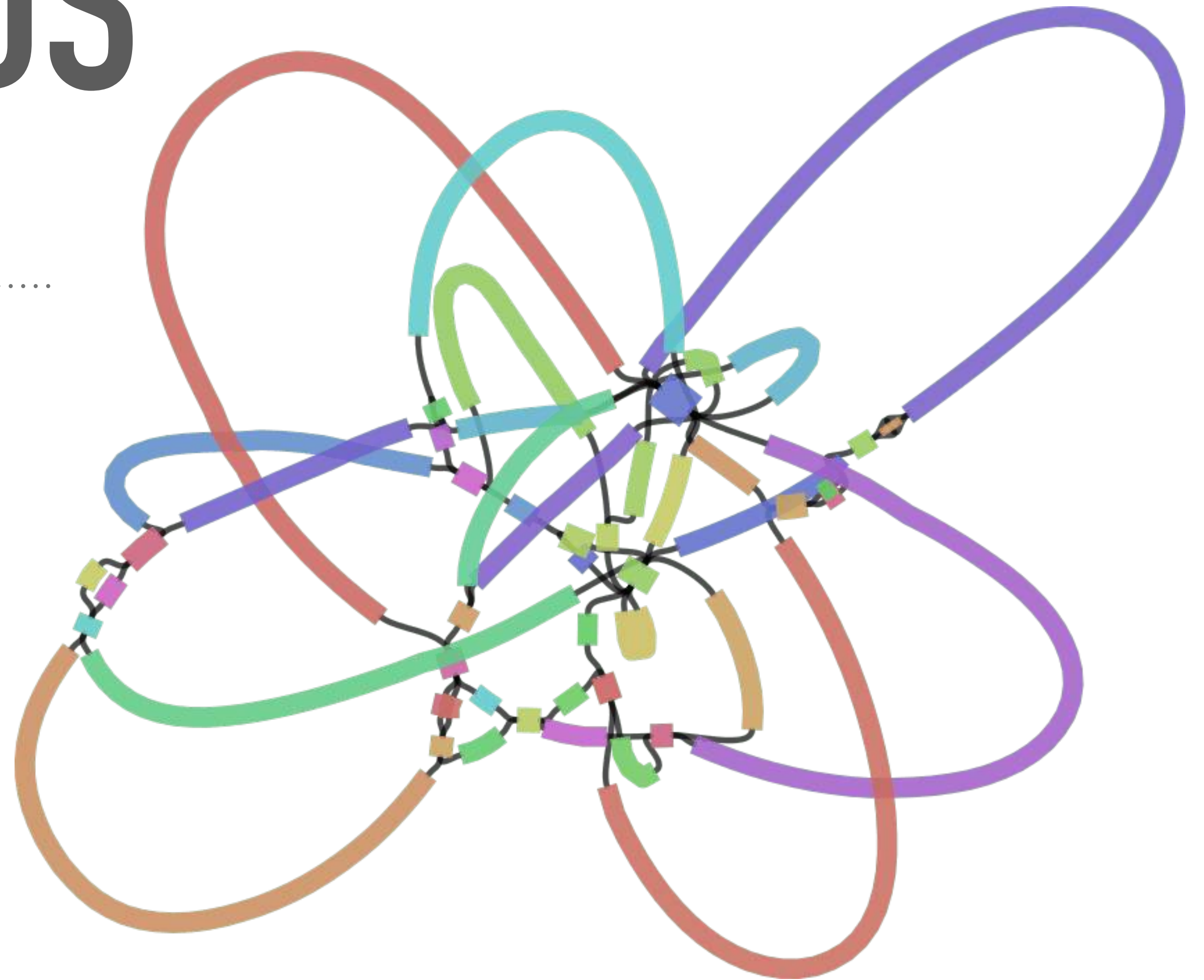


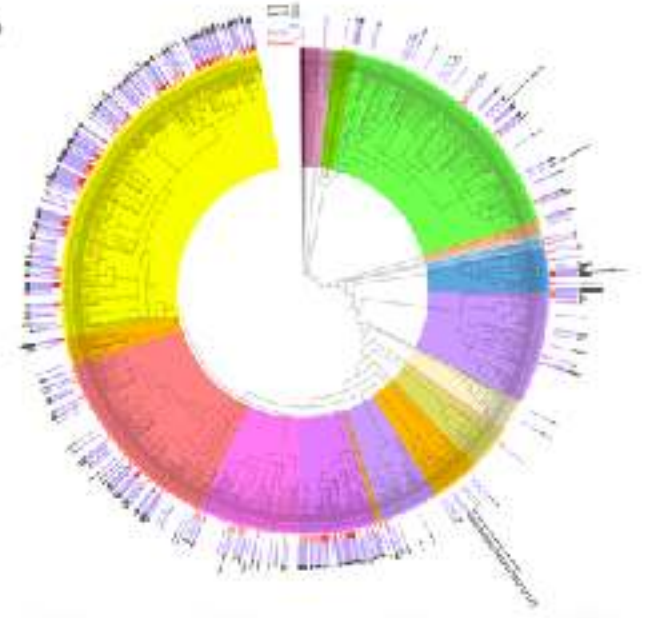
GENOME ASSEMBLY WITH LONG READS

Marcela Uliano-Silva



WHO AM I?

- Senior Bioinformatician Wellcome Sanger Institute - Darwin Tree of Life Project. Tree of Life Assembly Team (ToLA)
 - Churchill College Postdoctoral By-Fellow, University of Cambridge
-
- BSc in Biology (2010) - UFSC, Brazil
 - MSc in Biophysics (2013) - IBCCF UFRJ, Brazil
 - PhD in Biophysics (2017) - IBCCF UFRJ, Brazil
 - Horizon2020 Marie Curie PostDoc Fellow (2017-2019), IZW BeGenDiv (Germany)
 - TED Fellow





Darwin
TREE
of
LIFE

Tree of Life: Major Projects

Collaborating widely to deliver across diversity



★ Darwin Tree of Life Project

- 70,000 species from Britain and Ireland [Phase 1: 2,000 species]



★ Aquatic Symbiosis Genomics

- 1,000 species (500 symbiotic systems) from marine and freshwater



★ Vertebrate Genomes Project

- Realising VGP Phase 1 (ordinal - 260 species) and Phase 2 (family) goals



★ European Reference Genome Atlas

- Sequencing the genomes of all species in the European continent - Pilot 25 species



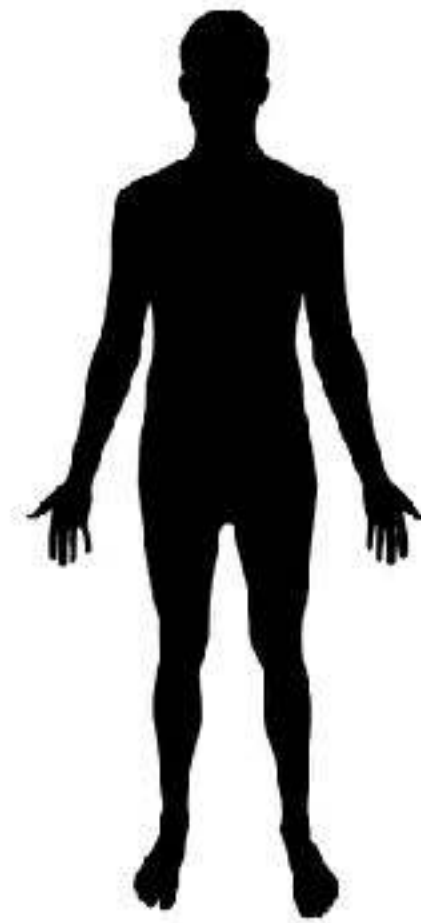
★ Earth BioGenome Project

- Working to deliver Phase 1 (family) goals, and to "sequence all life for the future of life"



Genome assembly: what is my goal?

- Understand variation in populations (disease-related SNPs etc...)
- Study the molecular profile of a species never before sequenced (evolutionary studies etc..)



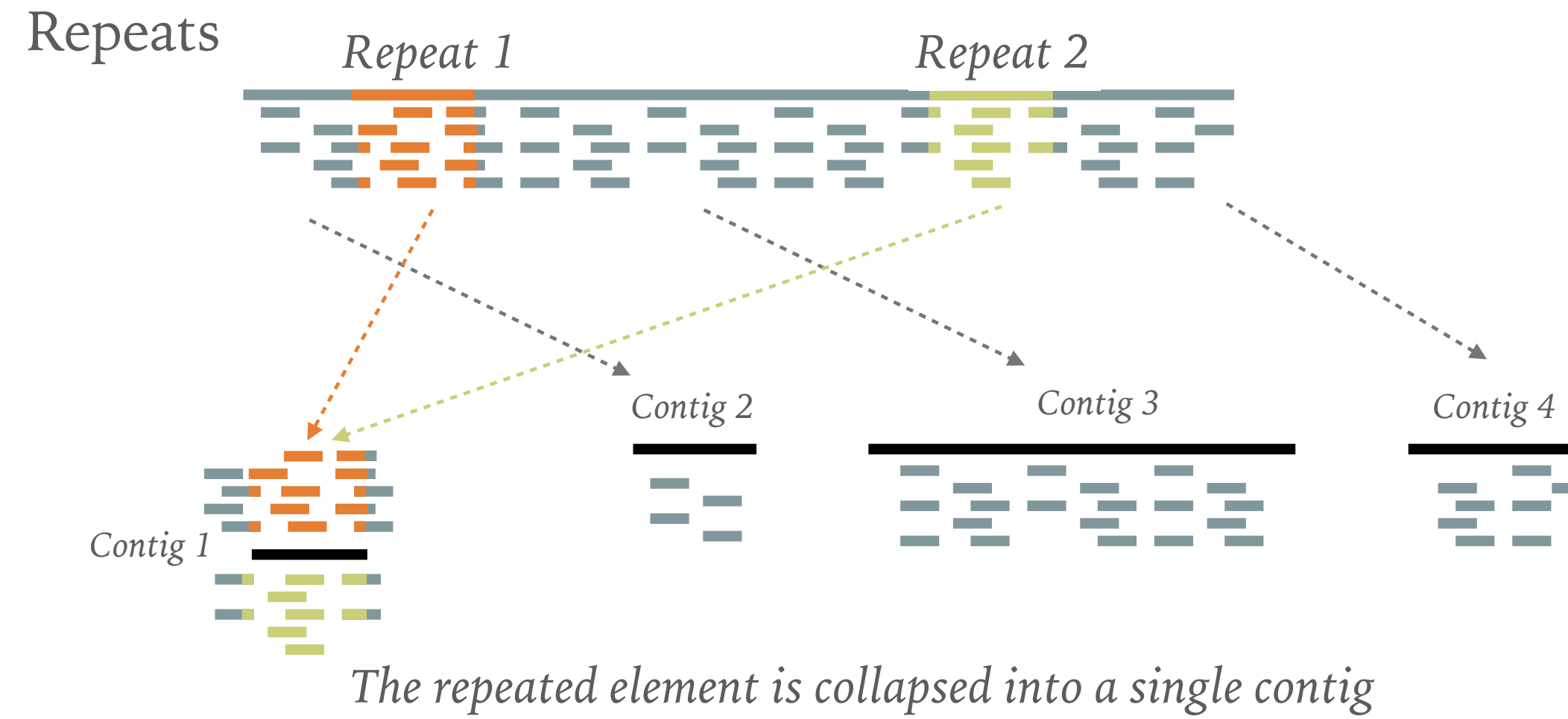
Genome re-sequencing
Assembly by mapping to a reference



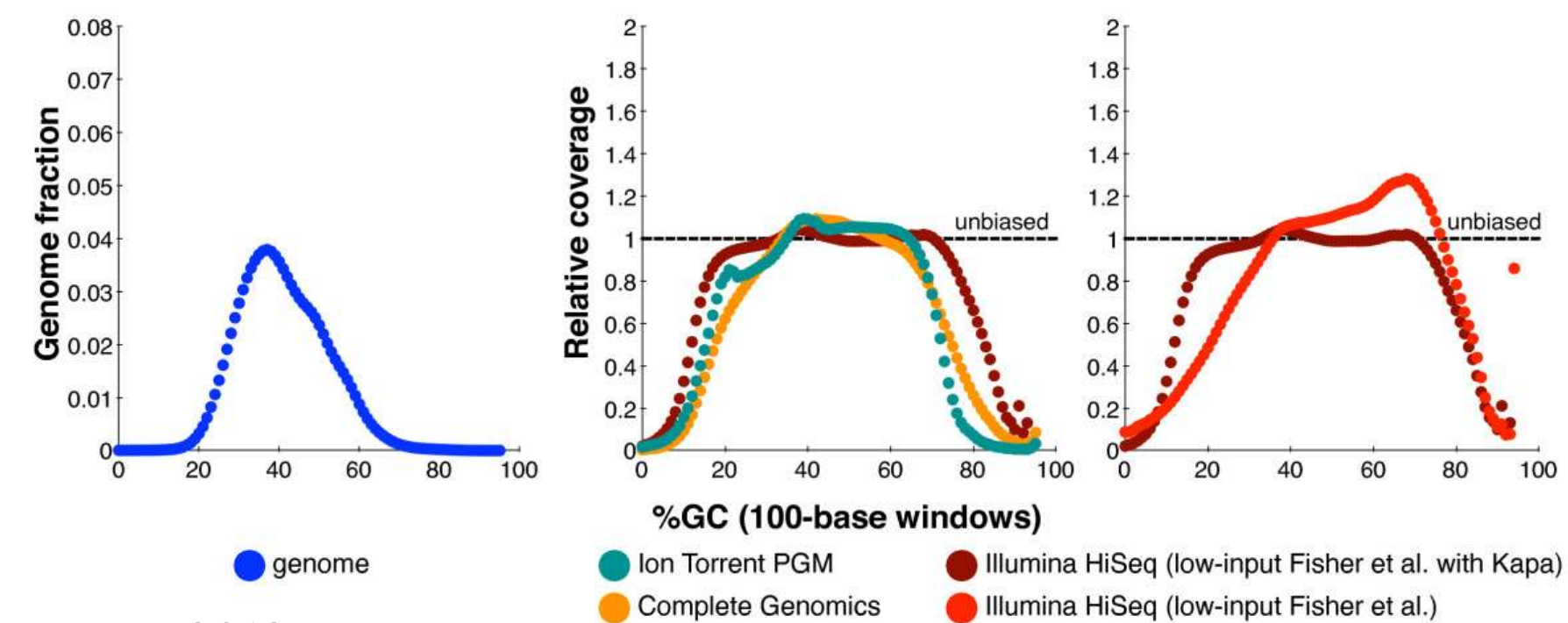
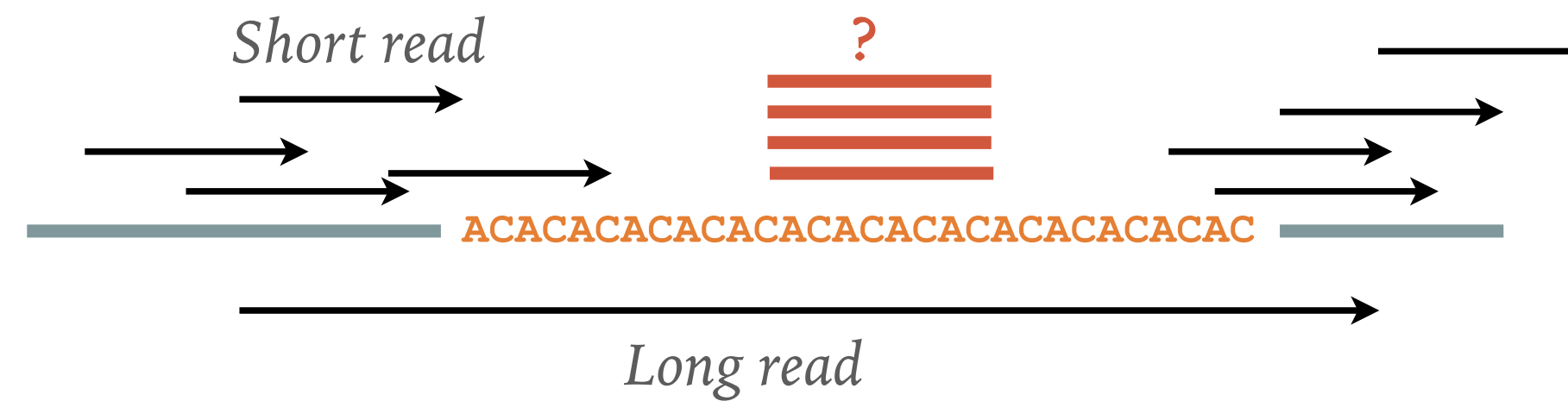
De novo assembly

Hurdles

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

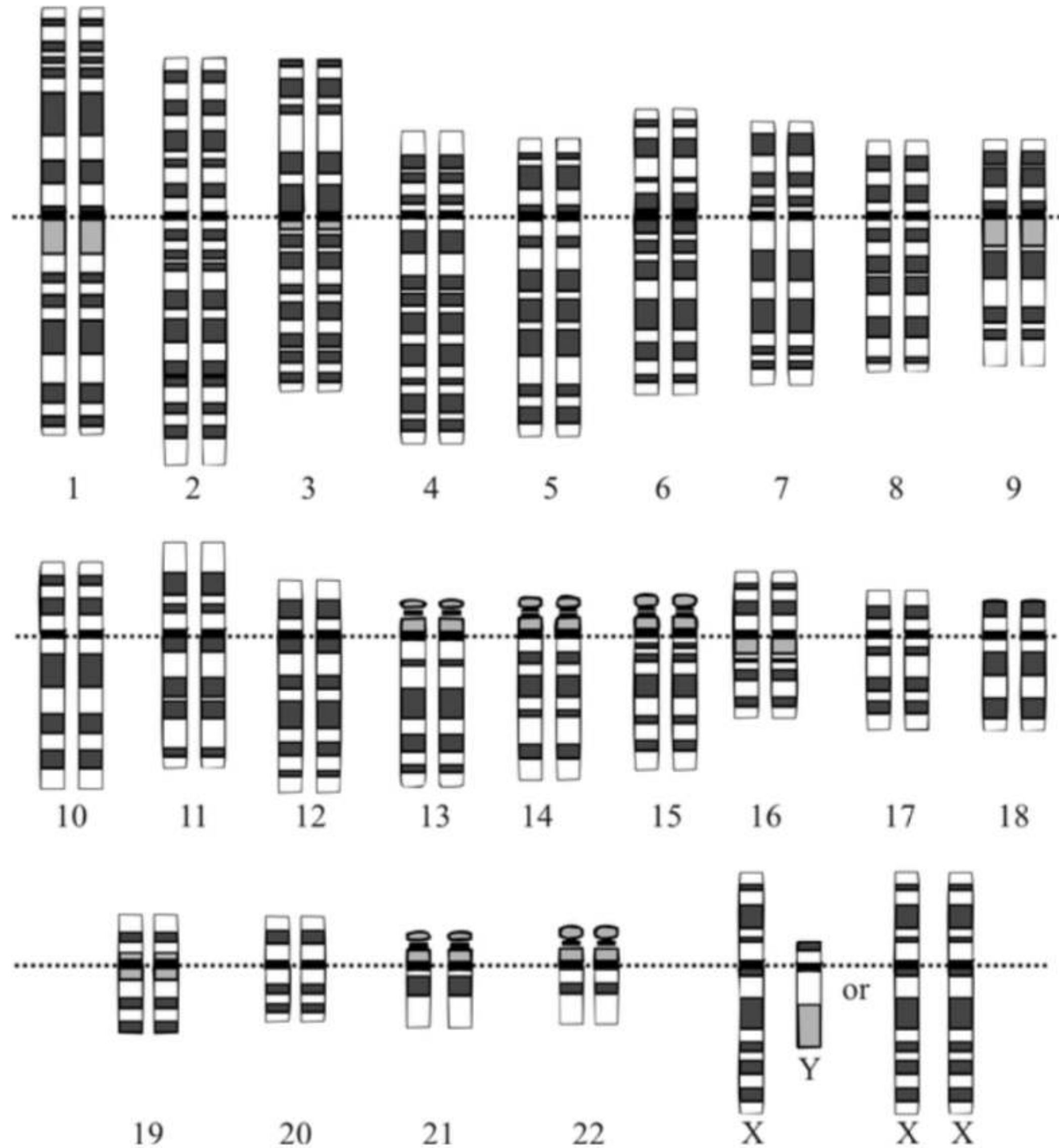


Adapted from Torsten Seemann (2014 talk)



