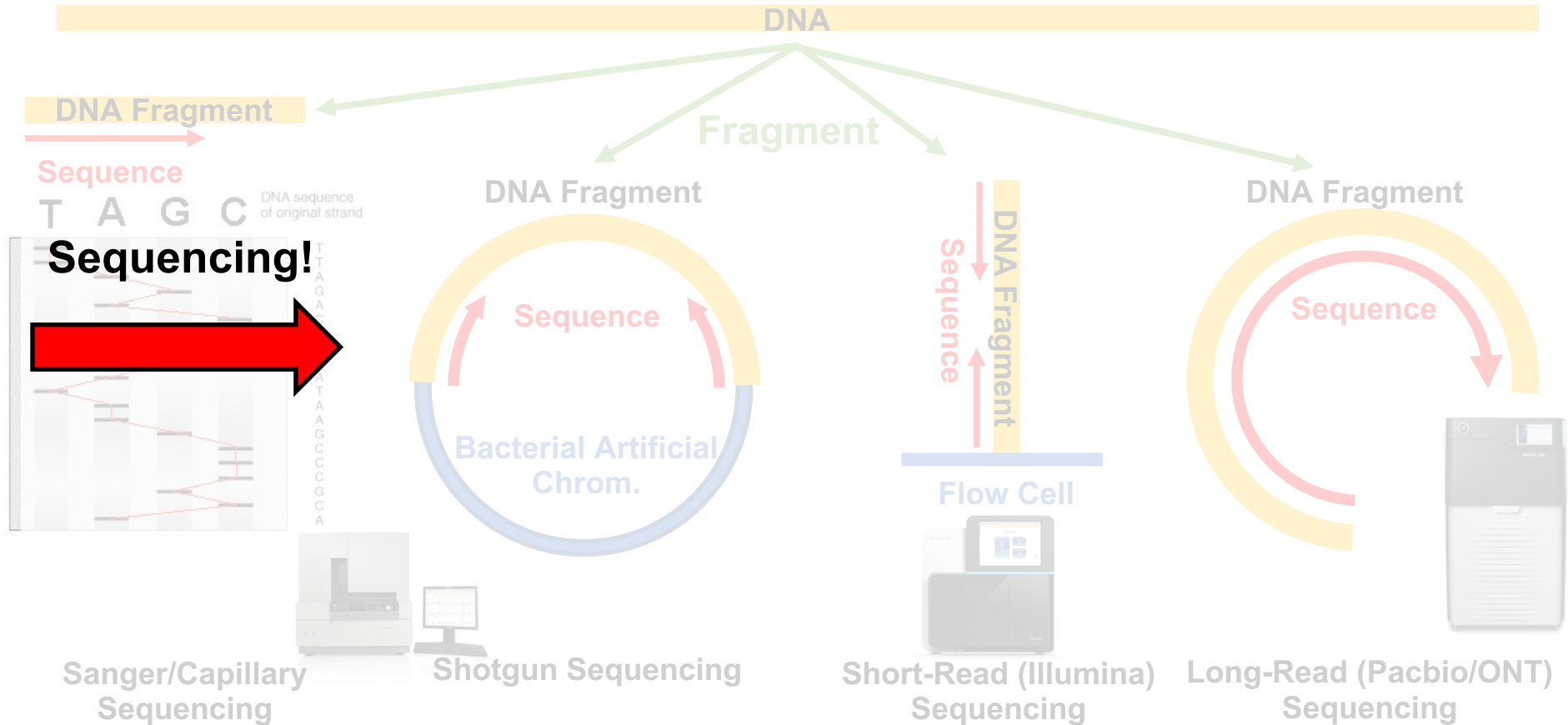
f) Scaffolds(Super contigs)



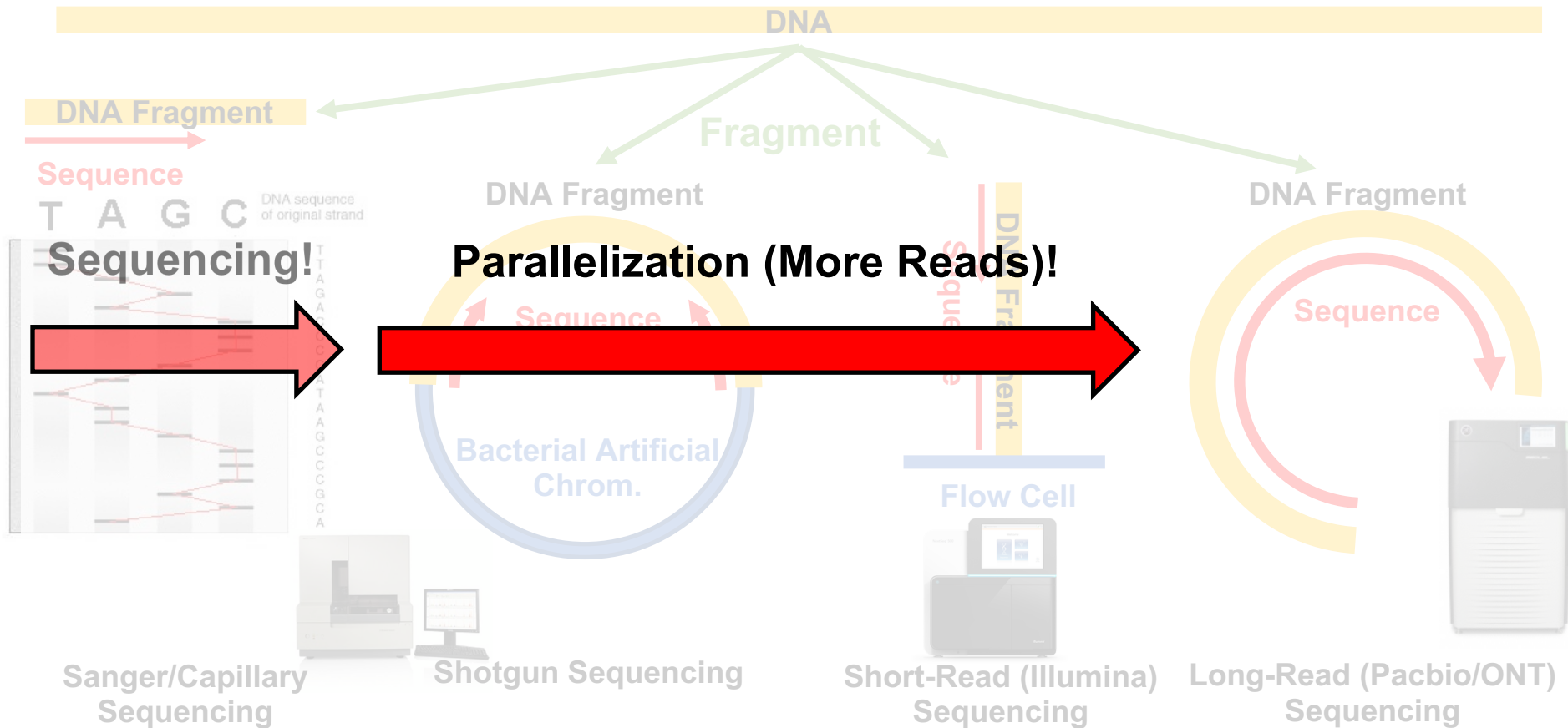
Brief History of Genome Assembly Technology