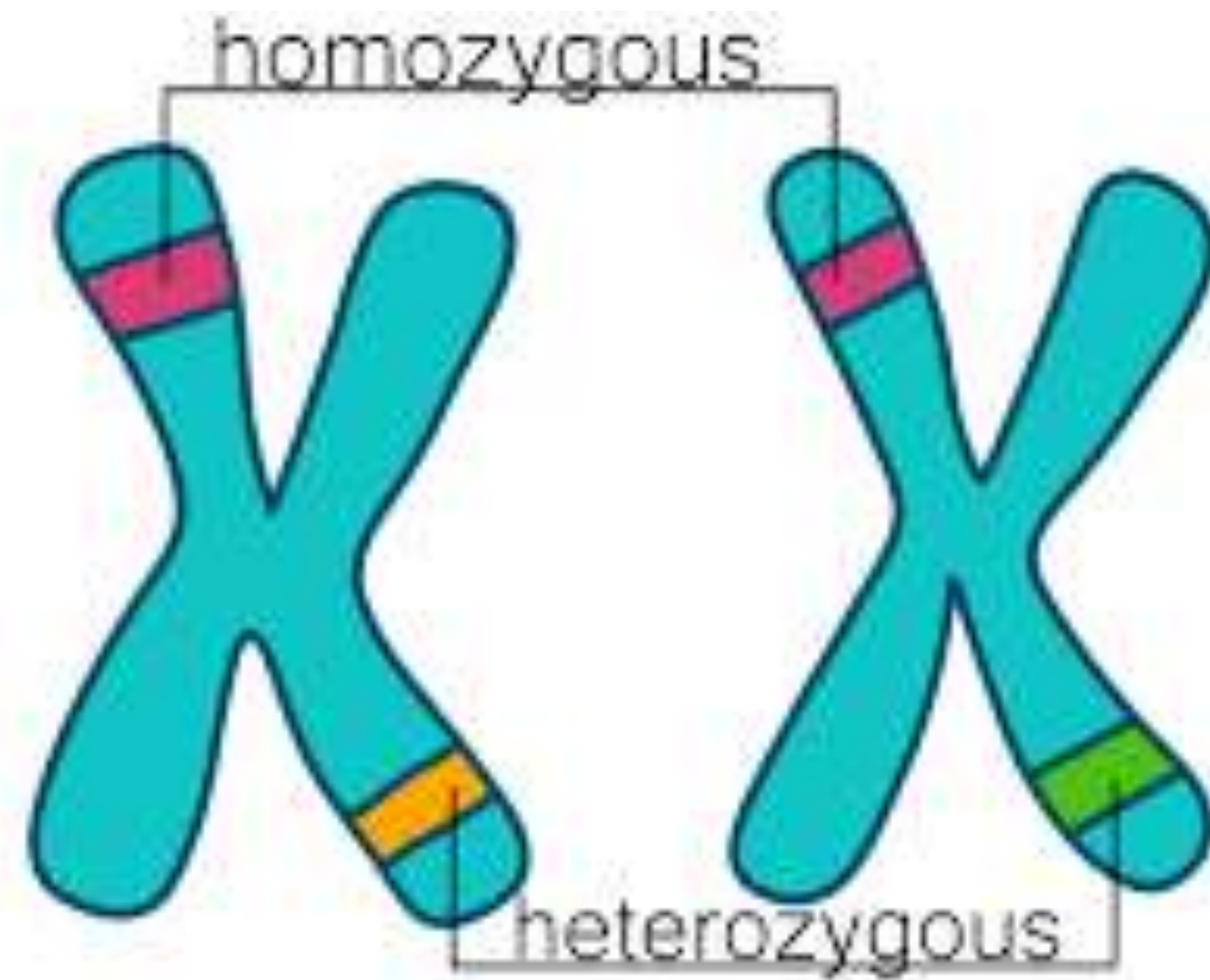
Ross, 2013

THE HUMAN GENOME – AGAIN



*Inside the nucleus of a somatic cell, we will have 6 billion bases of DNA from our genome, as we are **diploid** organisms. The reference human genome is the representation of one copy of each chromosome **allele**, thus 3 billion bases on average.*

ANOTHER PROBLEM – GENOME HETEROZYGOSITY



A. Diploid homologous chromosomes



B. Partially resolved de-novo assembled contigs

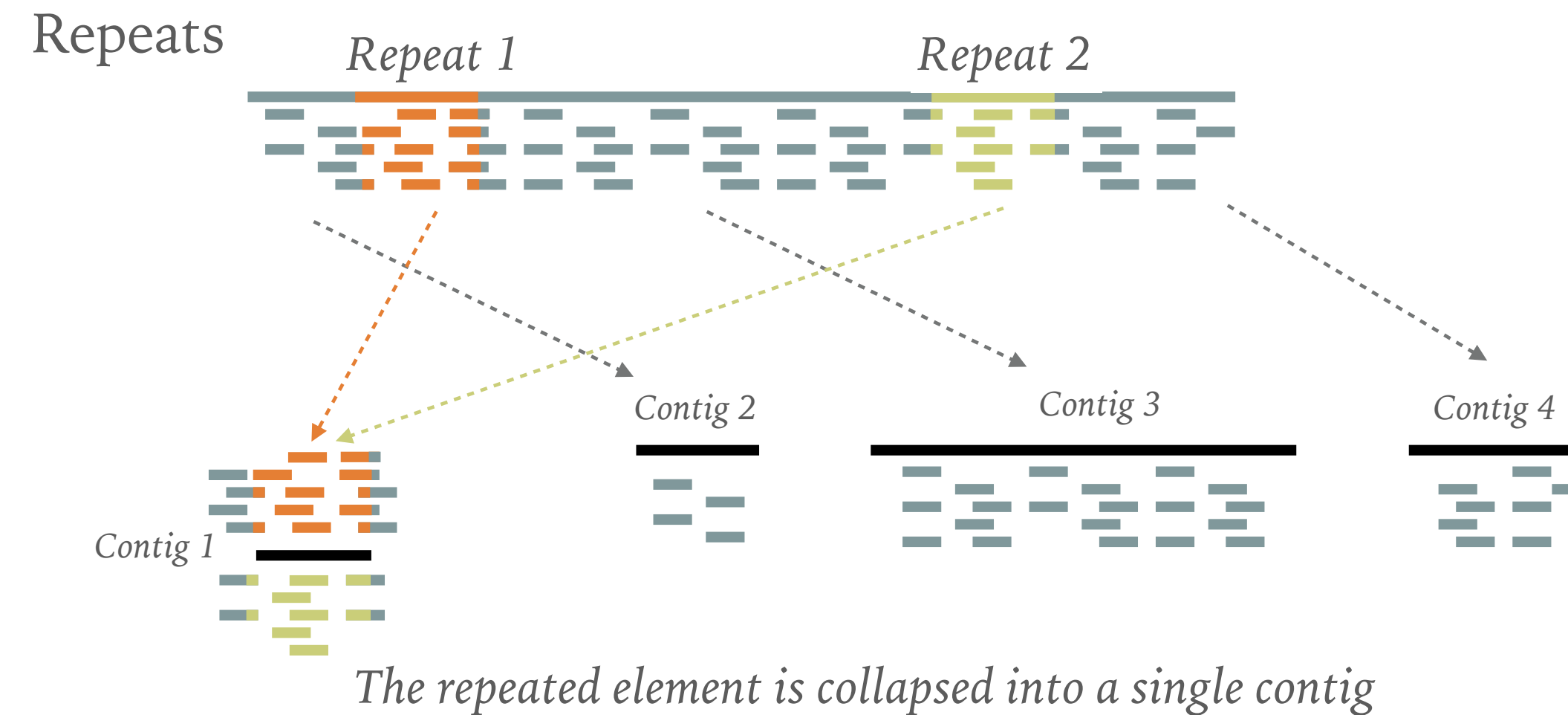


c. Pseudo-haploid Assembly

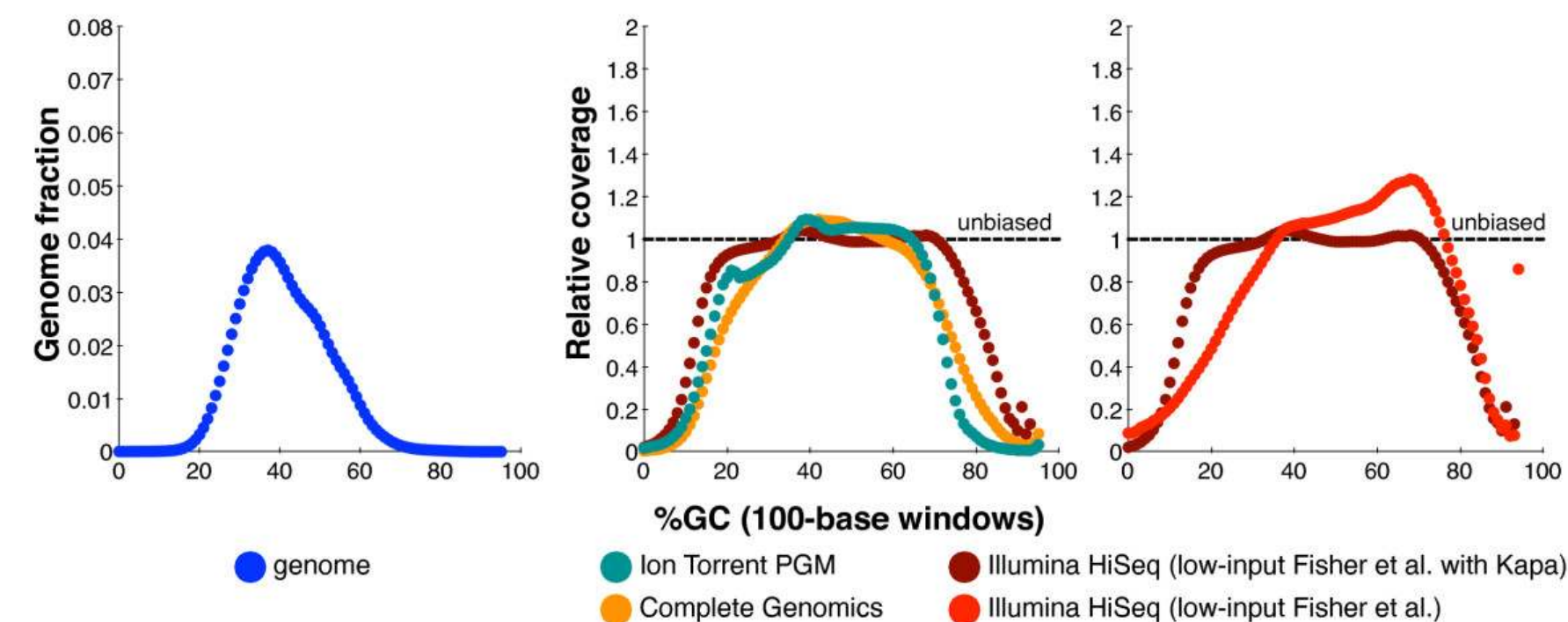
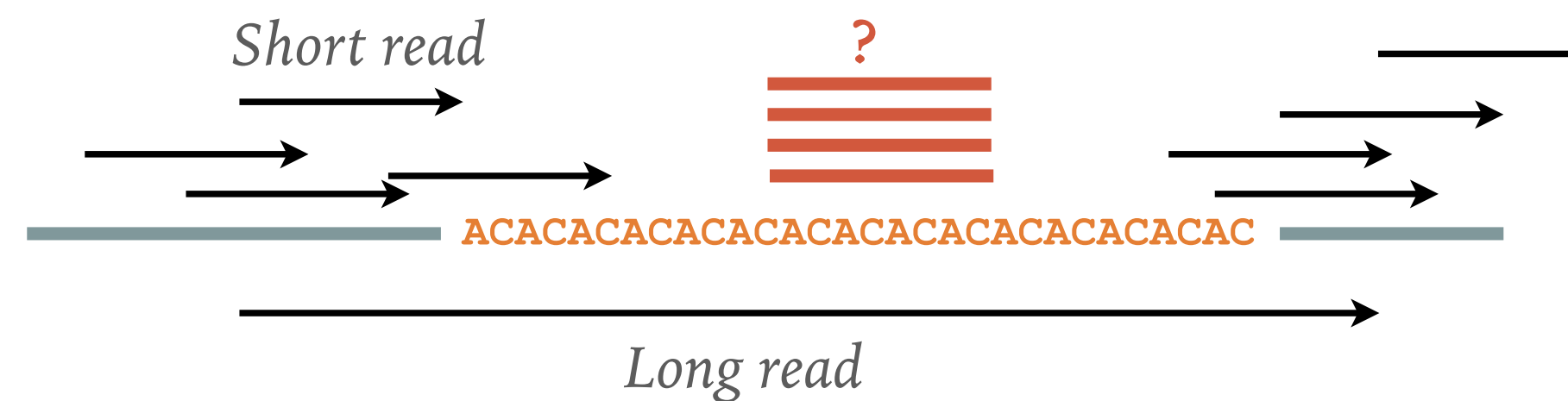


Hurdles

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



Adapted from Torsten Seemann (2014 talk)

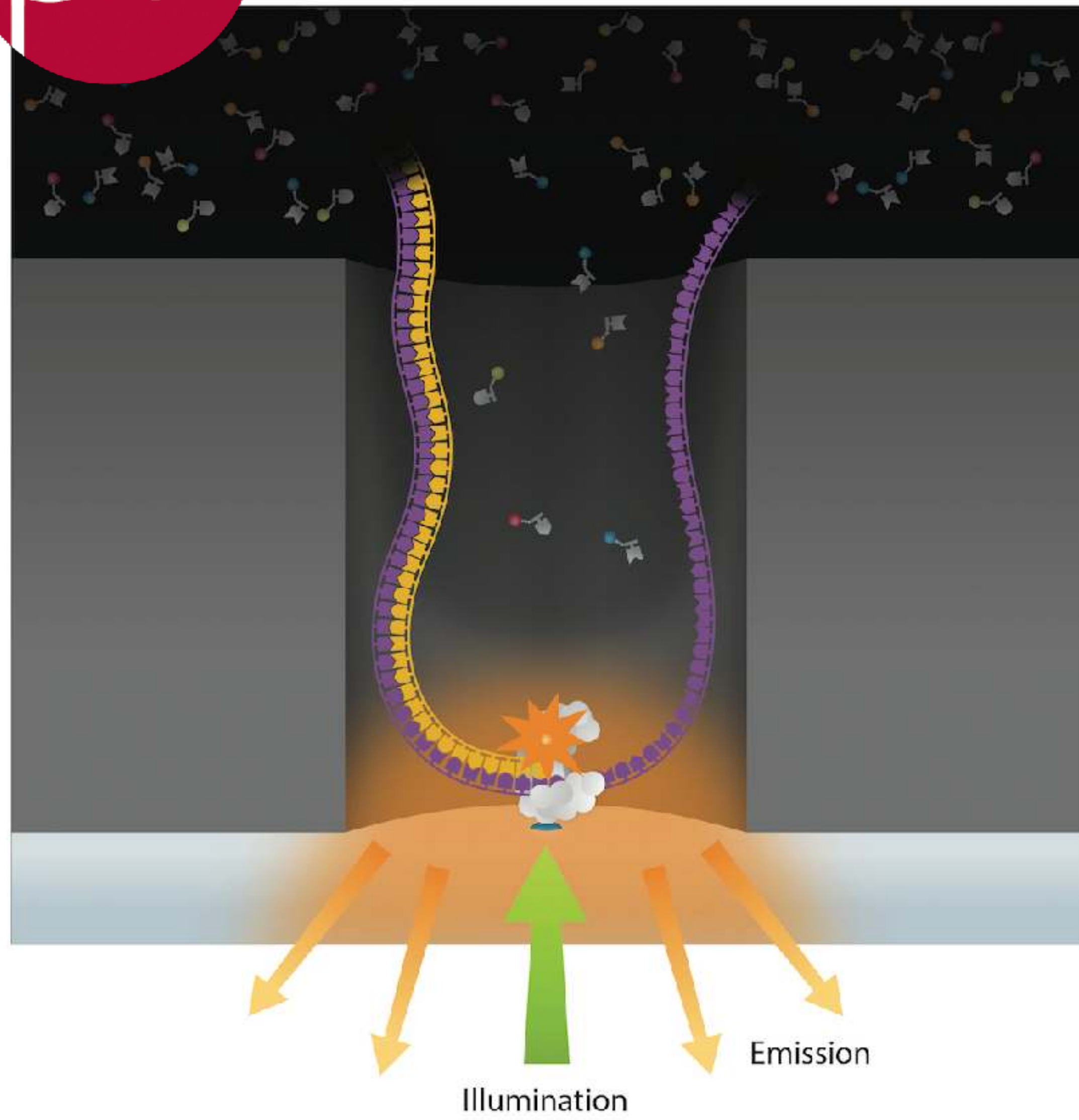


Ross, 2013

LONG READS



PACBIO®



Single molecule sequencing

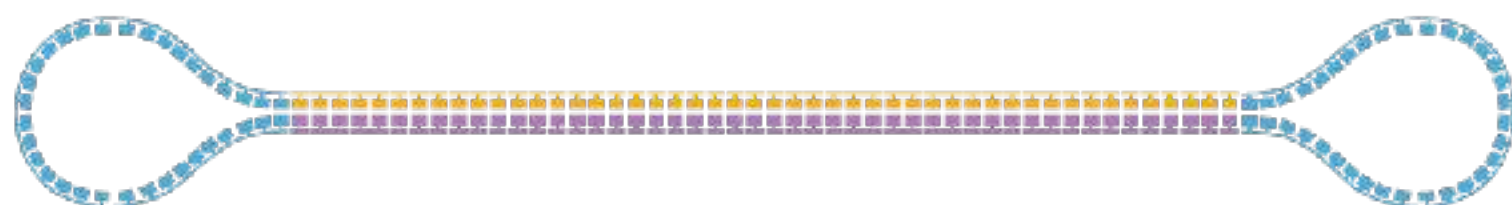
DNA Polimerase: 1000bp/s

PACBIO READS

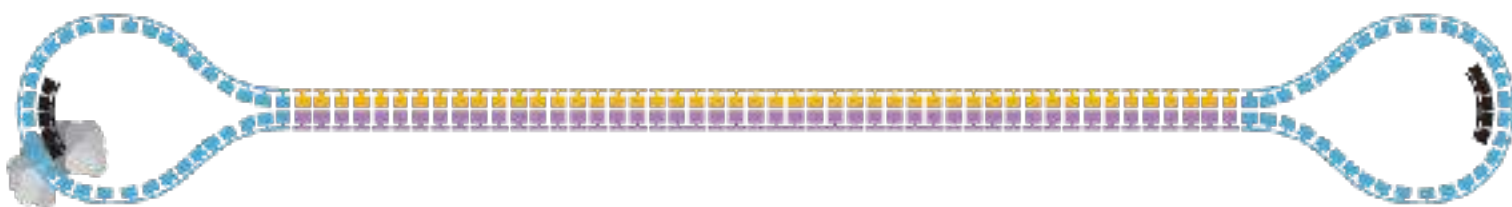
Start with high-quality double stranded DNA