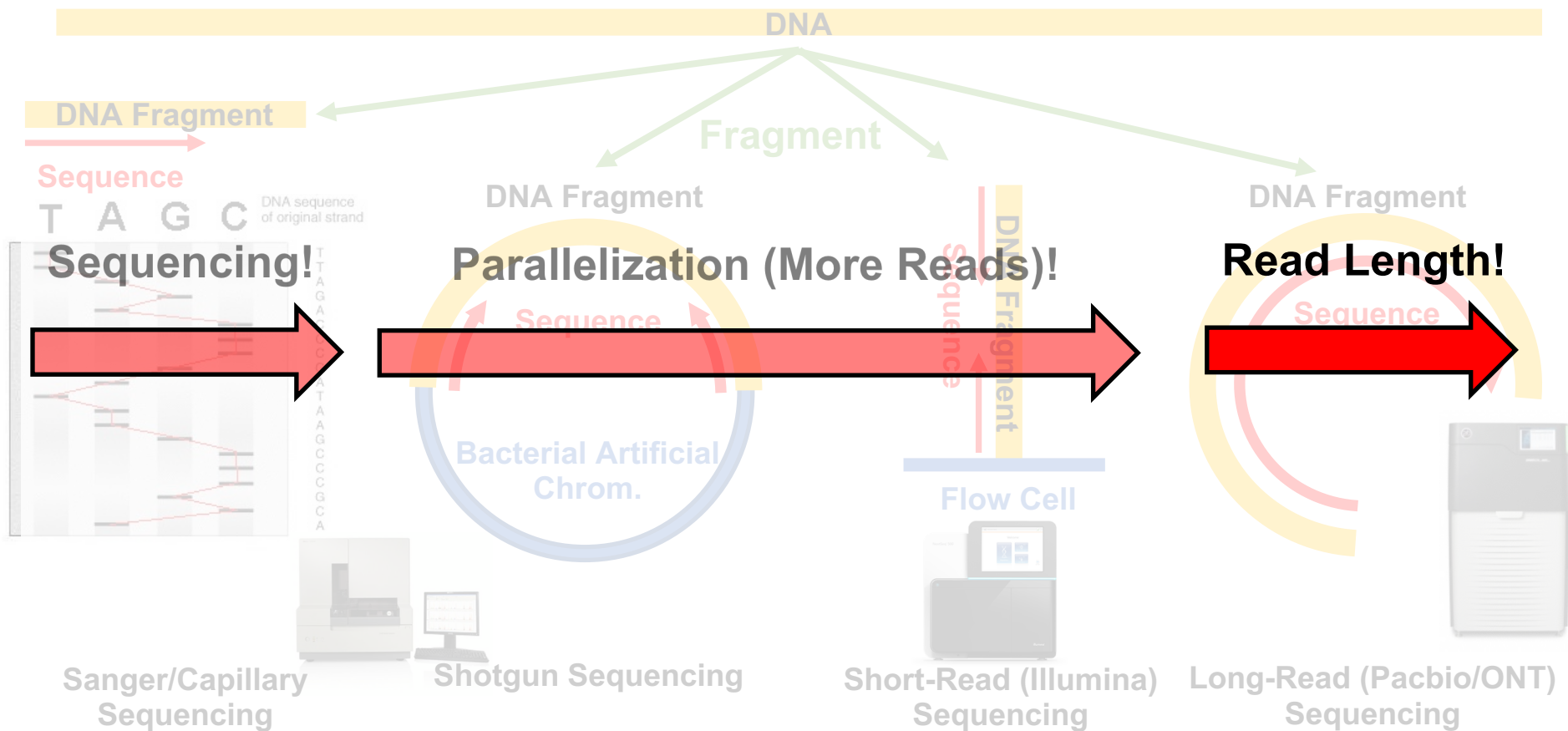
Brief History of Genome Assembly Technology



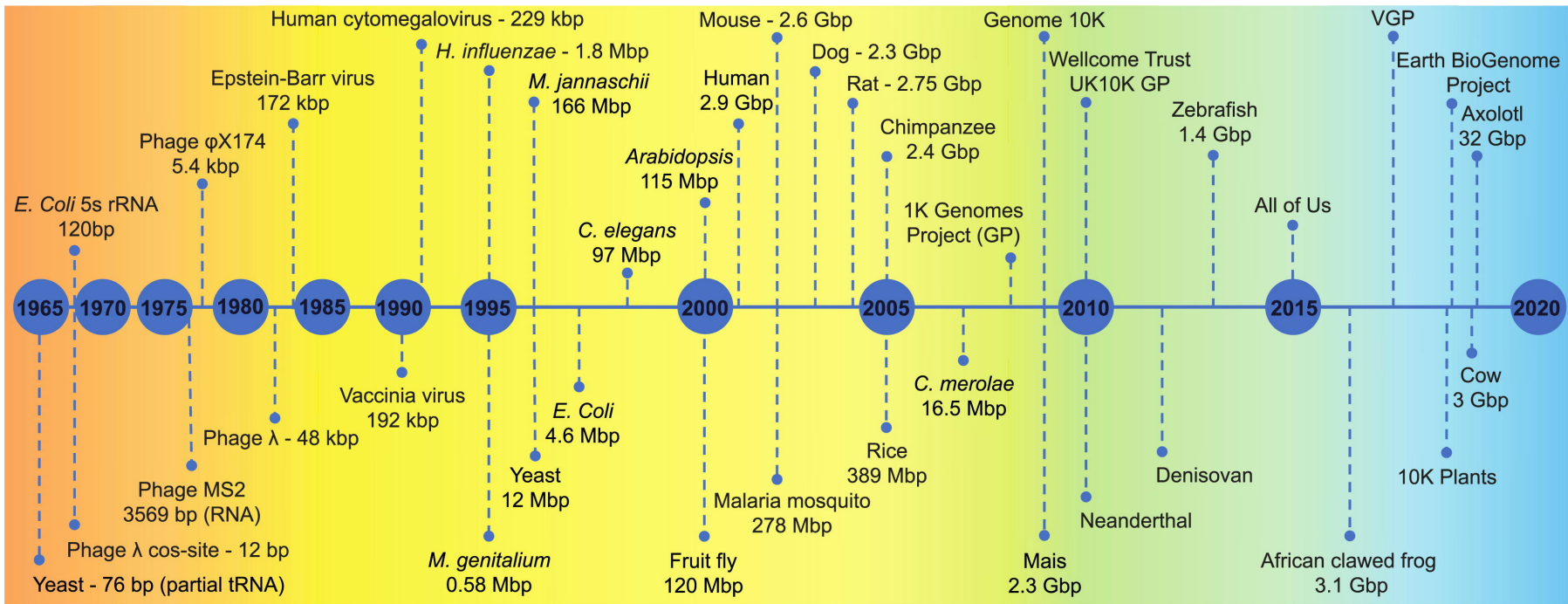
Brief History of Genome Assembly Technology



Brief History of Genome Assembly Technology



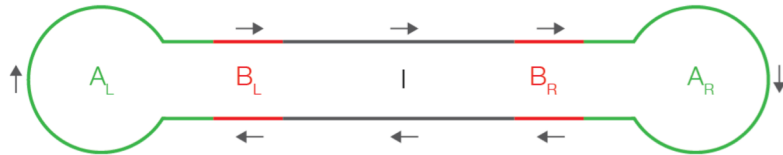
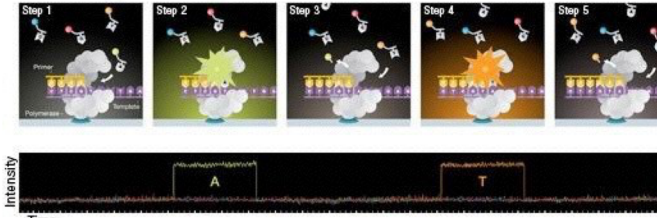
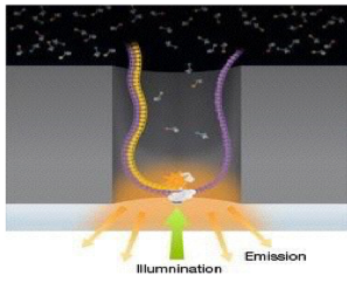
Brief History of Genome Assembly



Key considerations in assembly – Technology

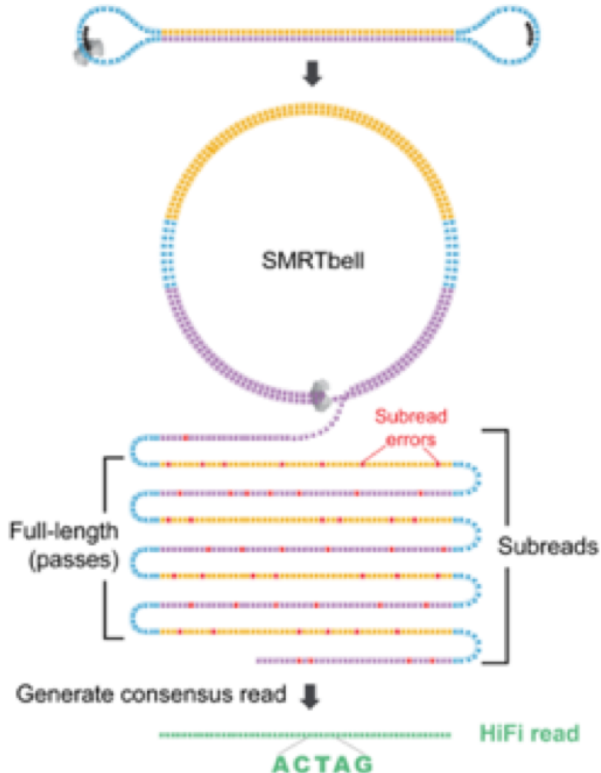
- Long read sequencing has become the dominant platform for genome assembly
- Illumina reads alone are rarely used now - while there used to be a large tradeoff in length/error rate of reads, improvements in tech and algorithms have made long reads common.