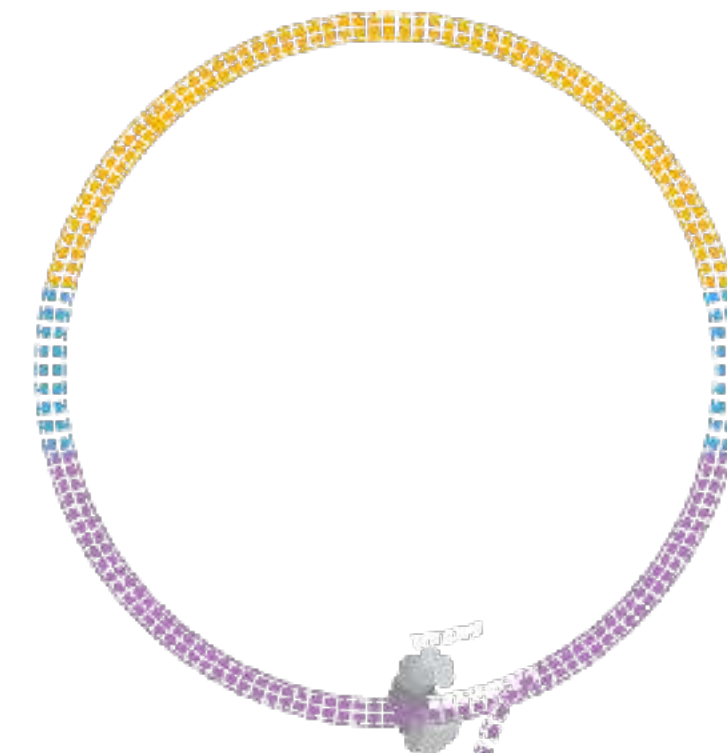
Ligate SMRTbell adapters and size select



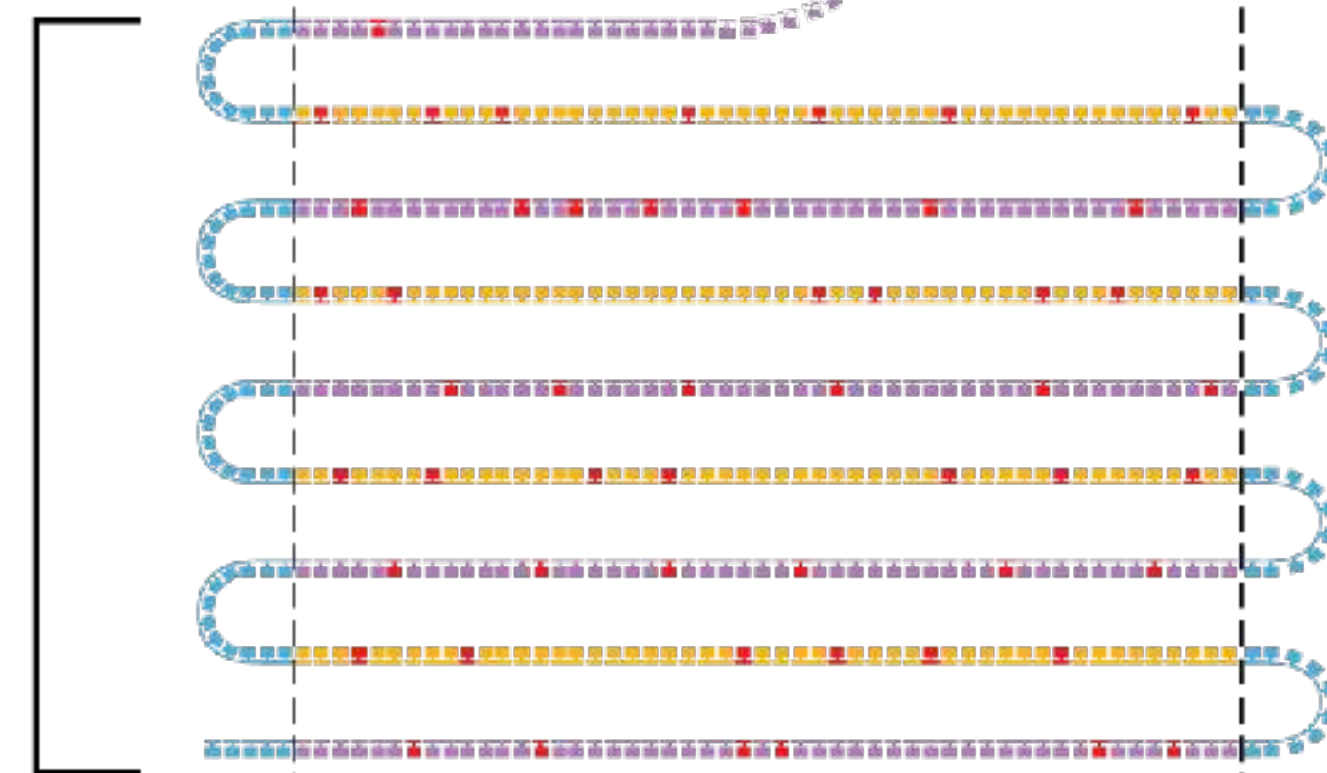
Anneal primers and bind DNA polymerase



Circularized DNA is sequenced in repeated passes



The polymerase reads are trimmed of adapters to yield subreads



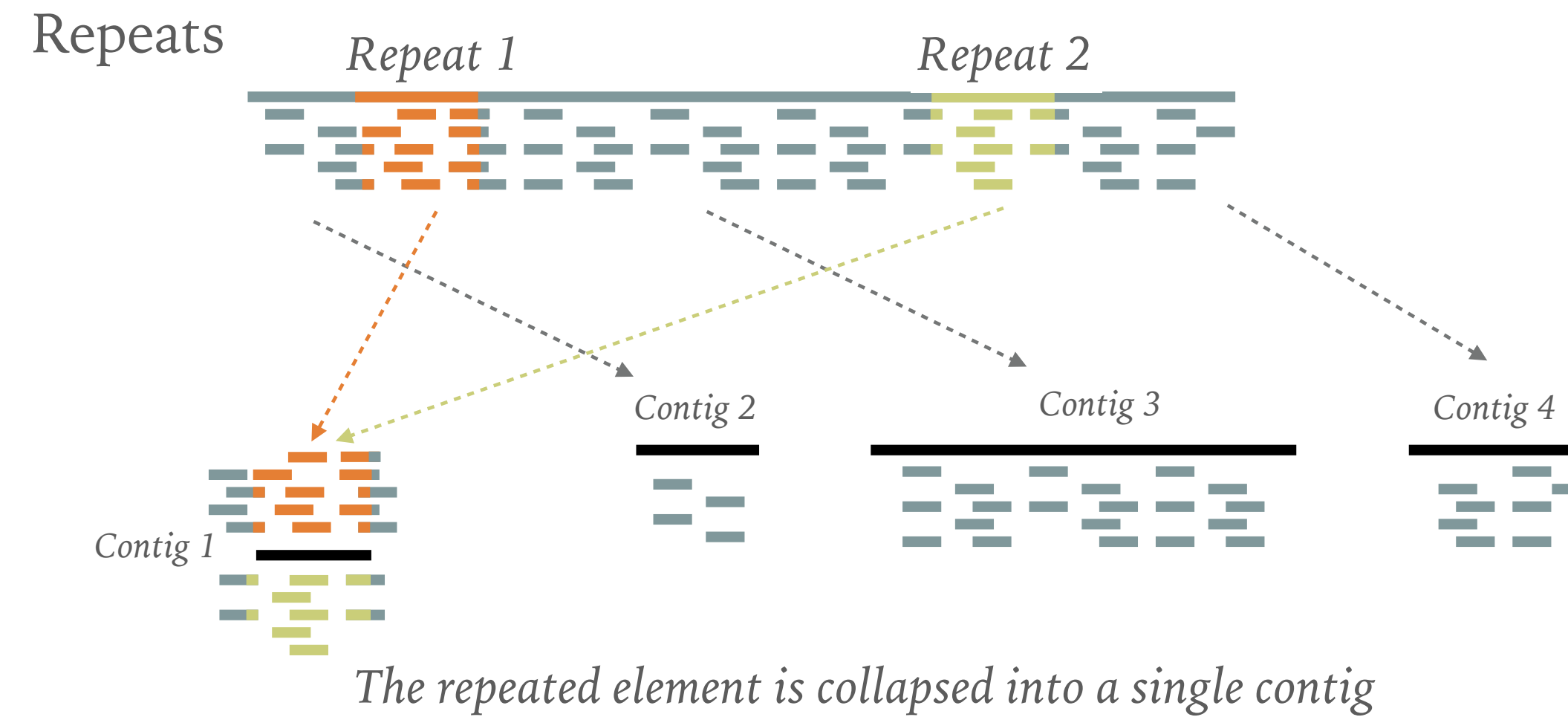
Consensus is called from subreads



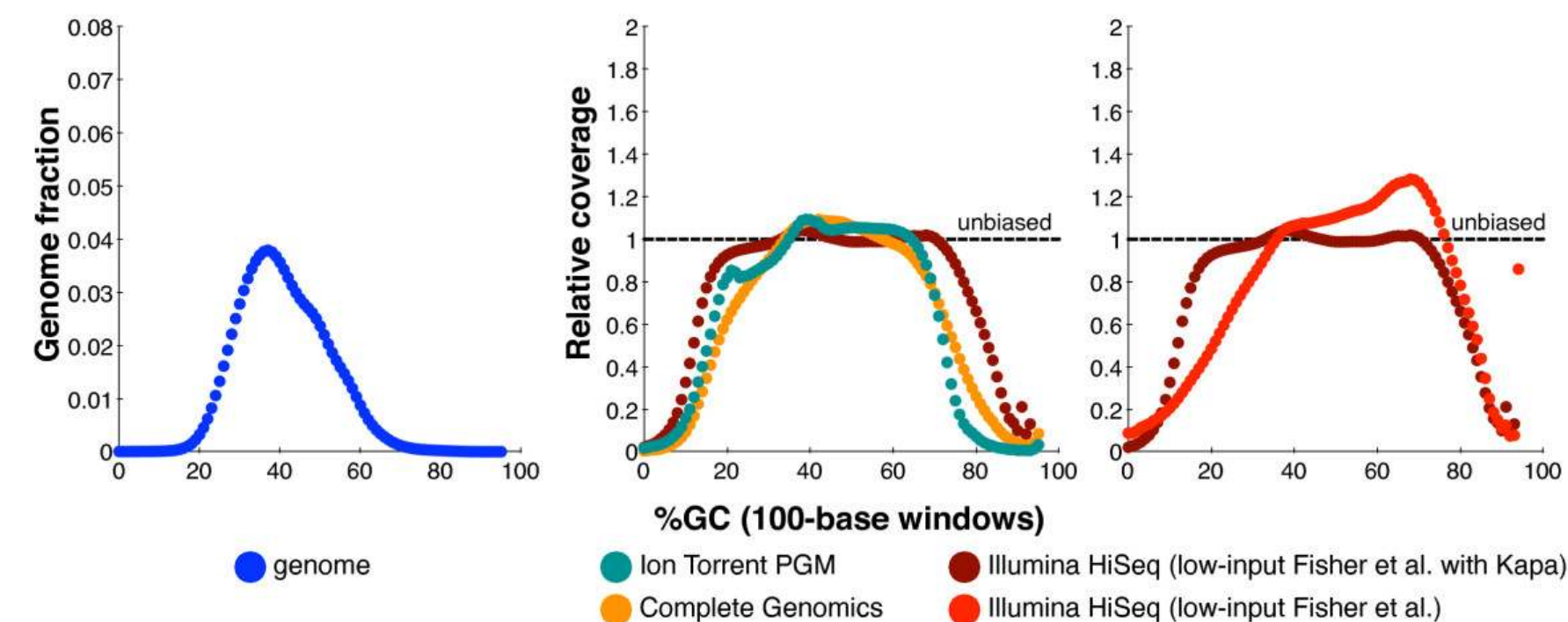
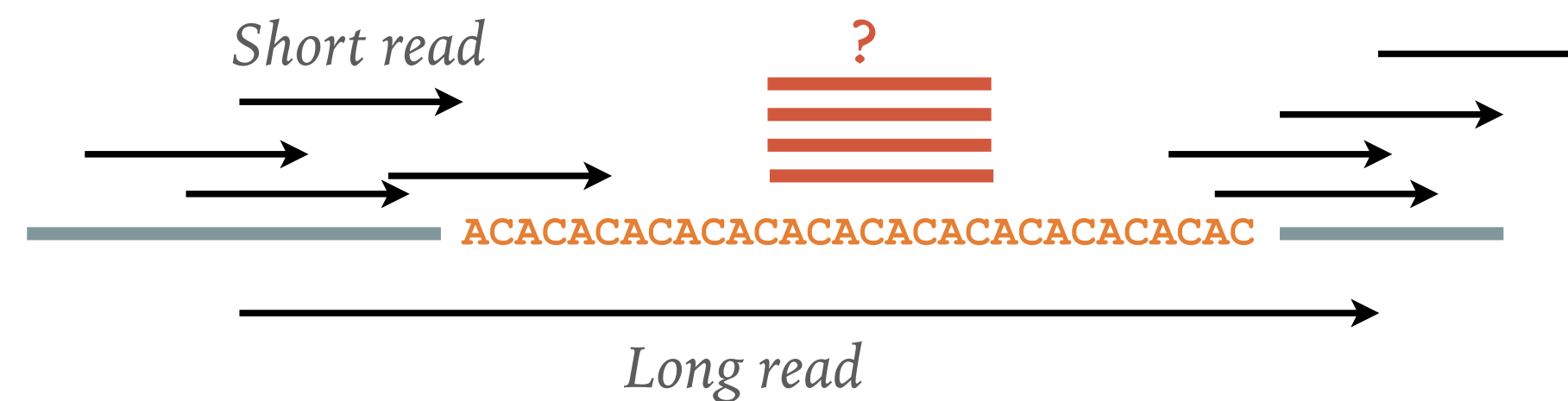
HiFi READ
(>99% accuracy)

Hurdles

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

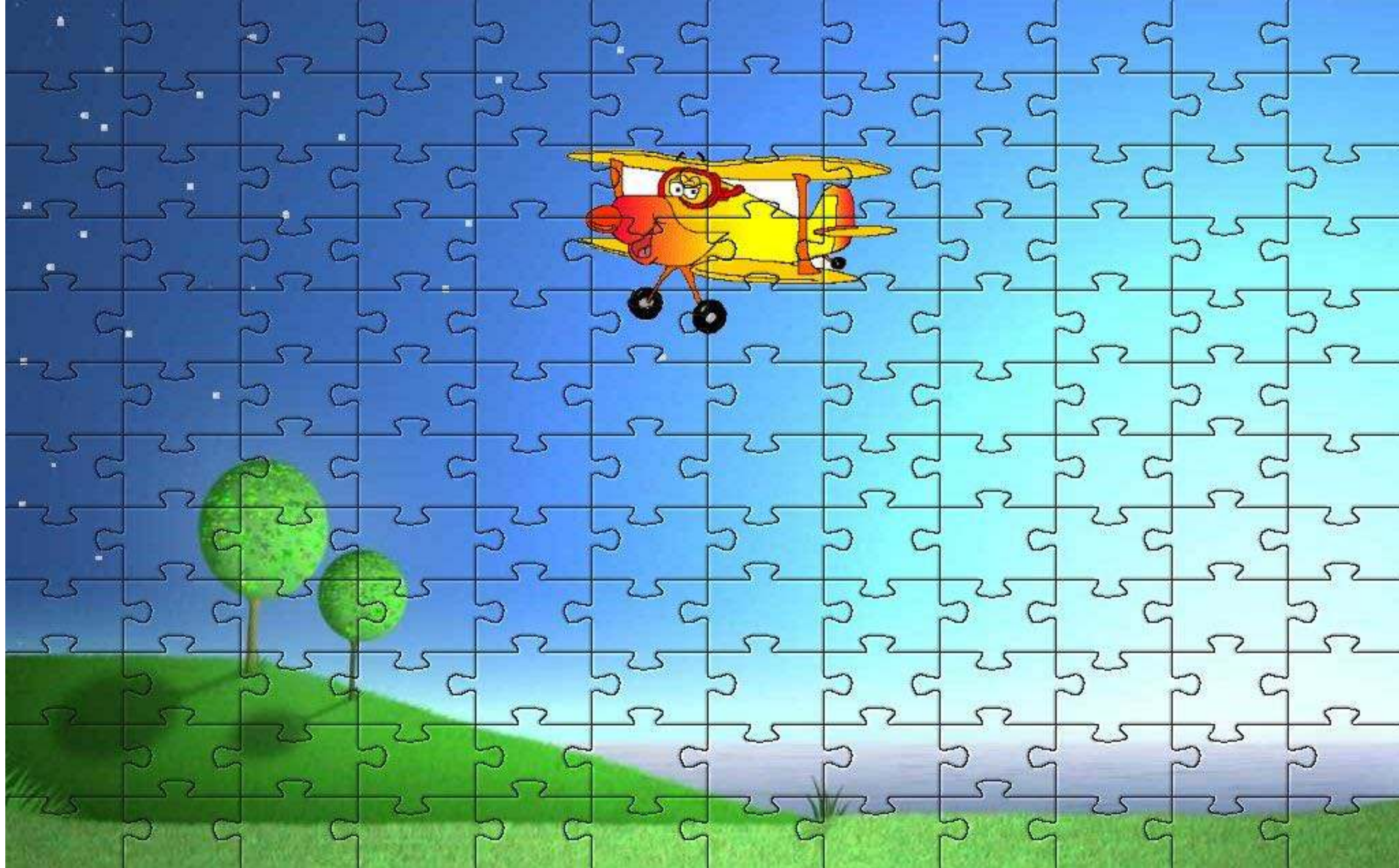


Adapted from Torsten Seemann (2014 talk)