PacBio sequencing



Key points:

- 1 DNA molecule and 1 polymerase in each well (zero-mode waveguide, ZMW)
- 4 colours flash in realtime as polymerase acts
- Methylated bases have distinct pattern
- No theoretical limit to DNA fragment length



PacBio sequencing and assembly

Advantages:

- Long read lengths
- Few systematic errors
- Can detect some base modifications
- Can get more accuracy, if you loop over the ZMW multiple times - circular consensus sequencing (CCS)

Disadvantages:

- High read error rate
- High cost per-base

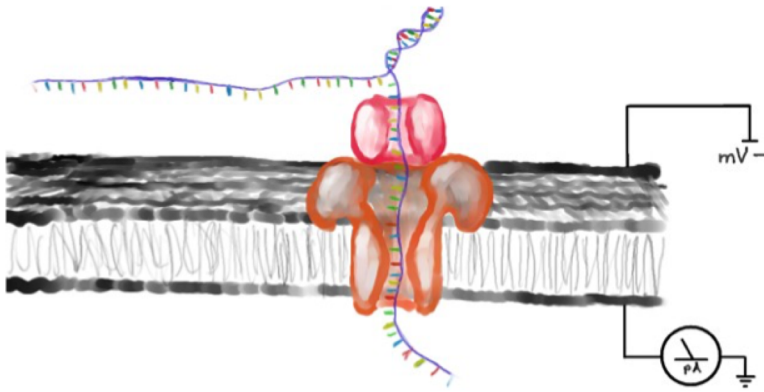
2019 Update:

- Sequel II launched with 8-fold increase in yield and improvement in CCS accuracy (HiFi reads).

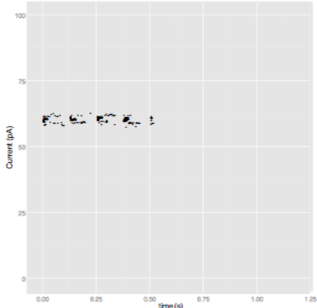


Pacbio Sequel
>10kbp read length
Up to 10GB yield

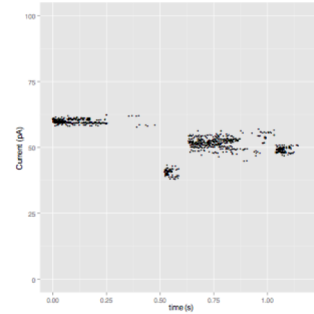
Nanopore sequencing



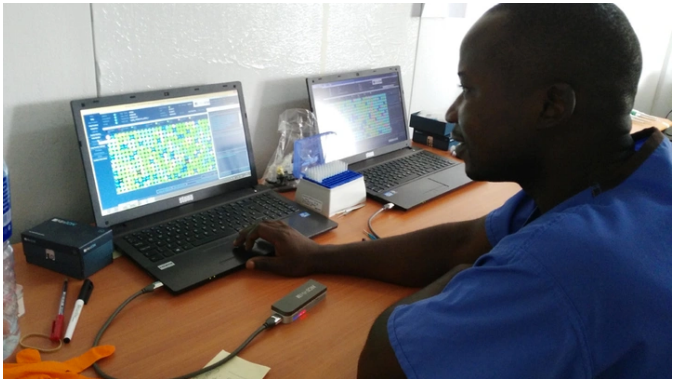
GCTACGATT →
Sample Current



CTACGATT →
Sample Current



Nanopore sequencing and assembly



Ed Yong. *The Atlantic*. September 16, 2015.

Advantages:

- Portability and low capital cost
- Read lengths up to 1Mbp reported
- Base modification detection
- Sequence RNA directly



Oxford Nanopore MinION

>10kbp read length
5-15GB yield



Oxford Nanopore PromethION

>10kbp read length
30-90GB yield per flowcell

Disadvantages:

- High error rate
- Systematic errors around homopolymers

Read clouds or linked-reads

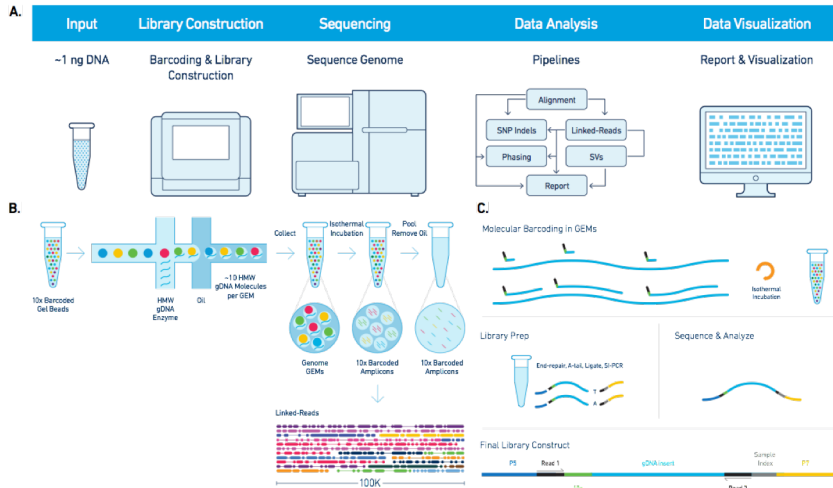
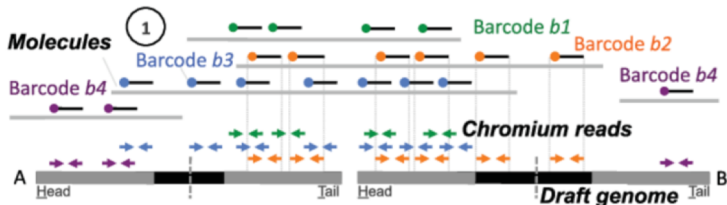


Figure 1. Chromium™ Genome Solution. (a) The Chromium Genome Solution provides a streamlined workflow and easy-to-use informatics pipeline and software for whole genome analysis. (b) An automated microfluidic system allows for functionalized gel beads to be combined with high molecular weight DNA (HMW gDNA) and oil to form a "Gel Bead in Emulsion (GEM)". Each GEM contains ~10 molecules of HMW gDNA and primers with unique barcodes. Isothermal incubation allows for the addition of a unique barcode to all DNA within the GEM. (c) Assay schematic overview.



Longer range “read clouds” e.g. from 10X Genomics Chromium, can bridge bigger repeats/gaps

Multiple reads from a long ~100kb template sharing a barcode



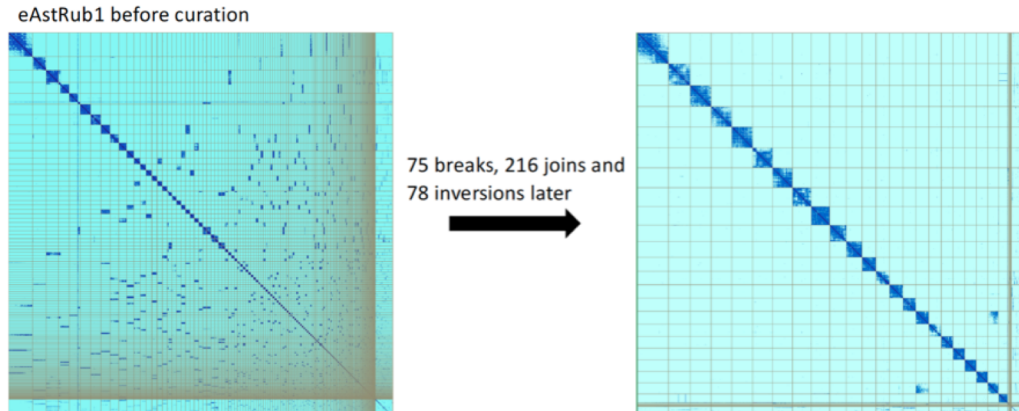
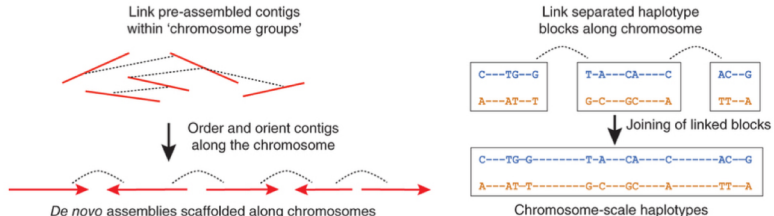
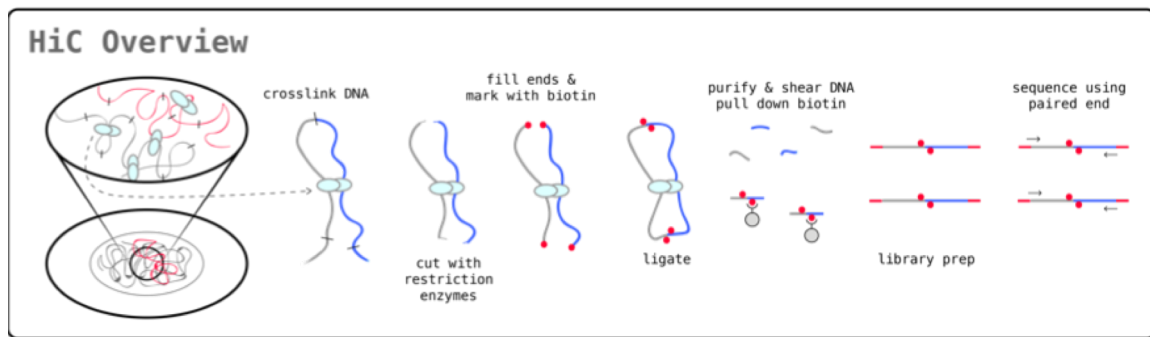
Yeo S, Coombe L, Warren RL, Chu J, Birol I. ARCS: scaffolding genome drafts with linked reads. Bioinformatics. 2018.

Hi-C

Hi-C “proximity ligation”

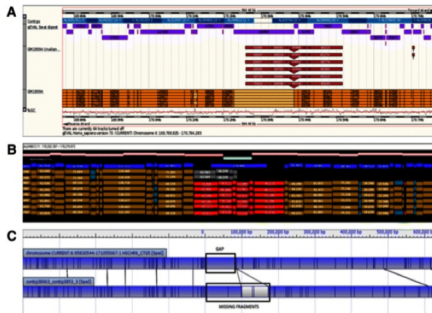
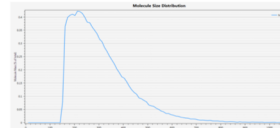
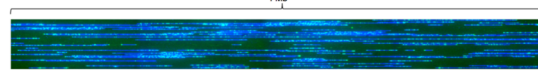
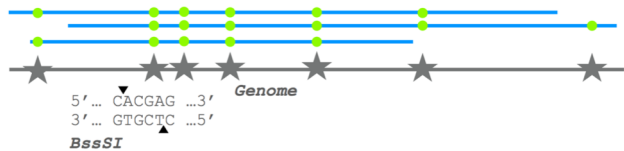
is used to look at local chromatin organisation, but globally most Hi-C links are on the same chromosome, and more are close than distant

- Automated tools (e.g. SALSA2, 3d-dna)
- Visualisers (e.g. Juicer, Hi-Glass)
- Editors (e.g. Juicer, Pretext)



Optical mapping

- Isolate high molecular weight DNA fragments
- Label specific sequence motifs across the entire genome
- Linearise and image labelled DNA
- Restriction digest maps give approximate spacing of restriction sites (short sequence motifs), skipping over repeats
- Now scaled up by dedicated optical imaging machines, e.g. BioNano



bionano
GENOMICS

Long-range technologies

Table 1 | Long-range sequencing and mapping platforms

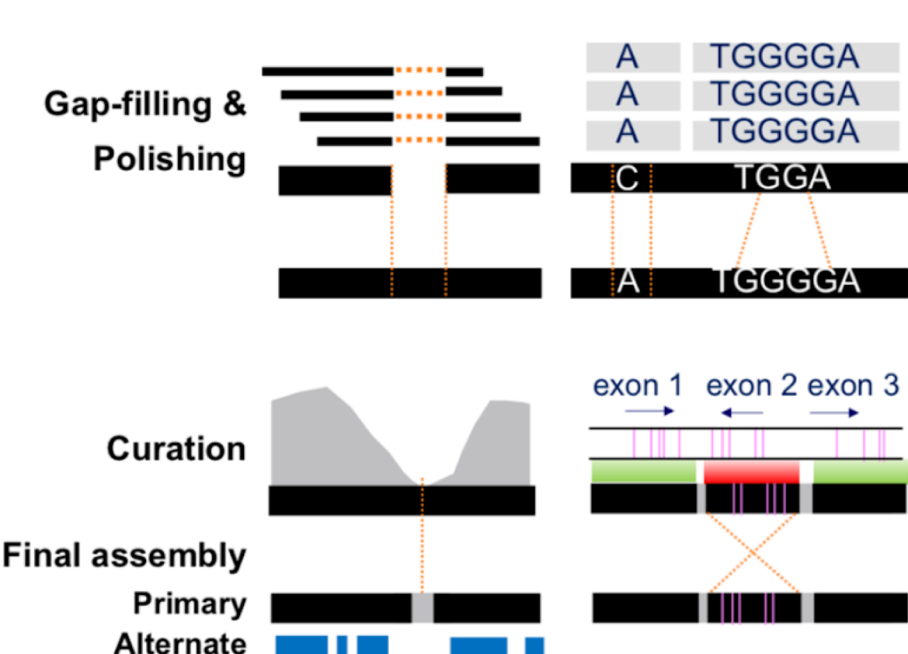
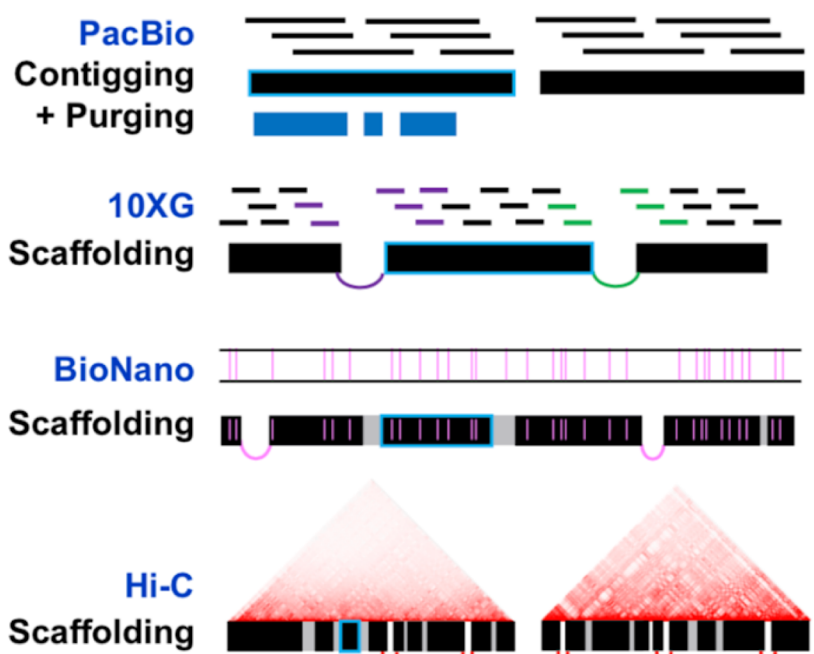
Platform	General characteristics and costs	Major applications	Bioinformatics challenges
PacBio SMRT sequencing	Single-molecule long reads averaging ~10 kb with some approaching 100 kb; several fold more expensive than short reads	De novo genome assembly, structural variant detection, gene isoform resolution and epigenetic modifications	Raw reads have high error rates dominated by false insertions; requires new alignment and error correction algorithms
Oxford Nanopore sequencing	Single-molecule long reads averaging ~10 kb with some >1 Mb; several fold more expensive than short reads	De novo genome assembly, structural variant detection, gene isoform resolution and epigenetic modifications	Raw reads have high error rates dominated by false deletions and homopolymer errors; requires new alignment and error correction algorithms
10X Genomics Chromium	Linked reads spanning ~100 kb derived from a collection of short-read sequences; moderately more expensive than short reads	De novo genome assembly and scaffolding, phasing, detection of large structural variants (>10 kb) and single-cell gene expression	Sparse sequencing rather than true long reads; more complicated to align, with poorer resolution of locally repetitive sequences
Hi-C-based analysis	Pairs of short reads (<100 bp) formed from crosslinking chromatin interactions; moderately more expensive than short reads	Genome scaffolding and phasing	Sparse sequencing with highly variable genomic distance between pairs (1 kb to 1 Mb or longer)
BioNano Genomics optical mapping	Optical mapping of long DNA molecules (~250 kb or longer) labelled with fluorescent probes; less expensive than short reads	Genome scaffolding and detection of large structural variants (>10 kb)	Limited algorithms to discover high-confidence alignment between an optical map and a sequence assembly