Ross, 2013

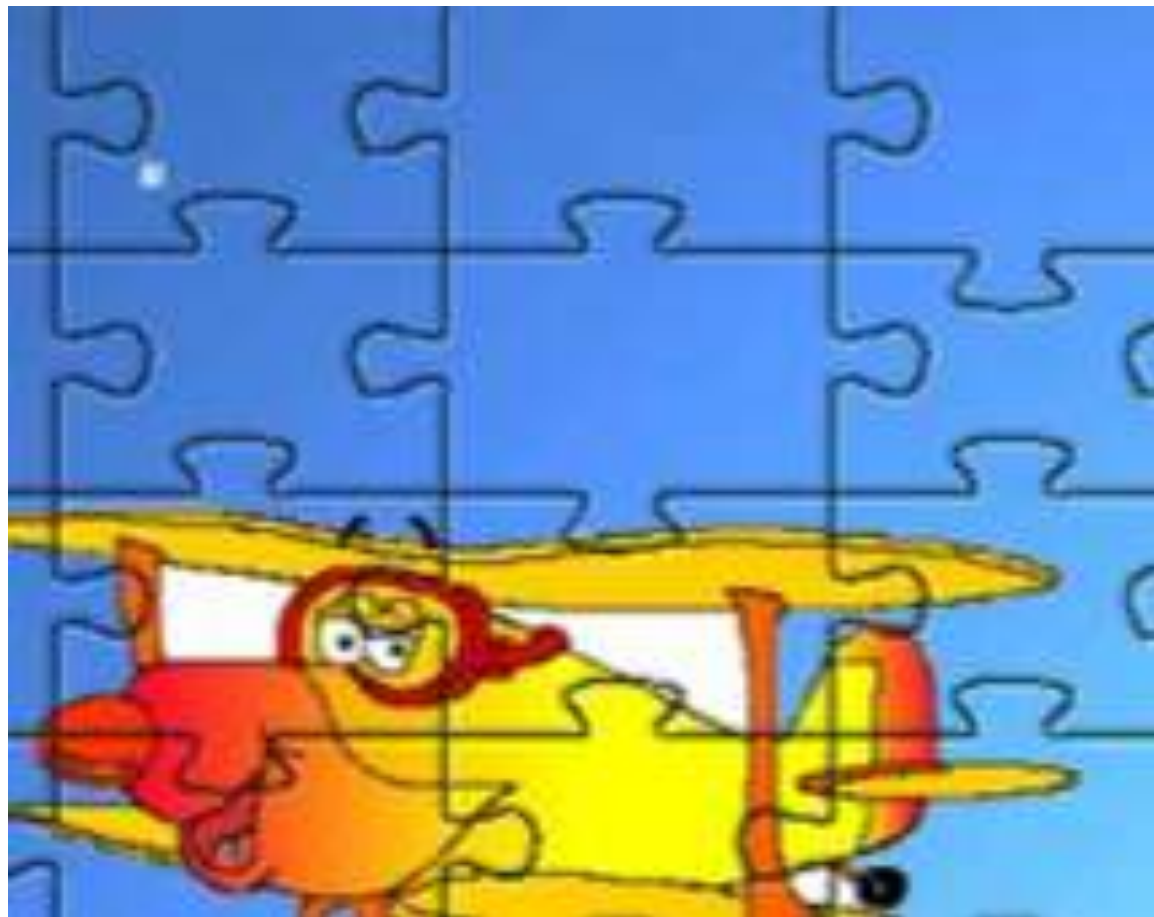
I'M A EUKARYOTIC GENOME – THE BLUE AND GREEN ARE MY REPEATS



THIS IS A SHORT-READS GENOME ASSEMBLY OF ME

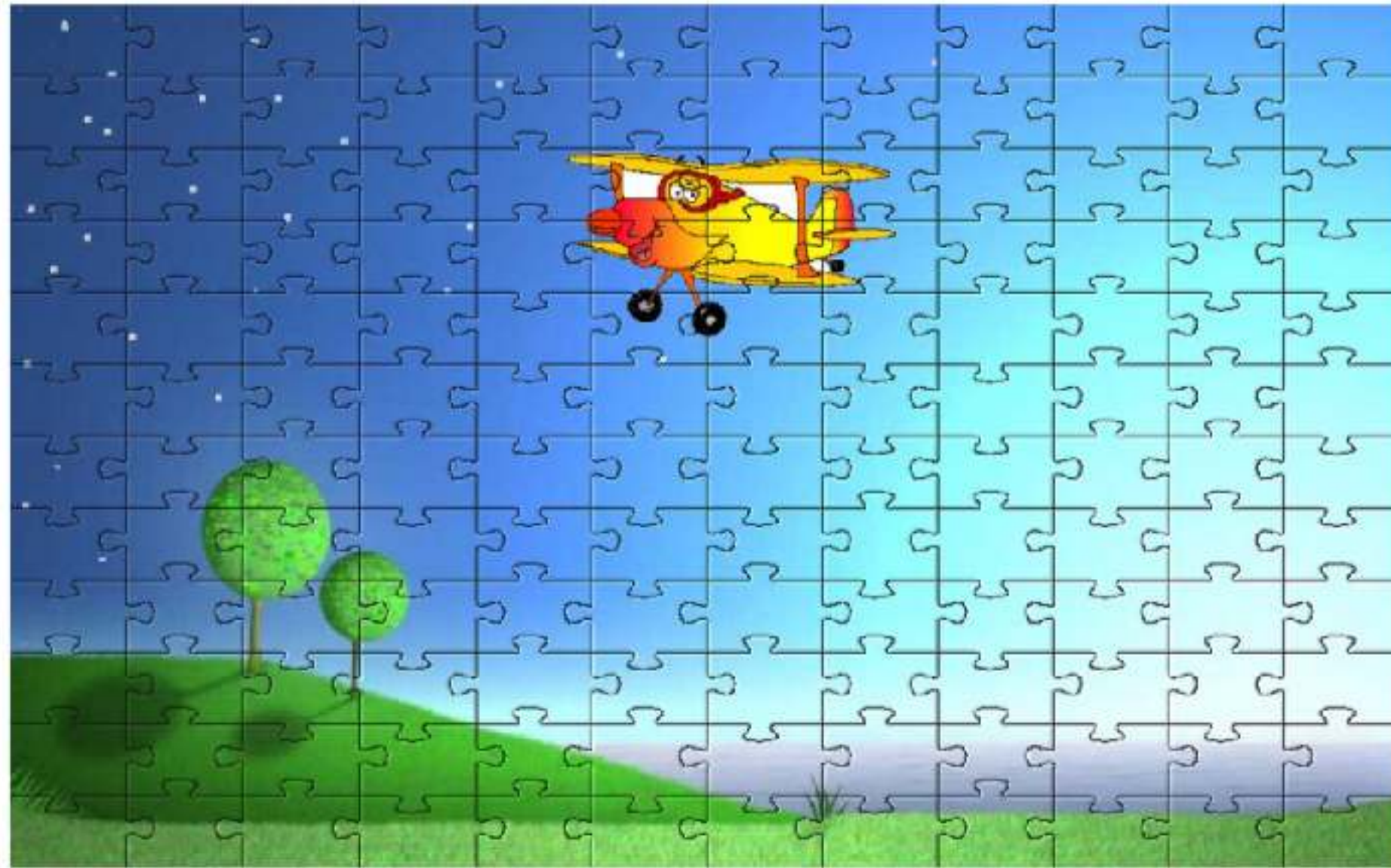


NNNNNN

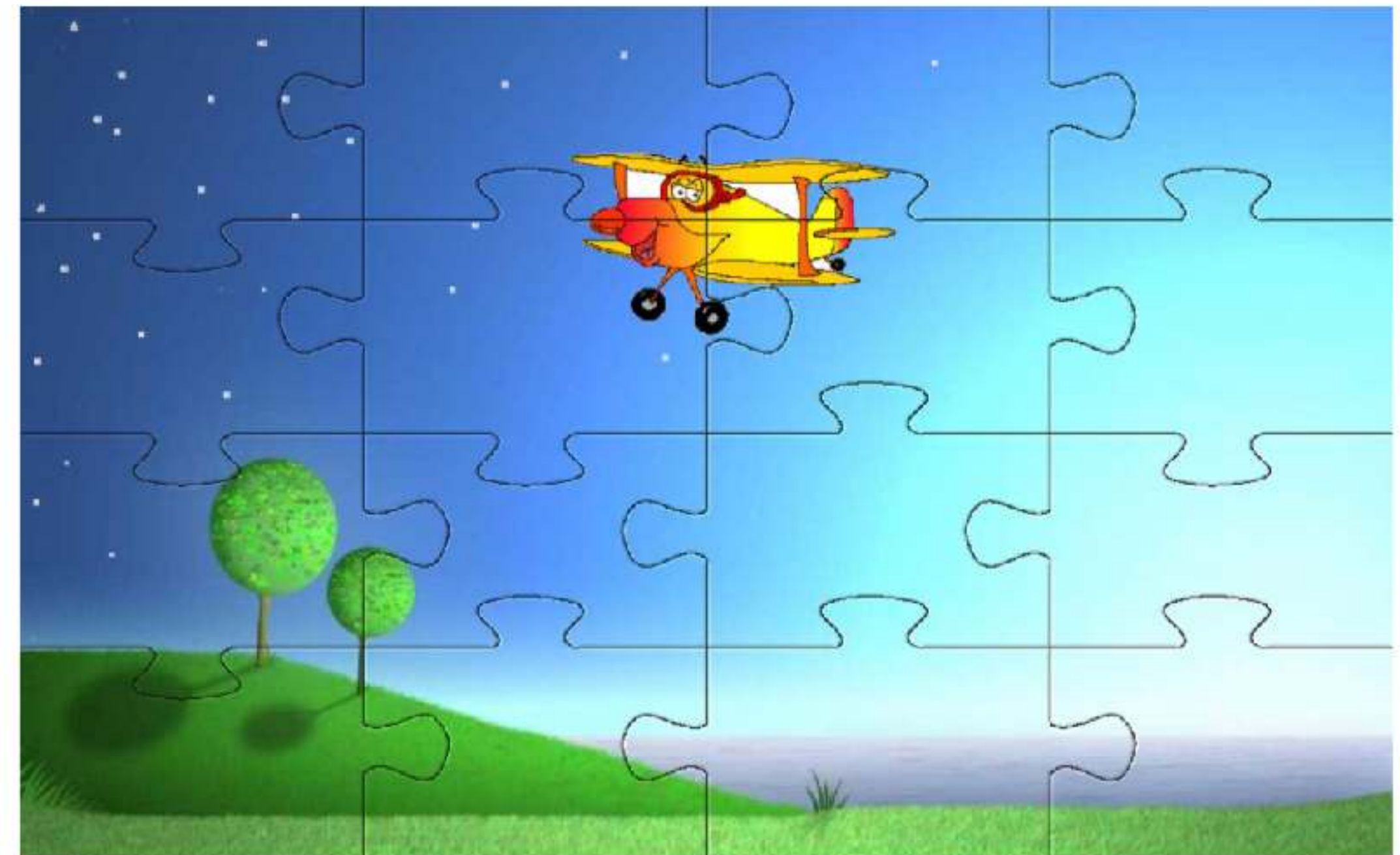


NNNNNN





Assembling with short reads



Assembling with long reads

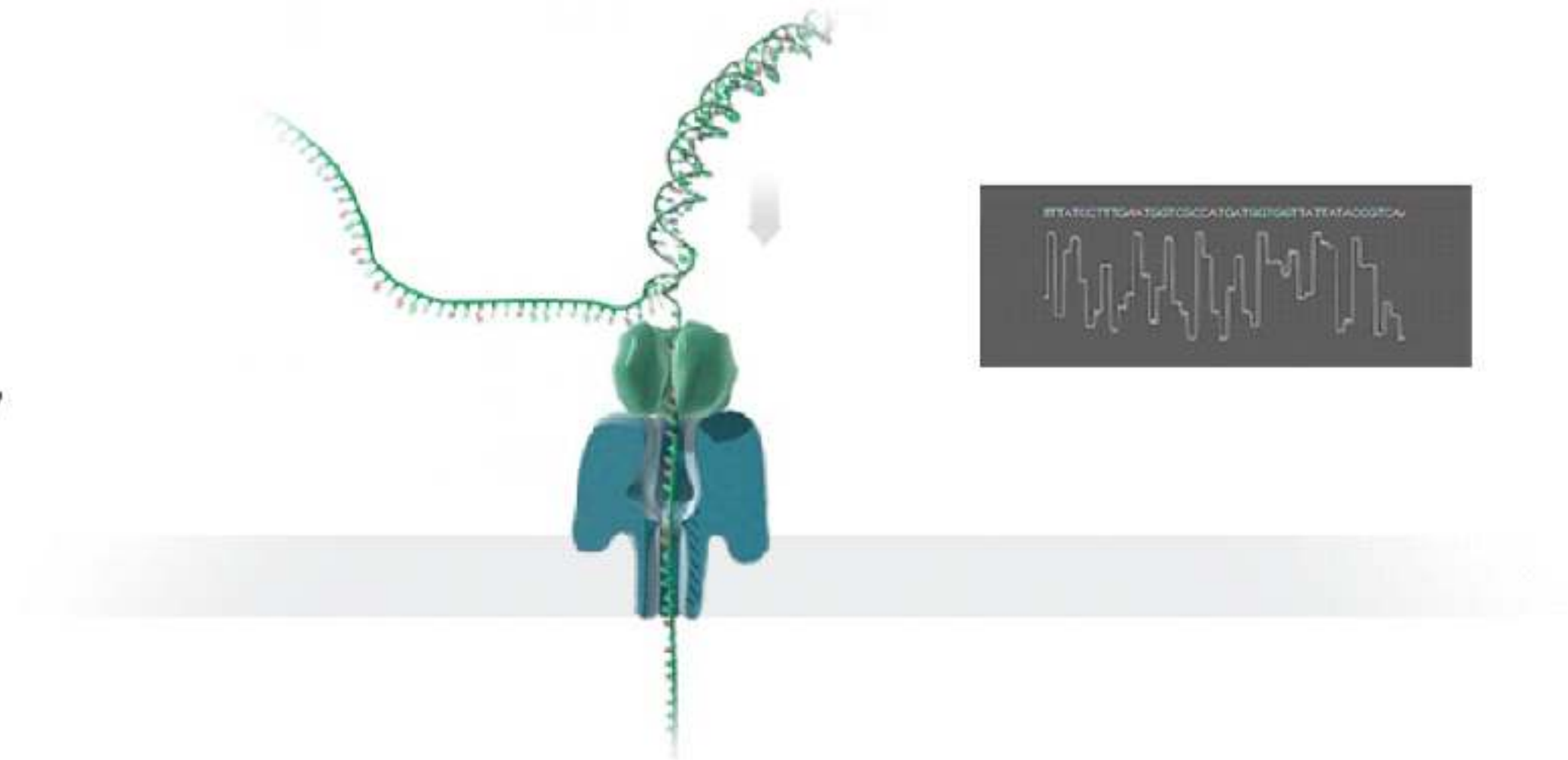
**WHAT SEQUENCING STRATEGY TO
CHOOSE?**

Nanopore ultra-long sequencing



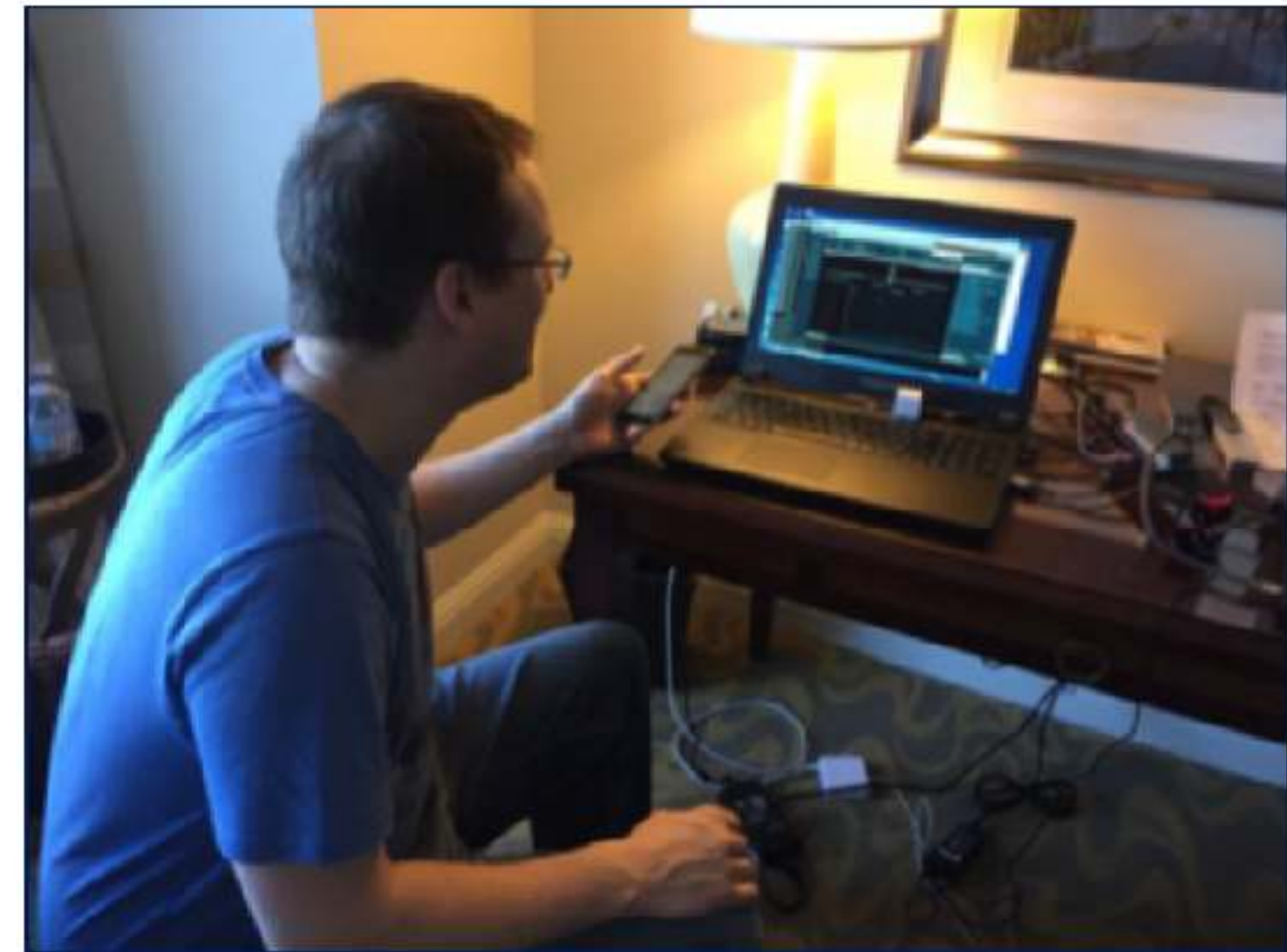
Slide Sergey Koren

- **Nanopore UL**
 - >100 kb reads, up to 1 Mb
 - 97% (Q15) read quality
 - 99.99% (Q40+) assembly quality
- **Pros**
 - Length and throughput
 - Reads *span* repeats
- **Cons**
 - Lower base quality