PacBio SMRT, Pacific Biosciences single-molecule real time.

Piercing the dark matter:
bioinformatics of long-range
sequencing and mapping

Fritz J. Sedlazeck¹, Hayan Lee², Charlotte A. Darby³ and Michael C. Schatz^{3,4*}

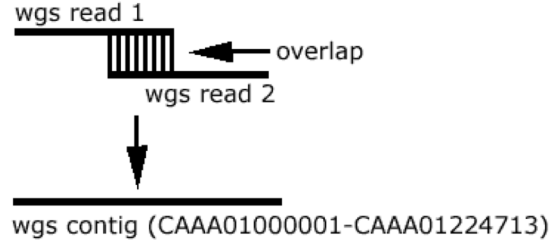
Example genome project workflow



Assembly: contig generation

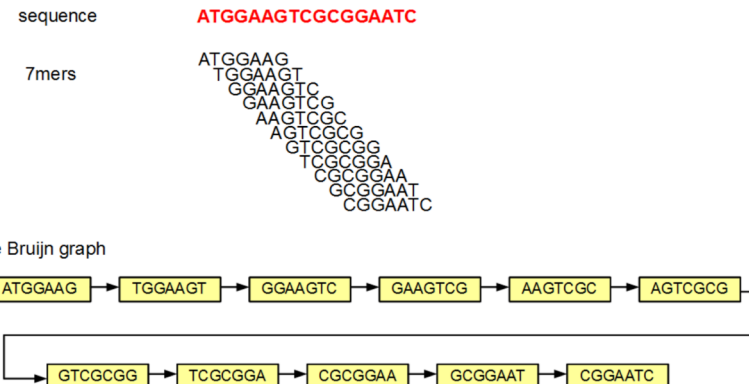
OLC (Overlap Layout Consensus):

- For all pairs x, y , of reads
 - determine if there is sufficient overlap
 - bundle stretches of overlap graph into contigs
- Computationally expensive (quadratic scaling with current approaches)
- Assembly software
 - Falcon (PacBio), Canu (PacBio, ONT), minimap/miniasm



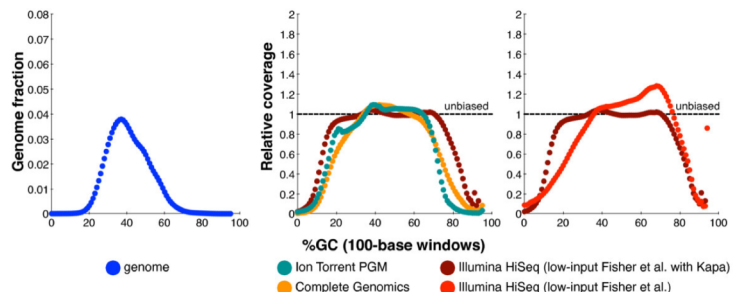
DBG: de Bruijn graph (k -mer):

- Build a graph of all subsequences ("kmers") of length k .
- Assembly software
 - velvet, ABySS, SPAdes, wtdbg2

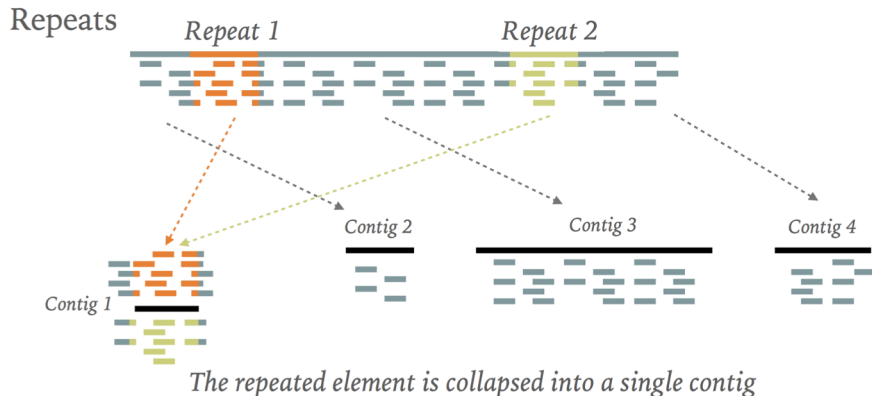


Hurdles

- Repeats / low complexity genomic regions
 - Heterozygosity
 - Sequencing errors
 - Base composition and sequencing bias



Ross, 2013



The repeated element is collapsed into a single contig

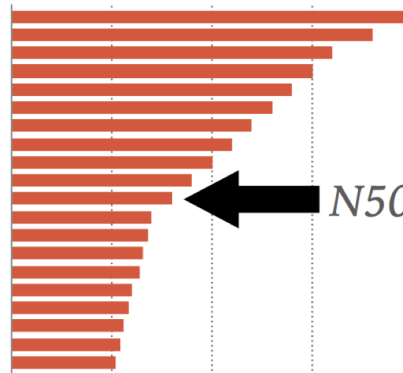
Adapted from Torsten Seemann (2014 talk)



Assembly metrics

- total length
- number of sequences (contigs and scaffolds)
- average length (contigs and scaffolds)
- largest/smallest (contigs and scaffolds)
- $N50 = X$ means 50% of the genome is in sequences larger than X
- NG50 (N50 scaled by the expected genome size)
- Gene content (% conserved core genes mapped)

$N50 = \text{what is the smallest contig at 50\% of genome?}$



Scaffolding

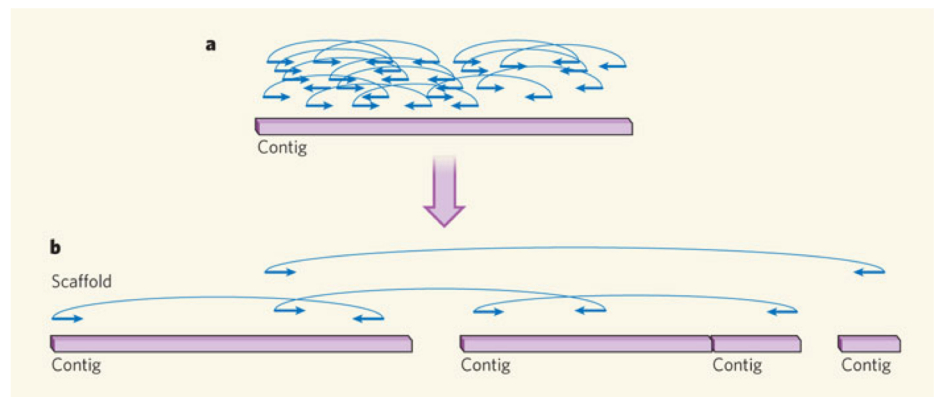
Goal: Order and orient the contigs into larger structure

Construct libraries of varying insert sizes

- Smaller size (2, 4 or 6 Kb), intermediate size (10-40 Kb), and libraries with large insert sequences (>100 Kb)
- Ends of these clones are sequenced, generating sequence reads.

Sources of evidence

- Mate-pair illumina libraries
- Fosmid ends
- Bacterial artificial chromosomes
- 10x Genomics linked reads
- Hi-C
- Optical maps



Gaps consisting of N (unknown) bases are inserted between contigs

(Pseudo) chromosome assignment

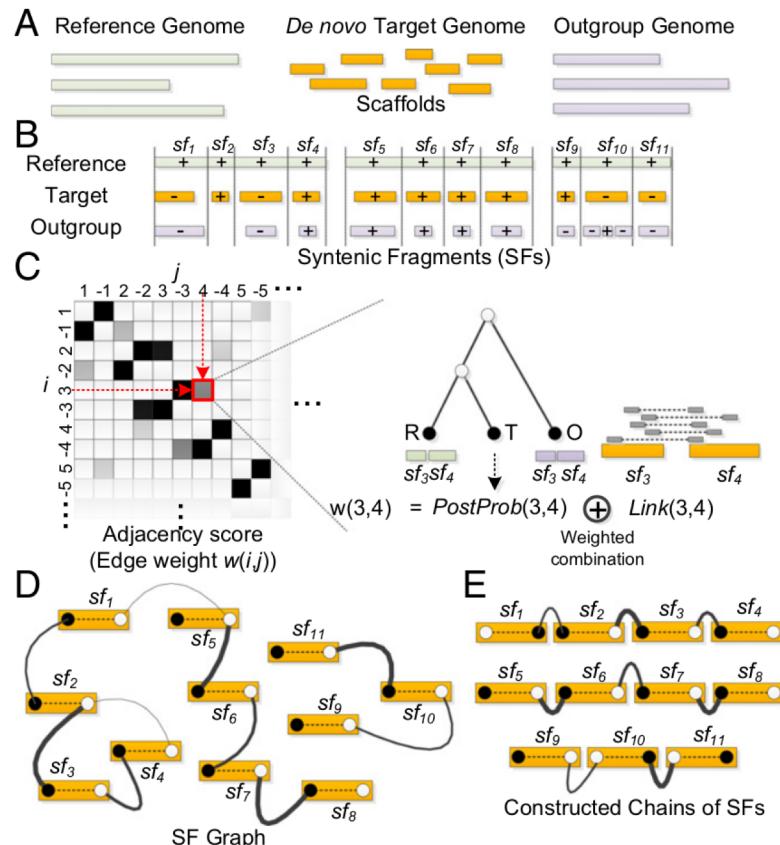
Goal: Assembly of chromosome-scale DNA fragments from scaffolds

Sources of evidence

- A close reference genome and/or outgroup genome
- Genetic markers or map

RACA software

- Input: Reference genome, de novo assembly, and one or more outgroups (A)
- Identify syntenic fragments (B)
- Calculate likelihood of adjacencies (C)



Reference-assisted chromosome assembly

Jaebum Kim^{a,b,1}, Denis M. Larkin^{c,1}, Qinglei Cai^d, Asan^e, Yongfen Zhang^d, Ri-Li Ge^{e,2}, Loretta Auvin^{f,g}, Boris Capitanu^{f,g}, Guojie Zhang^f, Harris A. Lewin^{a,b,2}, and Jian Ma^{a,1,2}

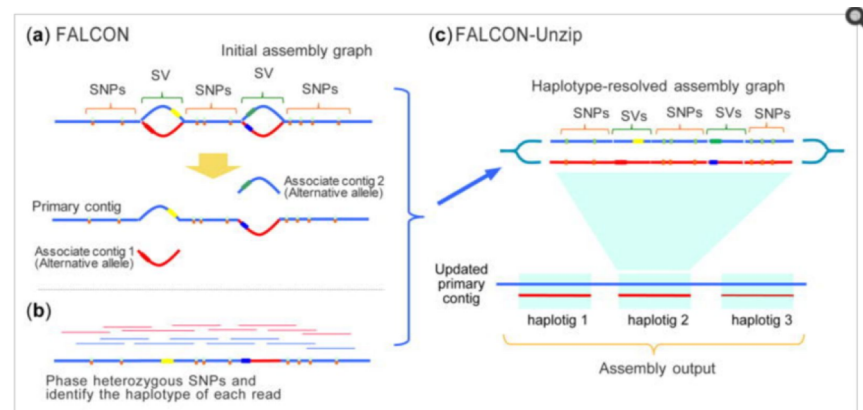
Diploid assembly

Almost all assemblers assume genome is haploid

- Ignore allelic variation between parental chromosomes

Diploid assembly: Produce two haplotype phased genomes

- e.g. Falcon-unzip



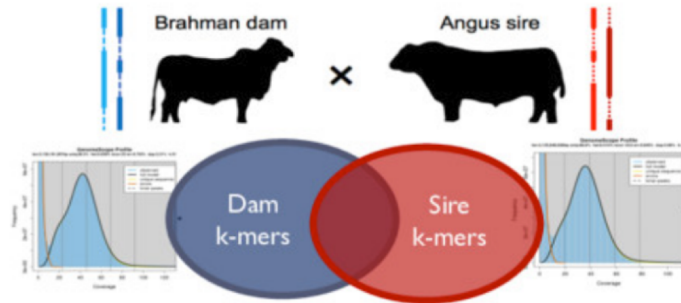
FALCON and FALCON-Unzip overview

(a) The initial assembly is computed by FALCON, which error corrects the raw reads (not shown) and then assembles using a string graph of the read overlaps. The assembled contigs are further refined by FALCON-Unzip into the final set of contigs and haplotypes. (b) Phase heterozygous SNPs and group reads by haplotype (c) The phased reads are used to open up the haplotype-fused path and generate as output a set of primary contigs and associated haplotypes.

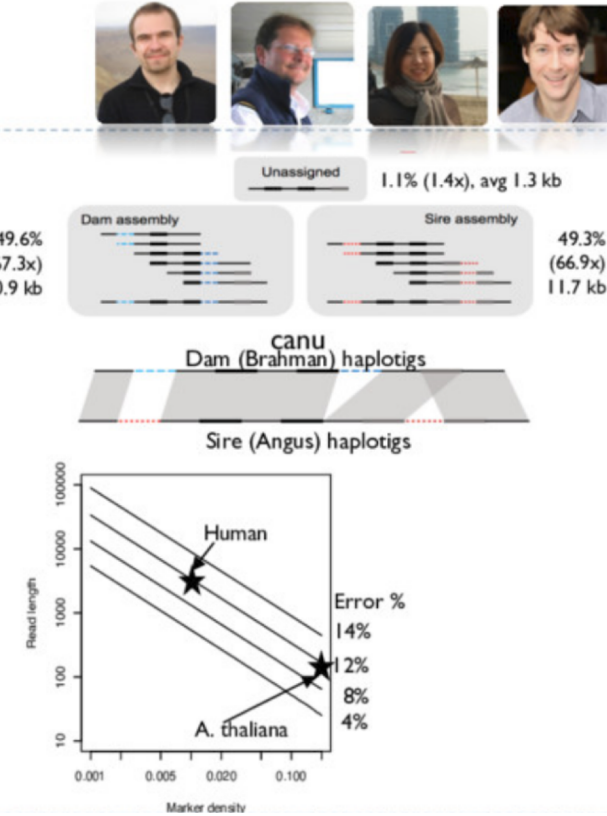
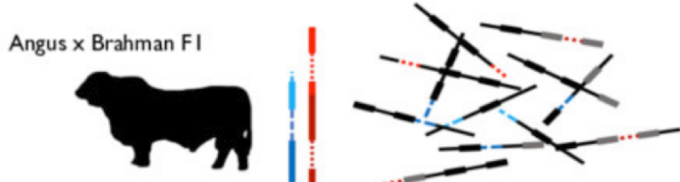
Diploid assembly

Trio Binning

- K-mer profiling of each parent (Illumina, 60x)



- K-mer profiling of the F1 (PacBio, 120x)