Nanopore sequencing and assembly of a human genome with ultra-long reads.
Jain et al. *Nature Biotechnology* (2018)

Nanopore sequencing and the Shasta toolkit enable efficient de novo assembly of eleven human genomes. Shafin et al. *Nature Biotechnology* (2020)



PacBio circular consensus sequencing

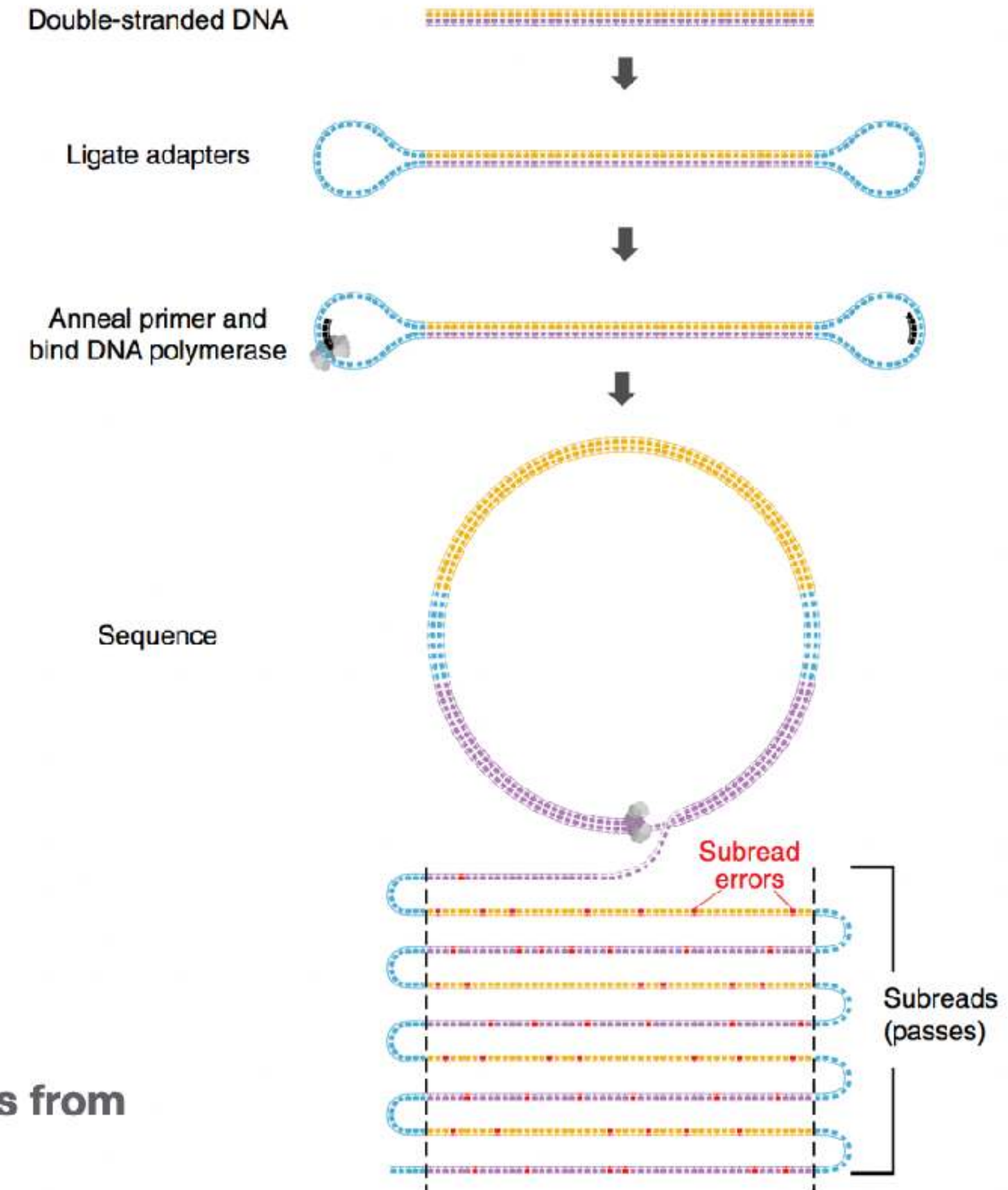


Slide Sergey Koren

- **PacBio HiFi**
 - 20 kb reads
 - 99.9% (Q30) read quality
 - 99.9999% (Q60+) assembly quality
- **Pros**
 - Near-perfect accuracy
 - Reads *distinguish* repeats
- **Cons**
 - Limited length and coverage

Accurate circular consensus long-read sequencing improves variant detection and assembly of a human genome. Wenger et al. *Nature Biotechnology* (2019)

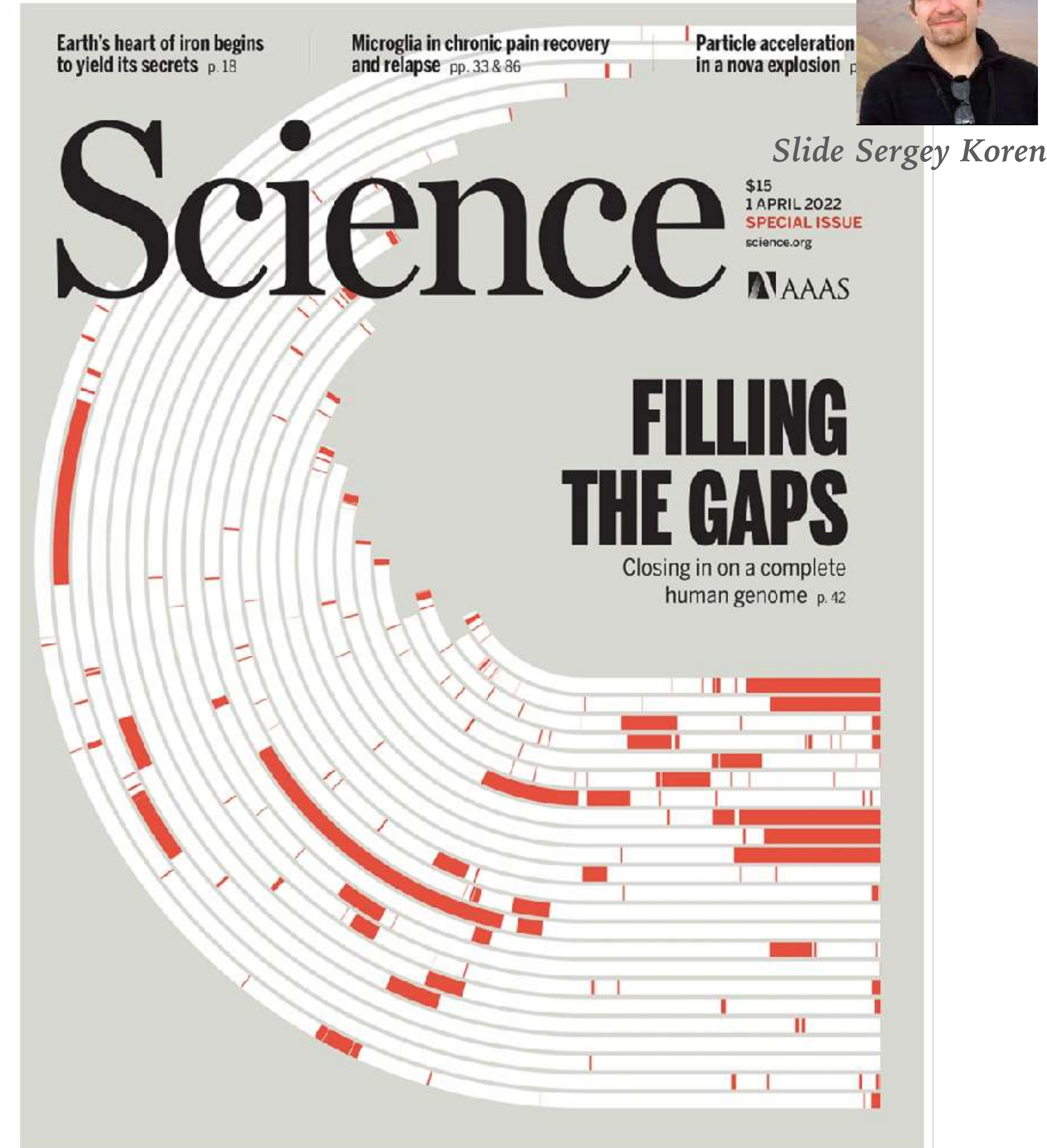
HiCanu: accurate assembly of segmental duplications, satellites, and allelic variants from high-fidelity long reads. Nurk et al. *Genome Research* (2020)



The best of both worlds

Telomere-to-Telomere

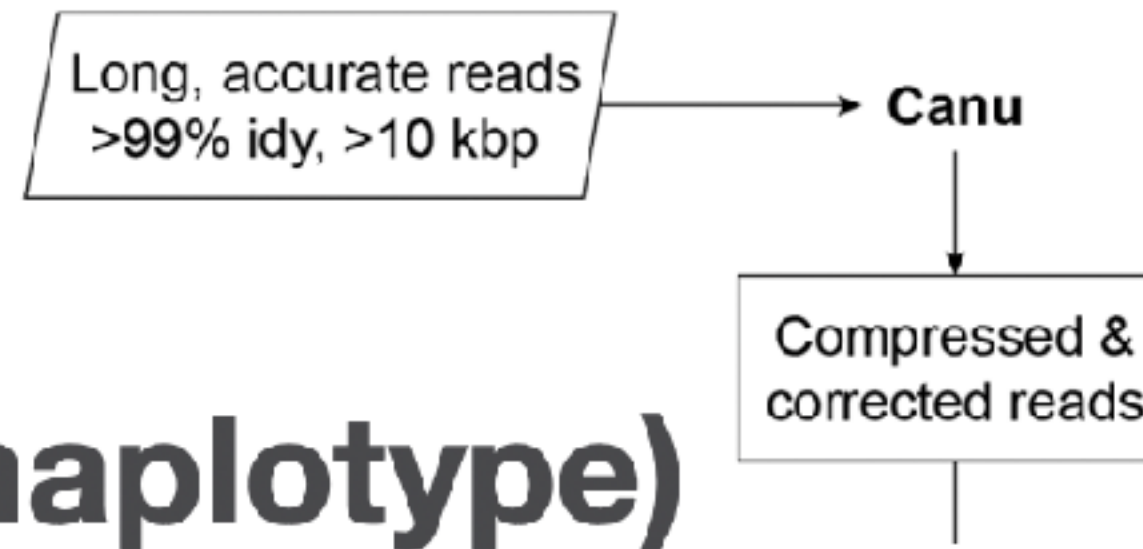
- The human genome is finally finished!
- 8% was left after HGP
- Solved with combination of HiFi + ultra-long ONT



Verkko!

- **Sequencing recipe (per haplotype)**

- 25 PacBio HiFi (20 kb)
- 25x ONT ultra-long (>100 kb)
- 30x Illumina Trio or Hi-C



The diagram illustrates how Verkko compresses repetitive sequences. It shows three levels of a sequence: 'Uncompressed' (TATTTTATACTCTACATGAAATATCAAA), 'Homopolymer compressed' (TATATACTCTACATGATATCA), and 'Microsatellite compressed' (TACTACATGATCA). Brackets indicate how the long runs of identical bases in the uncompressed sequence are collapsed into single bases or shorter motifs in the compressed versions.

Uncompressed
Homopolymer
compressed
Microsatellite
compressed

Telomere-to-telomere assembly of diploid chromosomes with Verkko
Rautiainen, et al. Nat Biotech (2023)



Slide Sergey Koren

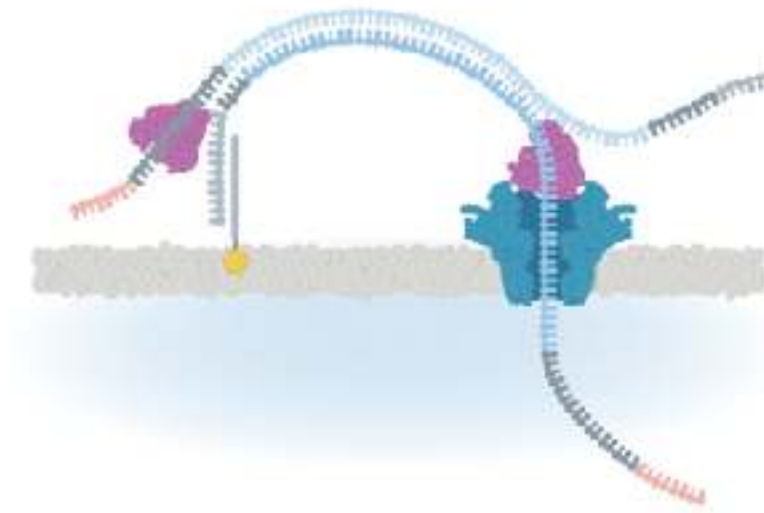
Nanopore duplex sequencing



Slide Sergey Koren

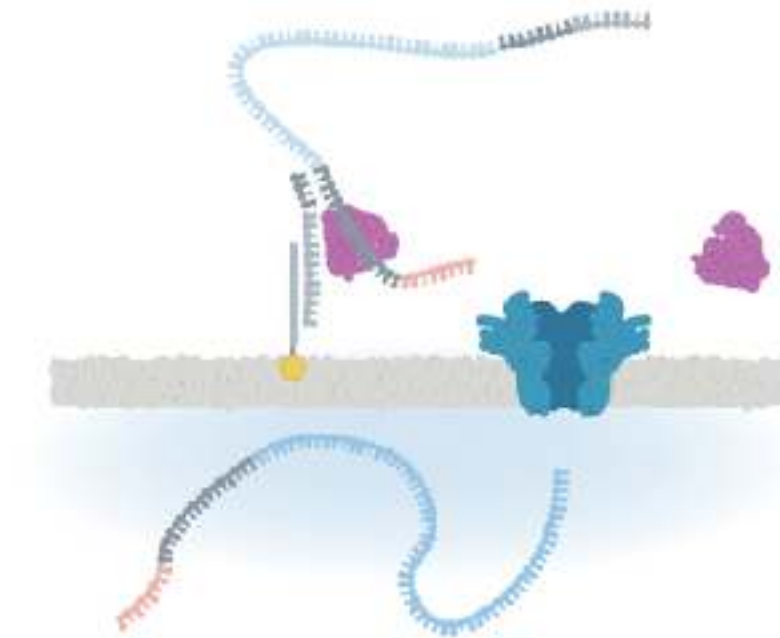
- **Nanopore Duplex**
 - >10 kb reads
 - 99.9% (Q30) read quality
 - 99.999% (Q50+) assembly quality
- **Pros**
 - Near-perfect accuracy
 - No size selection to limit length
 - Reads *distinguish* and *span* repeats
- **Cons**
 - Low throughput

Linear dsDNA molecule adapted on both ends and first strand sequenced

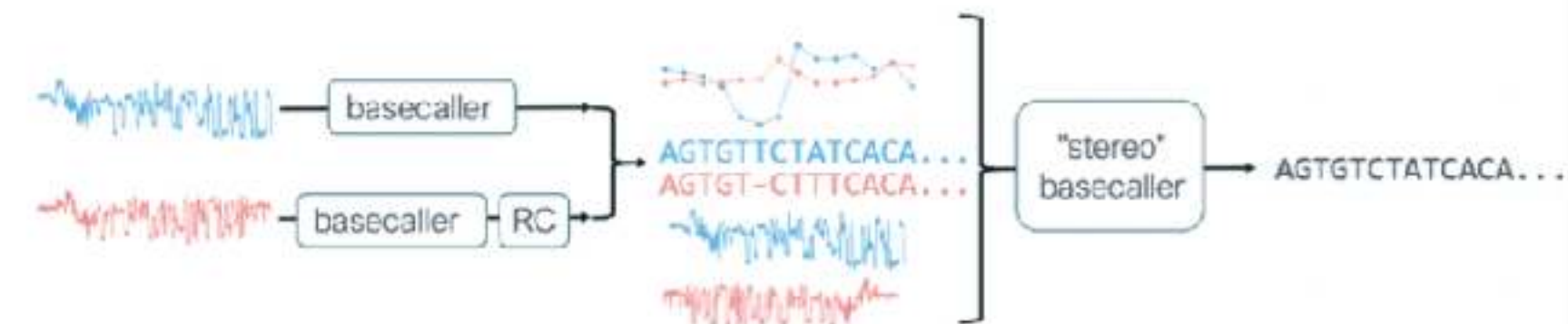


Dorado v0.0.3,
Super accuracy

Second strand captured and sequenced subsequently

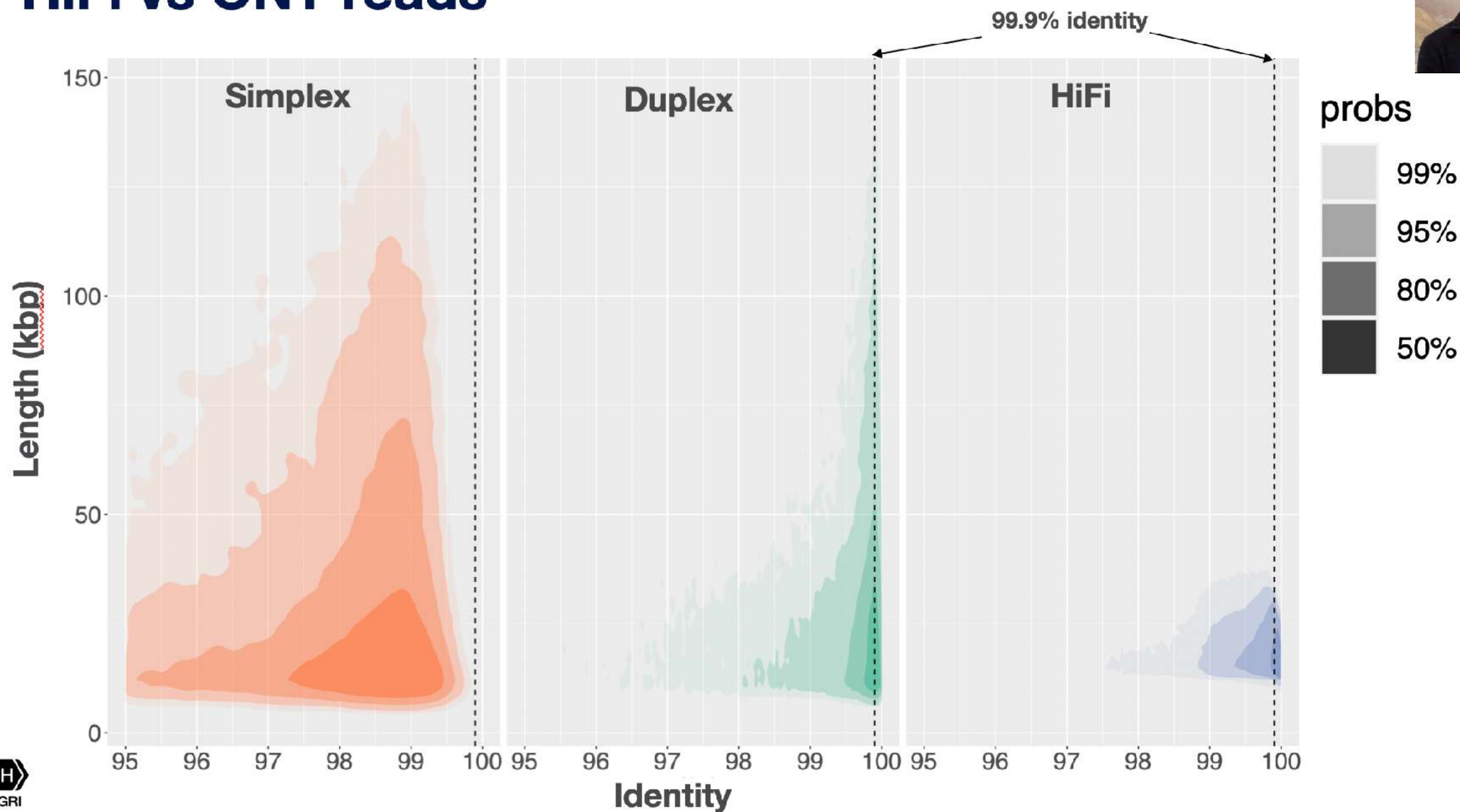


Duplex-tools: Stereo Method
post-basecalling read splitting





HiFi vs ONT reads





PacBio HiFi data at the core of DToL strategy



We generate

- HiFi CCS reads to ~25x
- Hi-C (Illumina) reads to ~100x
- linked reads (Illumina) to ~50x
- RNASeq (30 M Illumina read pairs)

also

- ONT long reads (*for large, repetitive genome*)
- isoSEQ RNASeq (*for gene finding in Family representatives*)



Wellcome Sanger Tree of Life Programme

@SangerToL

...