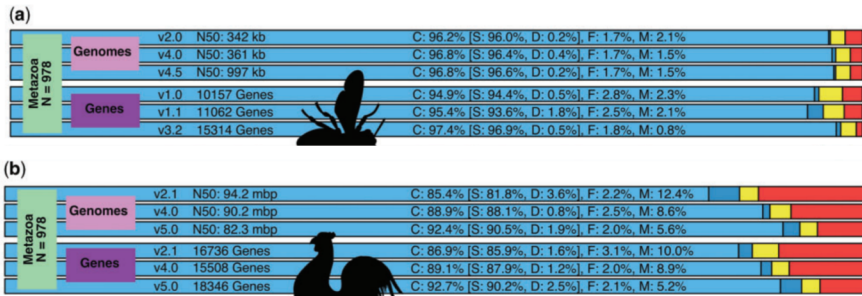
Genome assembly QC

Base Accuracy

- Realign reads from the same species
 - Identify SNPs+indels. Indels - known to be enriched in Pacbio+Nanopore assemblies
 - k-mer completeness with respect to Illumina data (KAT)

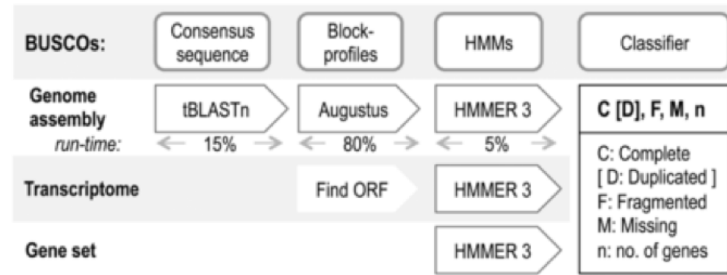
Local structure accuracy

- External evidence - e.g. sequencing data (REAPR, QUAST)
- Known adjacencies, e.g. PCR primers or structural variants



Gene content

- Order and orientation of genes/exons of known genes (e.g. housekeeping genes)
- BUSCO: Quantitative assessment of genome assembly and annotation completeness based on evolutionarily informed expectations of gene content



Benchmarking Universal Single-Copy Orthologs

- Start from set of conserved genes
- Find them in the assembly
- Assess whether they are Complete (C), Single copy (S) or Duplicated (D), Fragmented (F) or Missing (M)

BUSCO: assessing genome assembly and annotation completeness with single-copy orthologs

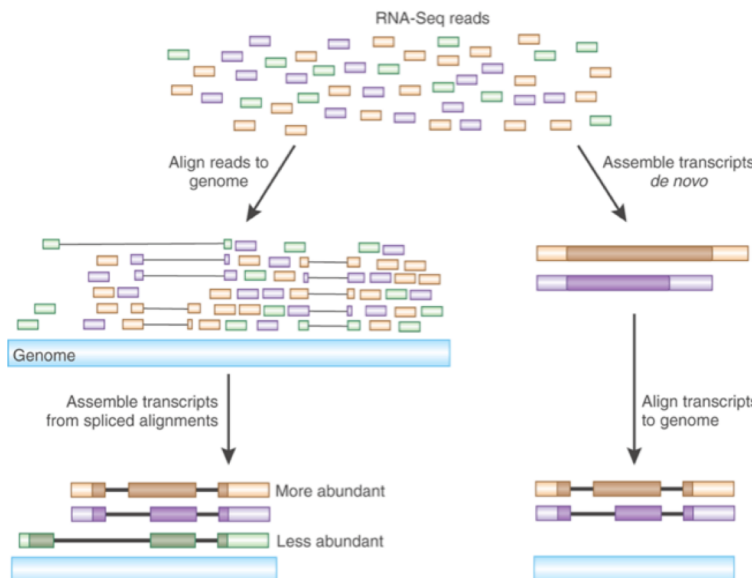
Felipe A. Simão, Robert M. Waterhouse, Panagiotis Ioannidis, Evgenia V. Kriventseva, Evgeny M. Zdobnov

Bioinformatics, Volume 31, Issue 19, 1 October 2015, Pages 3210–3212

Metagenomic assembly

- What if your DNA sample is a mix of uncultured (probably microbial) organisms?
- Abundances of the different components vary
 - Need to separate divergent sequences from errors
 - Output is typically a set of contigs, with coverage
 - Still a valuable simplification of a data set
- What scaffolding approaches are suitable?

Ayling M, Clark MD, Leggett RM. New approaches for metagenome assembly with short reads. *Brief Bioinform.* February 2019.
doi:10.1093/bib/bbz020.



Packages: Trinity, TopHat

Tree of Life Project



The Vertebrate Genomes Project (VGP), a project of the G10K Consortium, aims to generate near error-free reference genome assemblies of all 66,000 extant vertebrate species.

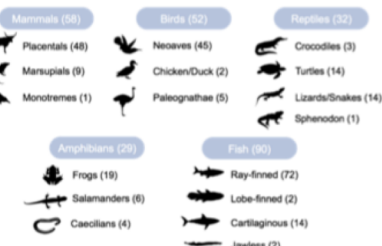


VertebrateGenomesProject.org



PACBIO® + 10X GENOMICS® +

BIONANO GENOMICS + ARIMA



Celebrate 25th anniversary of the Sanger Institute by sequencing 25 genomes representing five key areas of biodiversity in Britain

- **Cryptic:** Species that are out of sight, or have identical forms that are different in behaviour
- **Dangerous:** Invasive and harmful species
- **Floundering:** Endangered and declining species
- **Flourishing:** Species on the up in the UK
- **Iconic:** Species that represent the British countryside

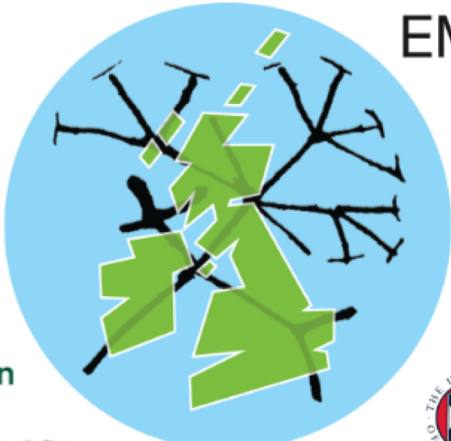


- New UK initiative to sequence all ~66,000 eukaryotic UK species.
- Collaboration between a number of UK scientific centres, including Wellcome Sanger Institute, Earlham Institute, Natural History Museum, Kew Gardens, Royal Botanic Garden Edinburgh

Darwin Tree of Life Project



Royal
Botanic Garden
Edinburgh



EMBL-EBI



UNIVERSITY OF
CAMBRIDGE

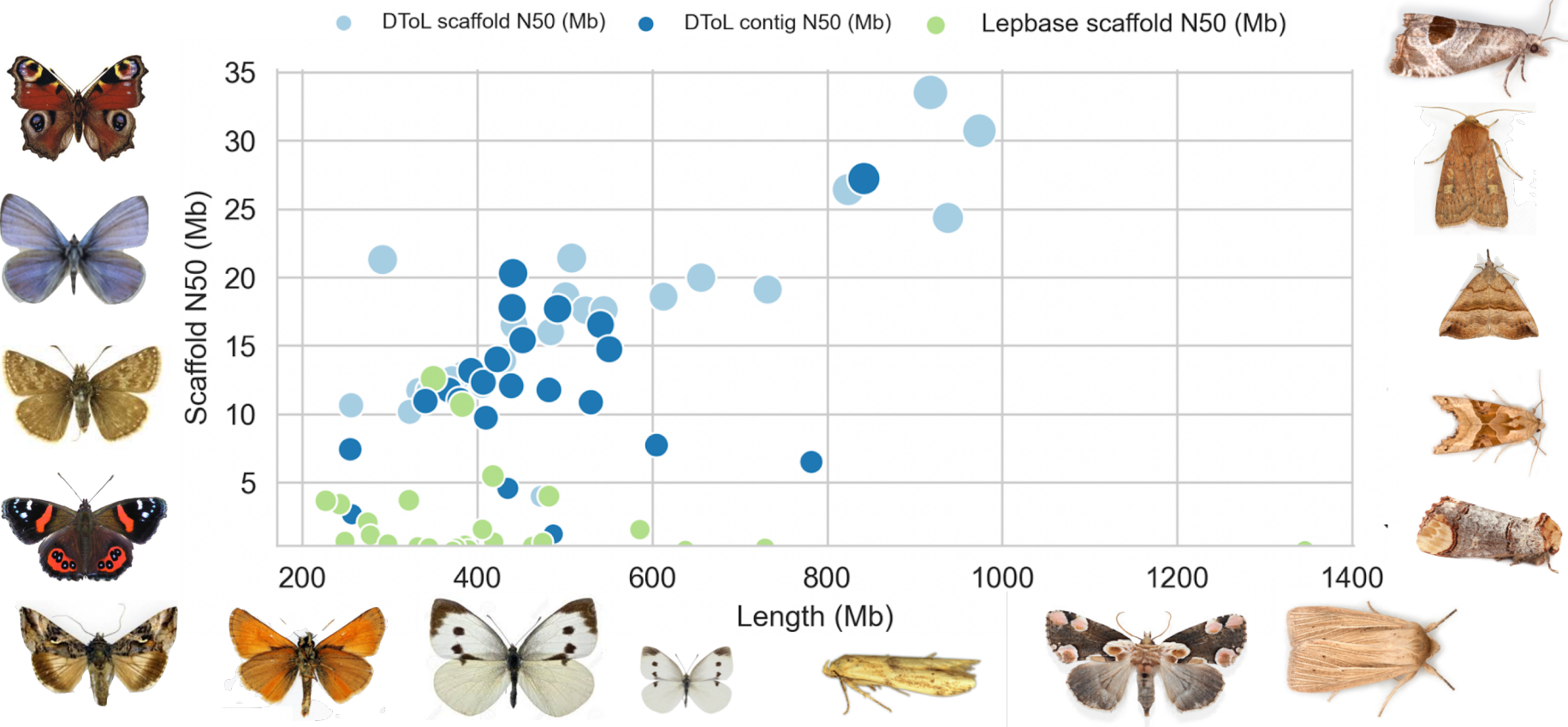


THE UNIVERSITY
of EDINBURGH

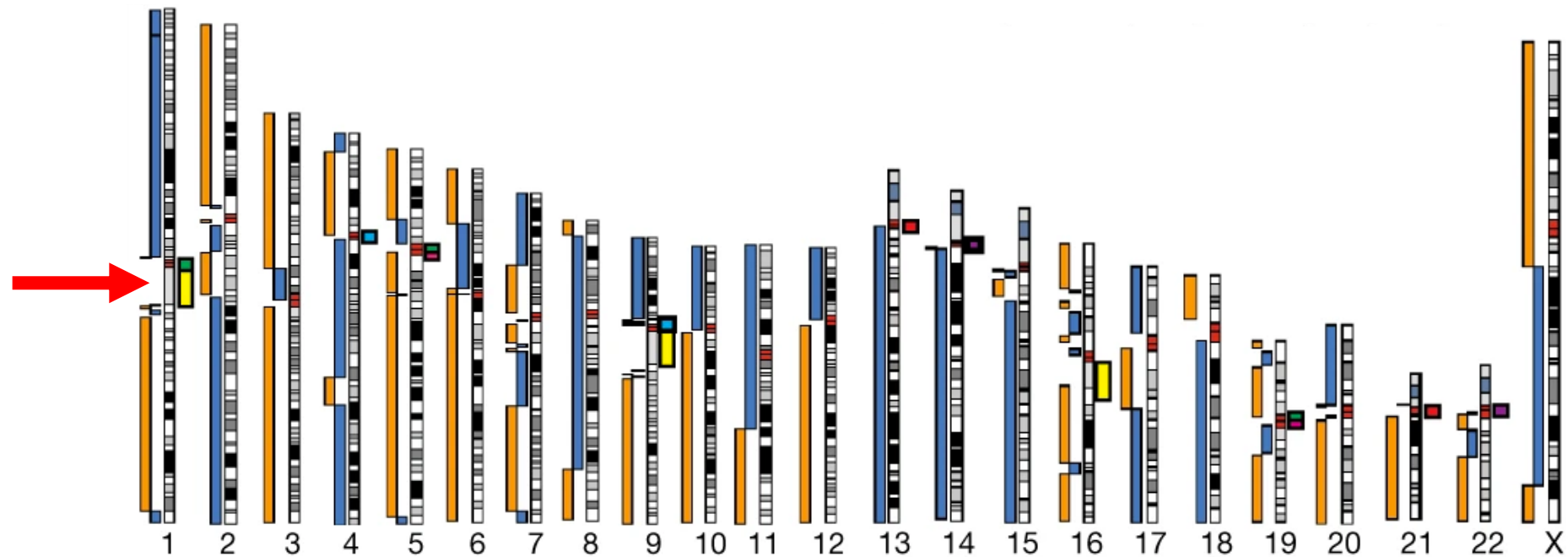


UNIVERSITY OF
OXFORD

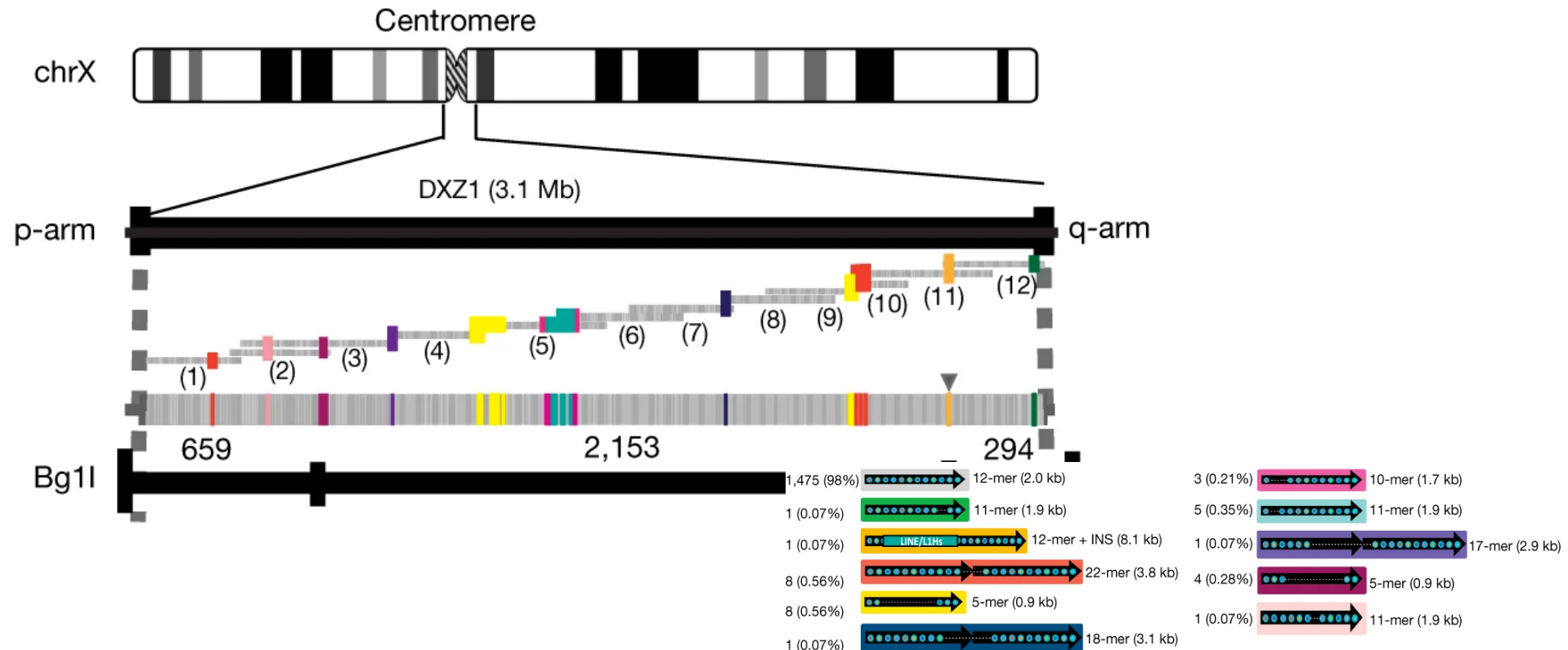
48 New Lepidoptera assemblies



Complete chromosome contigs



Complete chromosome contigs



Practical

- Assembly of a *Plasmodium falciparum* (Malaria) chromosome
 - Basics of assembly algorithms
 - Assembly with PacBio (get to this quick it may take awhile!)
 - Assembly with Illumina
 - Union (PacBio + Illumina) assembly and polishing
 - Assembly QC