The first of three @PacBio Revio systems has arrived @sangerinstitute



These will allow us to sequence #genomes for projects like @darwintreelife faster and at reduced cost - with each having up to 15x the throughput of current machines 🧬



Full details: pacb.com/revio



6:00 AM · Mar 22, 2023 · 17.1K Views

ASSEMBLY



Shane, Ksenia, Chenxi, Marcela, Eerik, Noah, James, Yumi, Willian

BREATHE

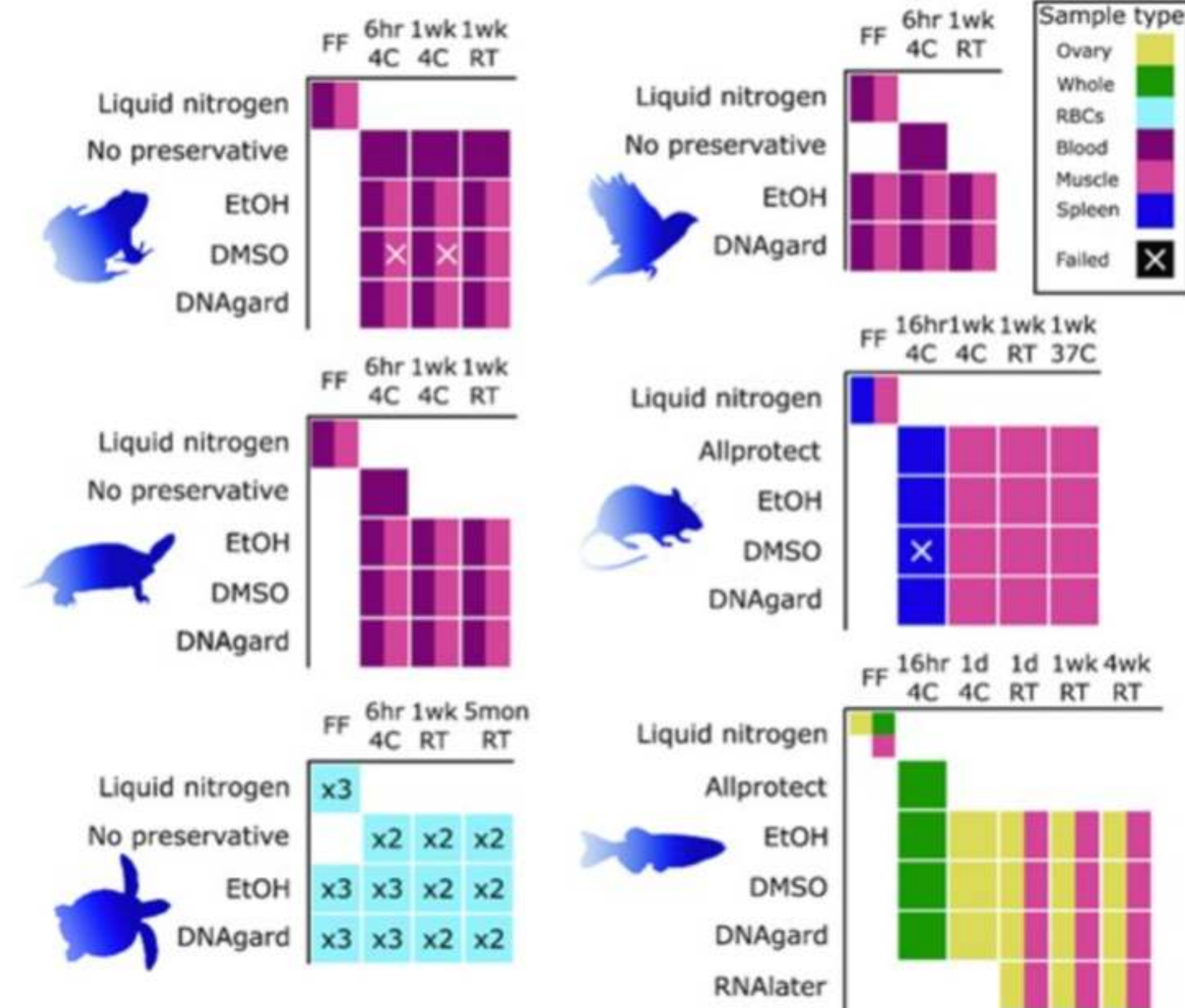
Benchmarking ultra-high molecular weight DNA preservation methods for long-read and long-range sequencing

Hollis A. Dahn^{1,†}, Jacquelyn Mountcastle^{2,†}, Jennifer Balacco², Sylke Winkler³, Iliana Bista^{4,5}, Anthony D. Schmitt⁶, Olga Vinnere Pettersson⁷, Giulio Formenti², Karen Oliver⁴, Michelle Smith⁴, Wenhua Tan³, Anne Kraus³, Stephen Mac⁶, Lisa M. Komoroske⁸, Tanya Lama⁸, Andrew J. Crawford⁹, Robert W. Murphy¹, Samara Brown², Alan F. Scott¹⁰, Phillip A. Morin¹¹, Erich D. Jarvis^{2,12} and Olivier Fedrigo^{2,*}

No one-size-fits-all protocol!

 slack Channel: all.things.up.to.assembly

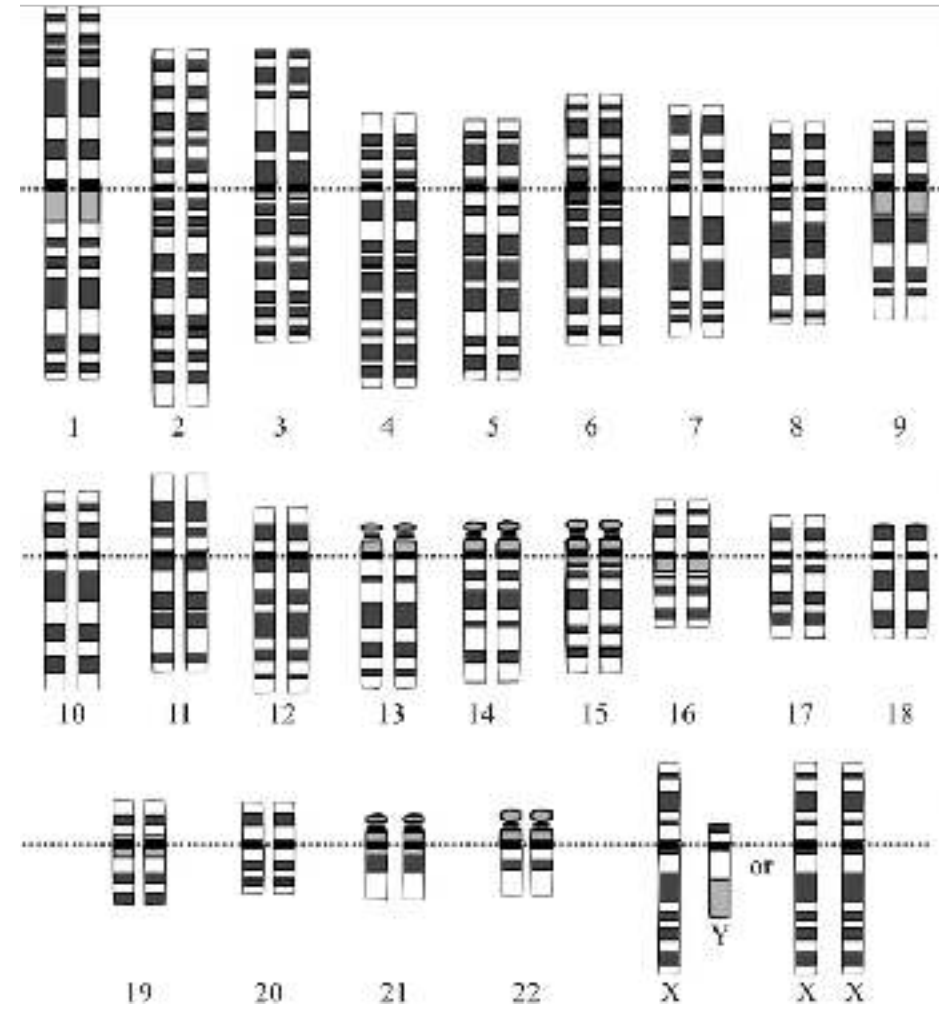
e 1:



TAKE HOME MESSAGE

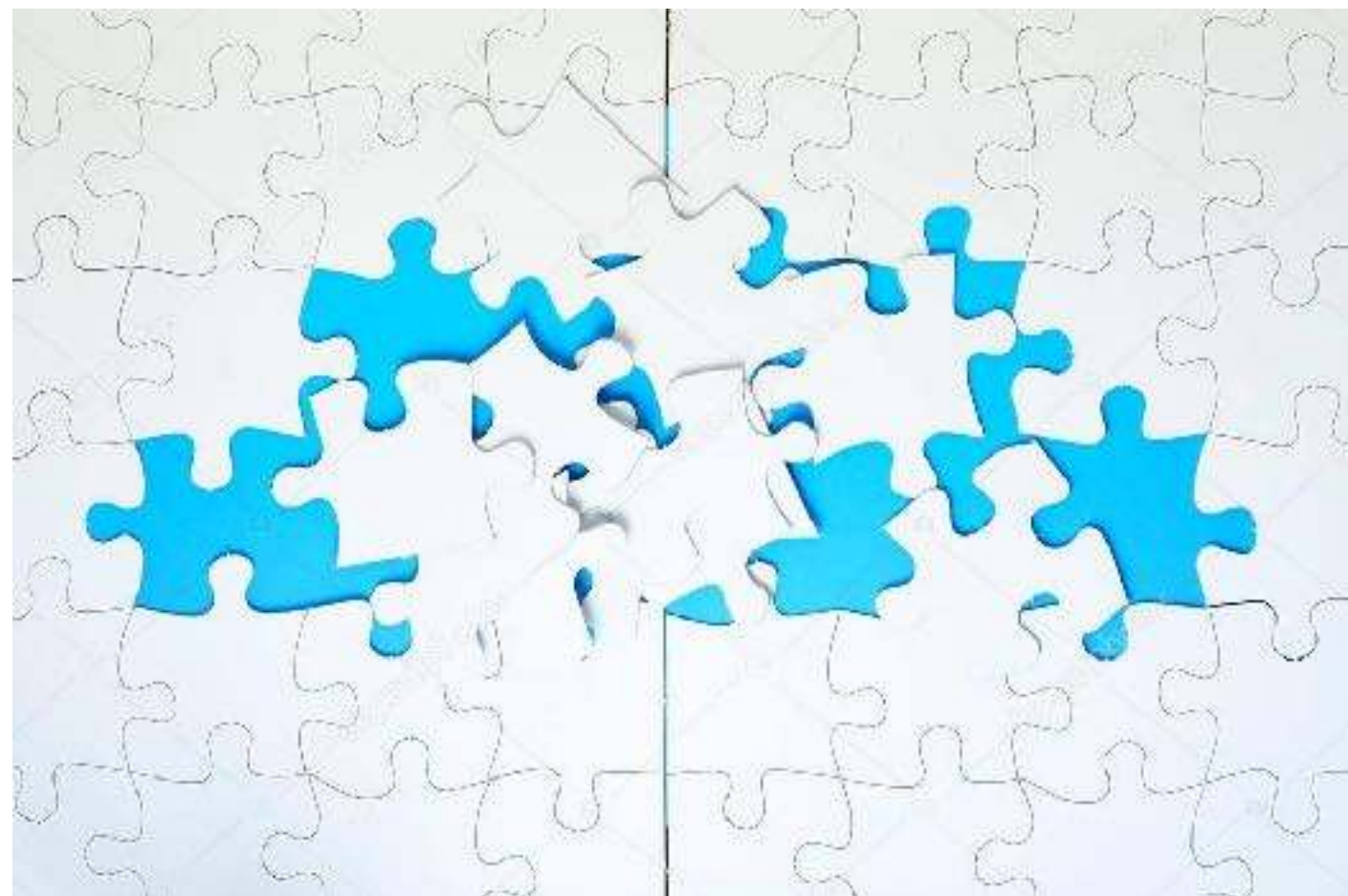
- Know your species! Do your research prior to sequencing. Try to have the best sample you can (fresh and immediately flash-frozen).
- **Estimated genome size, repeat content, heterozygosity**
- From our experience at the Darwin Tree of Life
- 25x coverage of PacBio HiFi (for both haplotypes) + 100x coverage of Hi-C is yielding high-quality assemblies
- Nanopore duplex is promising: not available to the public yet

WHEN WE ASSEMBLE A GENOME ...



What we would like to have

- *One DNA sequence for each chromosome*



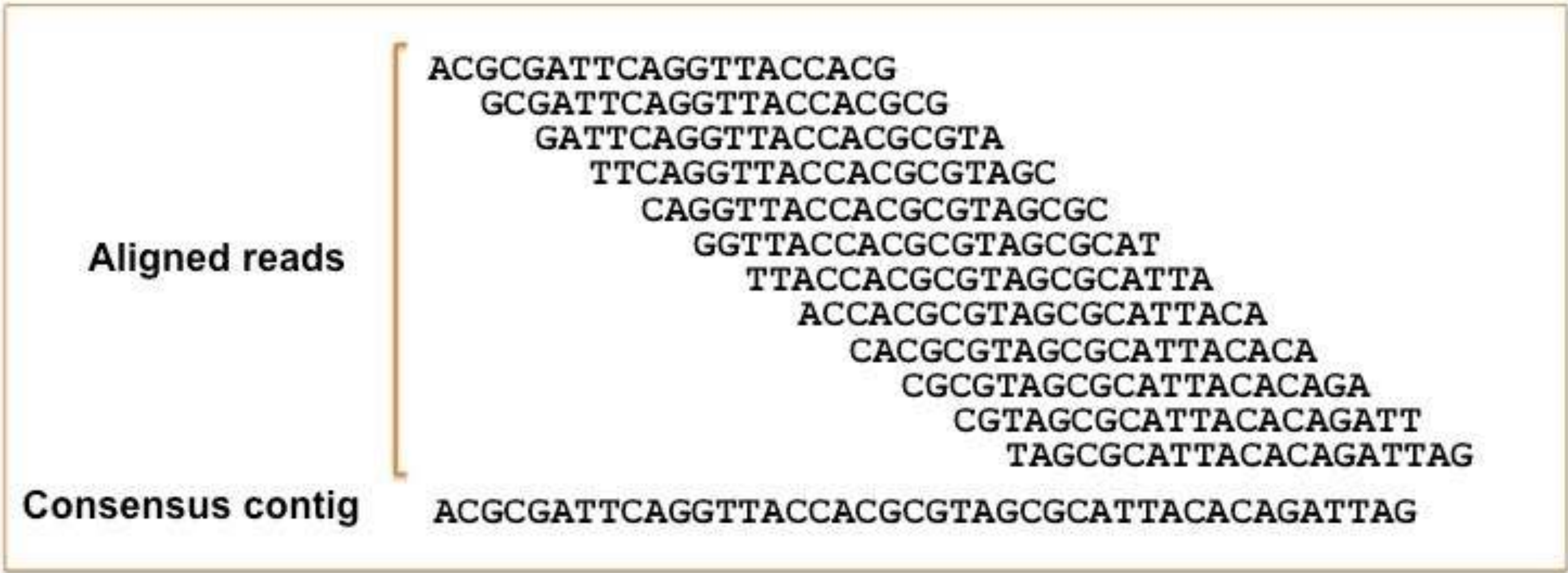
What we really have

- *Contigs, scaffolds, gaps, N50s*

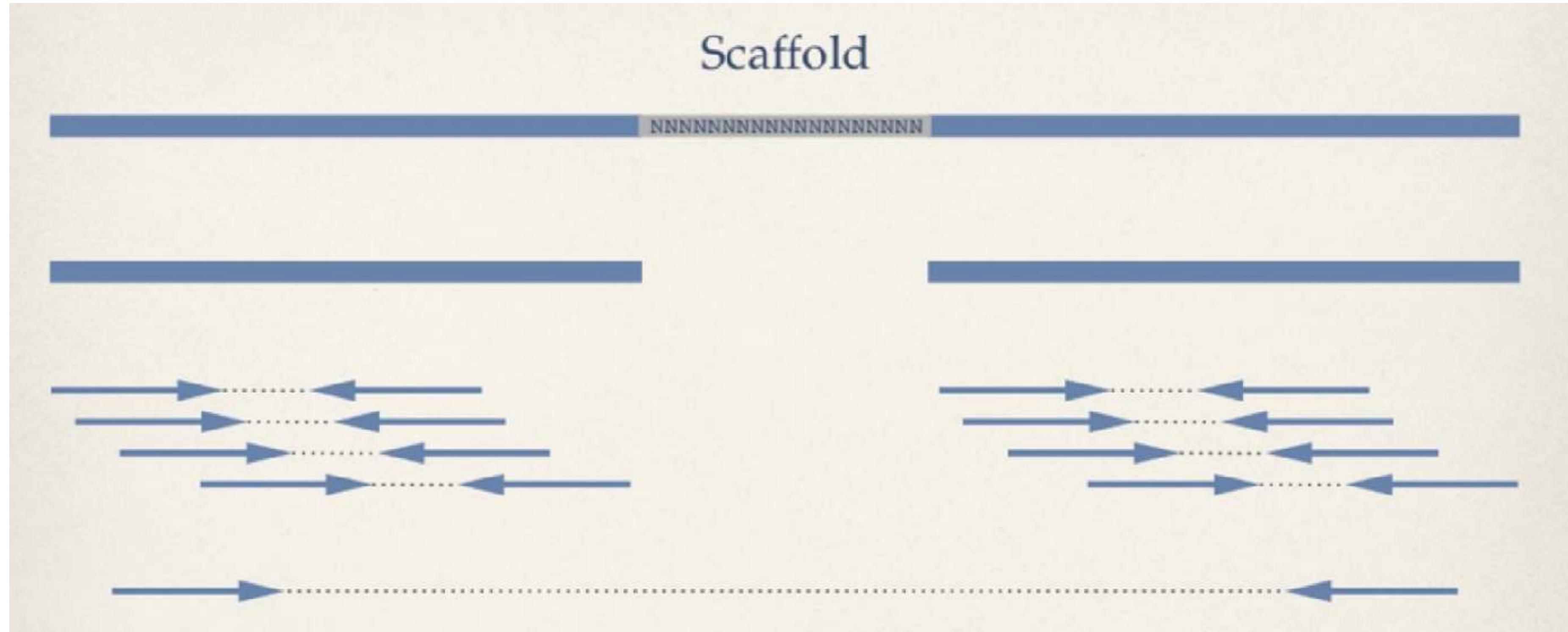
GENOME ASSEMBLY METRICS

A DNA sequence with gaps

CONTIG



Scaffolding methods

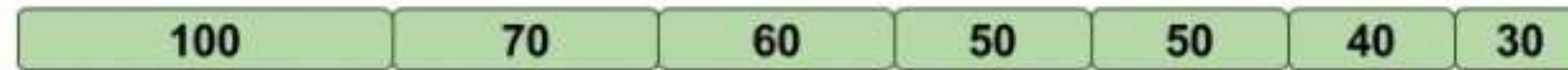


Scaffold: joining and orienting contigs

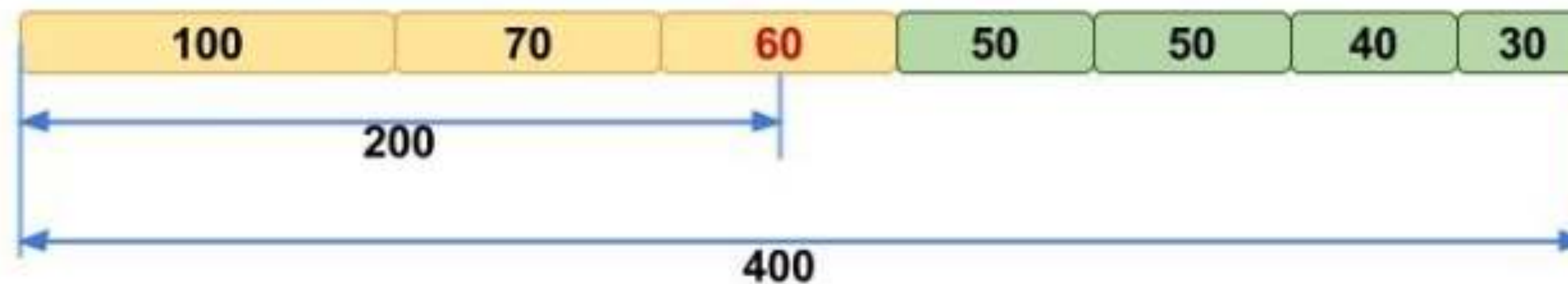
Scaffolding methods: mate-pairs (blerg), optical maps (bionano), Hi-C, Nanopore UltraLong reads

N50

- **N50: half of the genome is assembled in scaffolds that are the N50 size, or larger**



1a. Contigs, sorted according to their lengths.



1b. Calculation of N50 using sorted contigs.

Fig. 1. Example of calculating N50 for a set of seven contigs. Here N50 equals 60 kbp.

Quality metrics in genomics

- **N50: half of the genome is assembled in scaffolds that are the N50 size, or larger**

Chr Number	Chr Size	Accum Size	Genome % Coverage
1	197.61	197.61	18.82%
2	149.68	347.29	33.07%
3	110.84	458.13	43.63%
4	91.32	549.45	52.32%
Z	82.53	631.98	60.18%
5	59.81	691.79	65.88%
7	36.74	728.53	69.37%
6	36.37	764.9	72.84%
8	30.22	795.12	75.71%
9	24.15	819.27	78.01%
10	21.12	840.39	80.03%
12	20.39	860.78	81.97%
11	20.2	880.98	83.89%
13	19.17	900.15	85.72%
14	16.22	916.37	87.26%
20	13.9	930.27	88.58%
15	13.06	943.33	89.83%
18	11.37	954.7	90.91%
17	10.76	965.46	91.94%
19	10.32	975.78	92.92%
27	8.08	983.86	93.69%
33	7.82	991.68	94.43%
21	6.84	998.52	95.08%
W	6.81	1005.33	95.73%
24	6.49	1011.82	96.35%
23	6.15	1017.97	96.94%
31	6.15	1024.12	97.52%
26	6.06	1030.18	98.10%
22	5.46	1035.64	98.62%
28	5.12	1040.76	99.11%
25	3.98	1044.74	99.48%
16	2.84	1047.58	99.76%
30	1.82	1049.4	99.93%
32	0.73	1050.13	100.00%
MT	0.02		
Total	1050.15		



Scaffold N50
@ Chromosome level



N50 = 91Mb

Assembled size= 1Gb

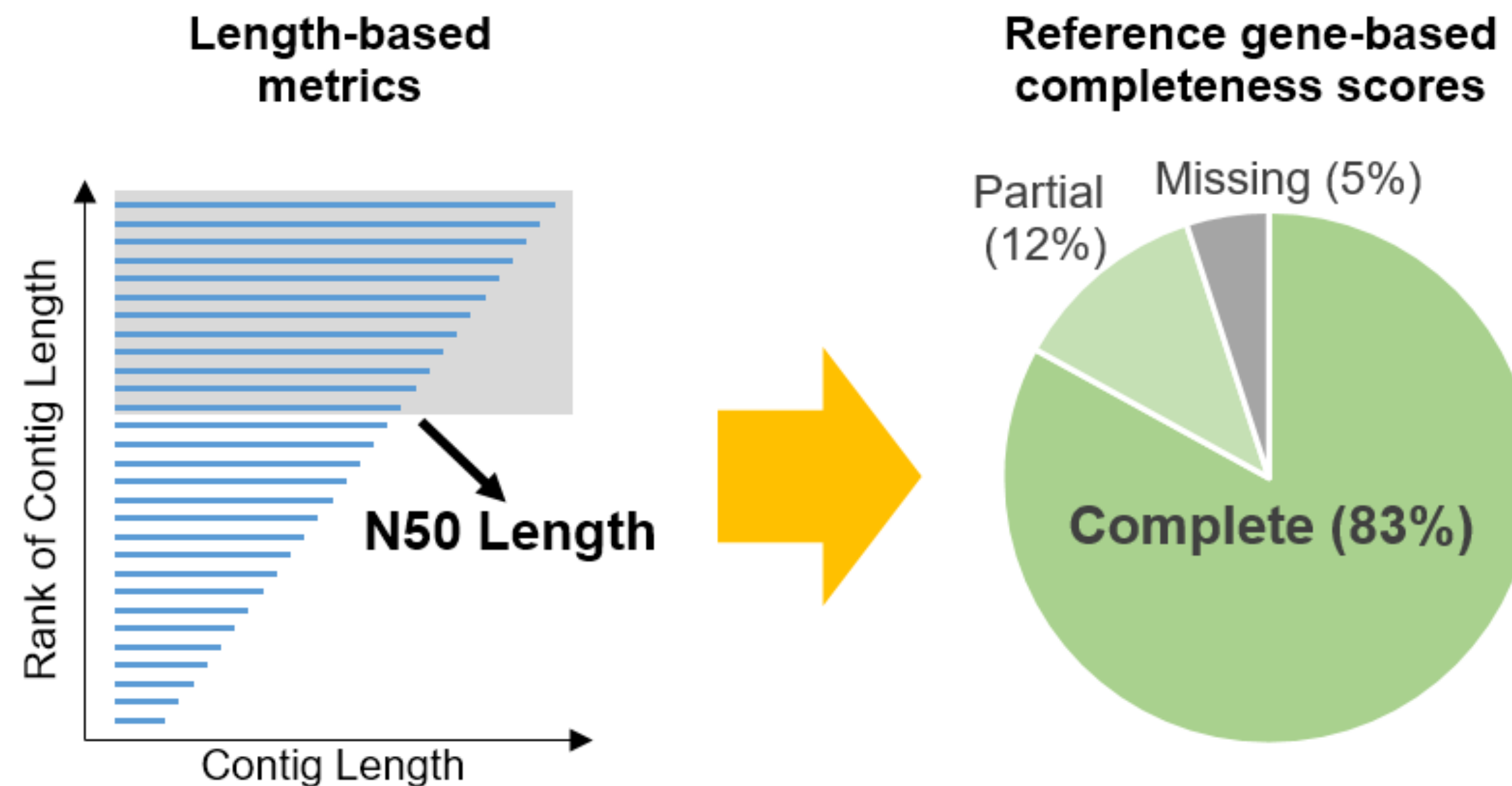
How many scaffolds= 32



Assessing genome assembly and annotation completeness with
Benchmarking **U**niversal **S**ingle-**C**opy **O**rthologs

- The quality metrics for genome assembly should not be only the ones related to contiguity, rather, the composition of the genes present in the assembly is also crucial

More accurate assessment for genome assembly!



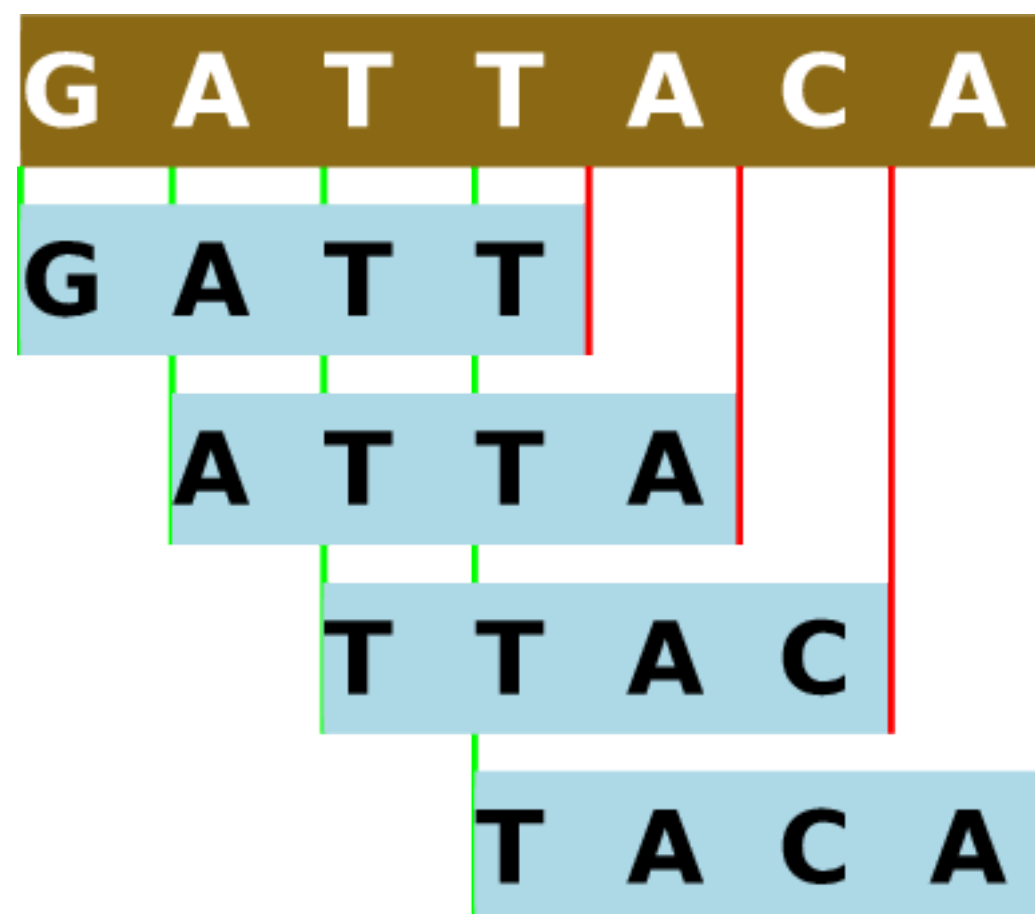
OUR LAB TODAY!

- Count Kmers
- Run general reads statistics
- Run genome assembly with Hifiasm
- Run MitoHiFi to assemble and annotate a mitochondrial genome!





KMER ANALYSIS



WHAT ARE K-MERS ?

- In biology, k-mers are unique subsequences of a sequence of length k

So, by way of example, the sequence **ATCGATCAC** contains the following *3-mers* (*k-mer* of size 3):

Sequence: ATCGATCAC

3-mer #0: ATC

3-mer #1: TCG

3-mer #2: CGA

3-mer #3: GAT

3-mer #4: ATC

3-mer #5: TCA

3-mer #6: CAC

APPLICATIONS OF K-MER ANALYSIS

- Genome assembly: K-mers used to construct De Bruijn graphs
- Detect bacterial contamination on eukaryotic genome assembly (CG content discrepancies)
- Correcting NSG data
- Detect horizontal gene transfers
- Identification of CpG Islands
- Estimation of genome size and heterozygosity
- Genome assembly k-mer completeness

WHY ARE K-MERS SO POPULAR?

“Decomposing a sequence into its *k-mers* for analysis allows this set of fixed-size chunks to be analysed rather than the sequence, and this can be more efficient.” (Bernardo Cavijo)

<https://bioinfologics.github.io/post/2018/09/17/k-mer-counting-part-i-introduction/>

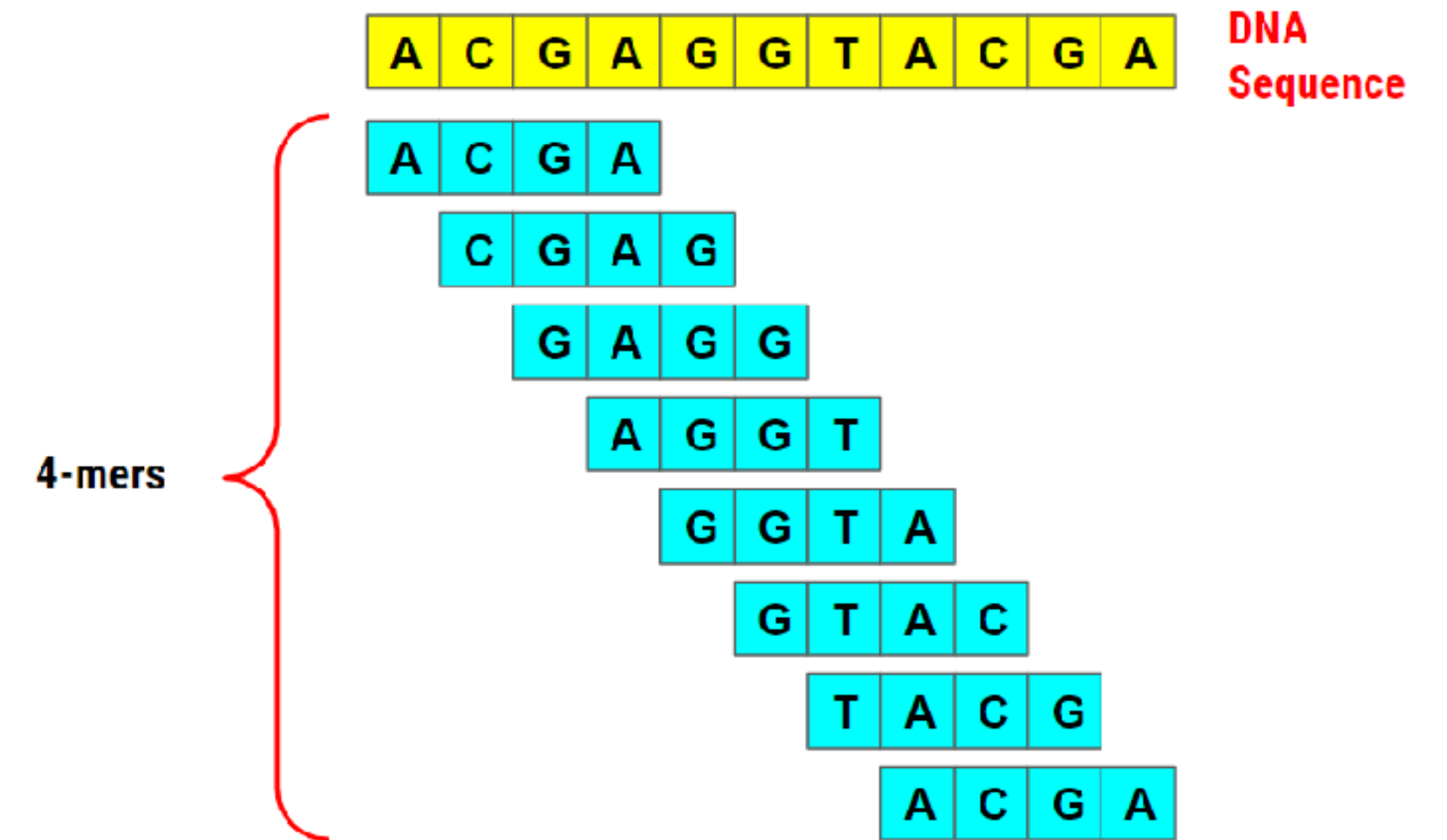
KMER SIZE

Choosing k : specificity vs. Sensitivity

- Using a k that is too small will result in many unrelated sequences being composed of the same k -mers, in a textbook case of specificity loss because there being very few possible k -mers of that size. In the extreme of the small k , $k=1$ only distinguishes two *canonical k-mers*: A and C. 1-mer analysis is incredibly popular in biology, but it is best known by the name of *GC content analysis*.
- Using extremely large k values would sacrifice many of the benefits and sensitivity of k -mer analyses in the first place. (Bernado Caviyo's post)

Why do we chose $k=31$ so often?

One reason is: it is specific enough that a large number of them are unique both in mammalian-sized genomes and in bacterial genome databases.



COUNT AND HISTO

Counting *k*-mers in a (small) genome

We will start with an easy example first: the [phi-X174 genome](#) has 5386 bp and is a simple non-repetitive genome.

We can use `kat hist` to count *27-mers* on the genome and check how many times each *27-mer* appears (we start with `k = 27` because KAT uses that as default):

```
$ kat hist -o phiX.hist phiX.fasta
```

Checking the `phiX.hist` histogram (A.K.A. kmer spectrum) file, every *27-mer* in the genome appears only once. After the header lines starting with `#`, every line has a copy number (A.K.A. frequency) and a number of *k-mers*.

```
# Title:27-mer spectra for: phiX.fasta
# XLabel:27-mer frequency
# YLabel:# distinct 27-mers
# Kmer value:27
# Input 1:../genomes/phiX.fasta
###
1 5360
2 0
3 0
4 0
...
```

COUNT AND HISTO

```
$ kat hist -o phiX_9mer.hist -m 9 phiX.fasta
```

Then the `phiX_9mer.hist` file looks like this:

```
# Title:9-mer spectra for: phiX.fasta
# XLabel:9-mer frequency
# YLabel:# distinct 9-mers
# Kmer value:9
# Input 1:phiX.fasta
###
1 4972
2 189
3 8
4 1
5 0
6 0
7 0
8 0
9 0
...
```

```
$ kat hist -o phiX_8mer.hist -m 8 phiX.fasta
```

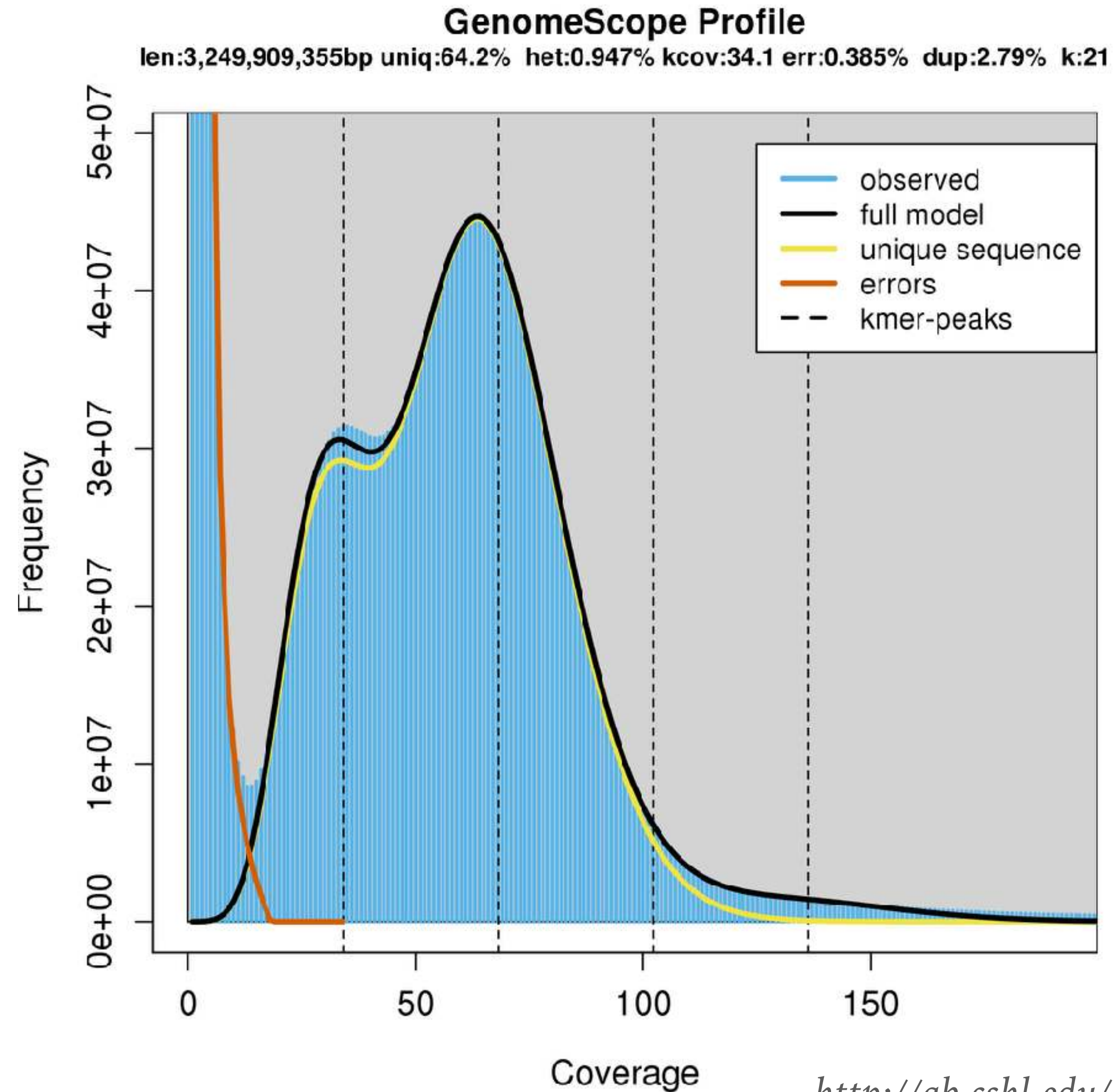
Now the histogram file looks like this:

```
# Title:8-mer spectra for: phiX.fasta
# XLabel:8-mer frequency
# YLabel:# distinct 8-mers
# Kmer value:8
# Input 1:phiX.fasta
###
1 4159
2 491
3 67
4 8
5 1
6 0
7 0
8 0
9 0
```

Here, only **4159** *8-mers* are *unique*, out of **4726** *distinct 8-mers*, that are present in the genome's **5377** *total 8-mers*.

Bernardo Cavijo's post

A TYPICAL KMER PLOT FOR A DIPLOID SPECIES



Choloepus didactylus (VGP)



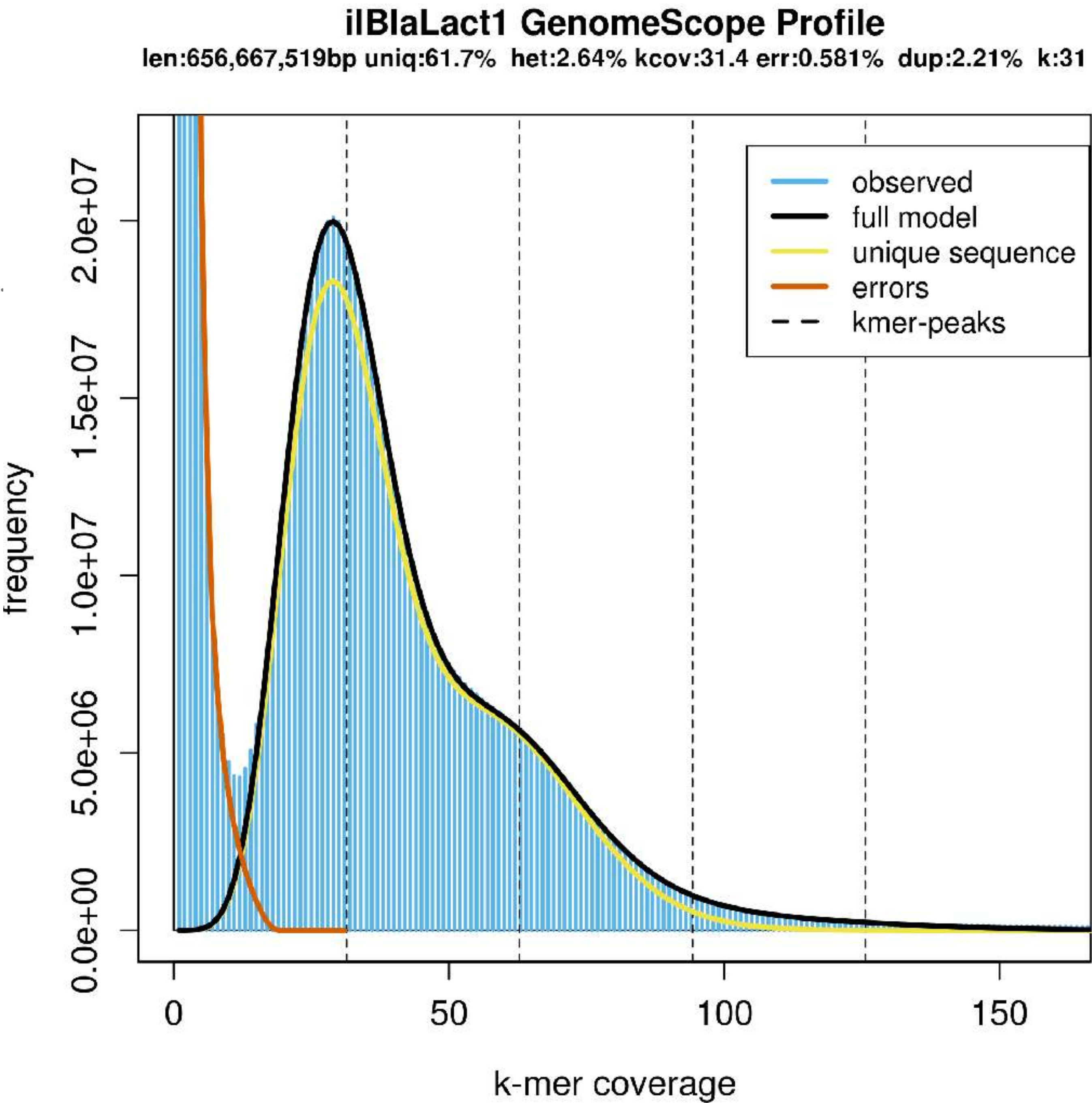
<http://qb.cshl.edu/genomescope/analysis.php?code=bVuZNLhwn2tVCHhRN71I>

A TYPICAL KMER PLOT FOR A DIPLOID SPECIES WITH HIGH HETEROZYGOSITY

Blastobasis lacticolella (DToL)

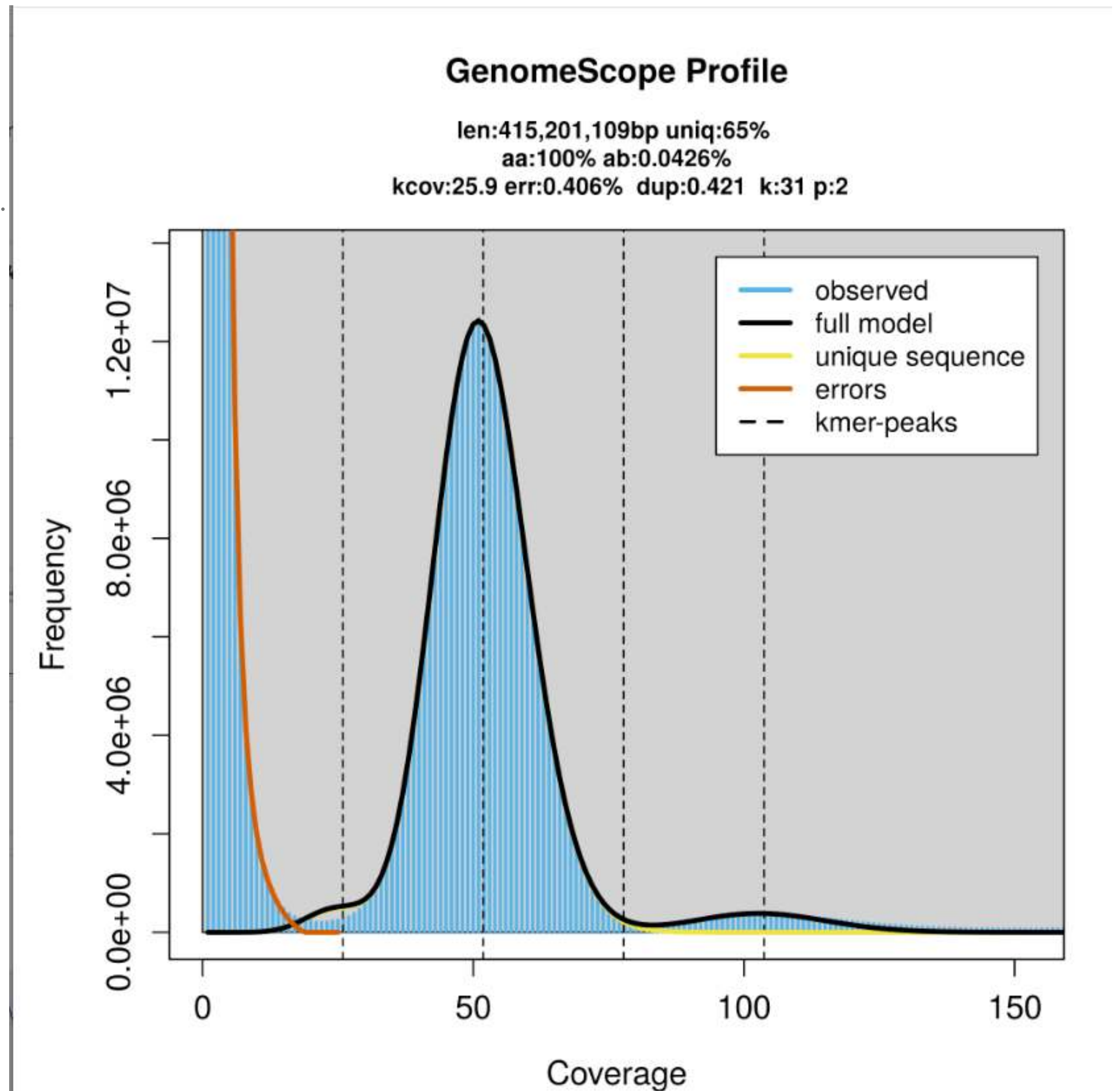


Wakely's dowd



A TYPICAL KMER PLOT FOR A DIPLOID SPECIES WITH LOW HETEROZYGOSITY

Rhytidiadelphus loreus
Little Shaggy-moss



KMERS CAN BE ANALYSED ONLY FOR HIGH-QUALITY DATA

This means that:

- *If you have sequenced PacBio CLR, you should have short-read sequencing for kmer analysis (and for polishing)*
- *But with Pacbio HiFi that you can count kmers as you do with short-reads!*





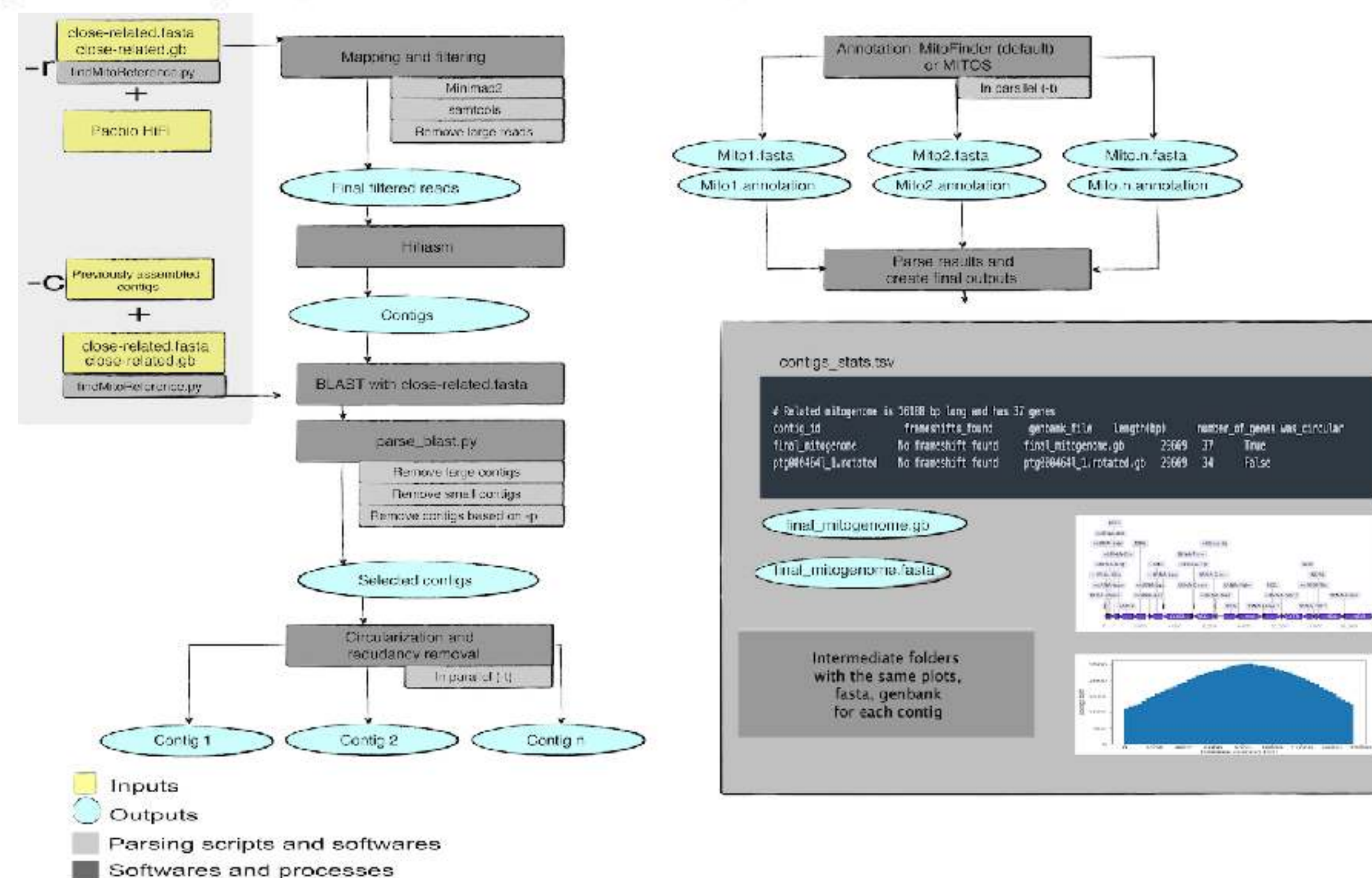
JORGE CHAM © 2005

www.phdcomics.com

MitoHiFi: a python pipeline for mitochondrial genome assembly from PacBio High Fidelity reads

Marcela Uliano-Silva, João Gabriel R. N. Ferreira, Ksenia Krasheninnikova, Darwin Tree of Life Consortium, Giulio Formenti, Linelle Abueg, James Torrance, Eugene W. Myers, Richard Durbin, Mark Blaxter, Shane A. McCarthy

doi: <https://doi.org/10.1101/2022.12.23.521667>

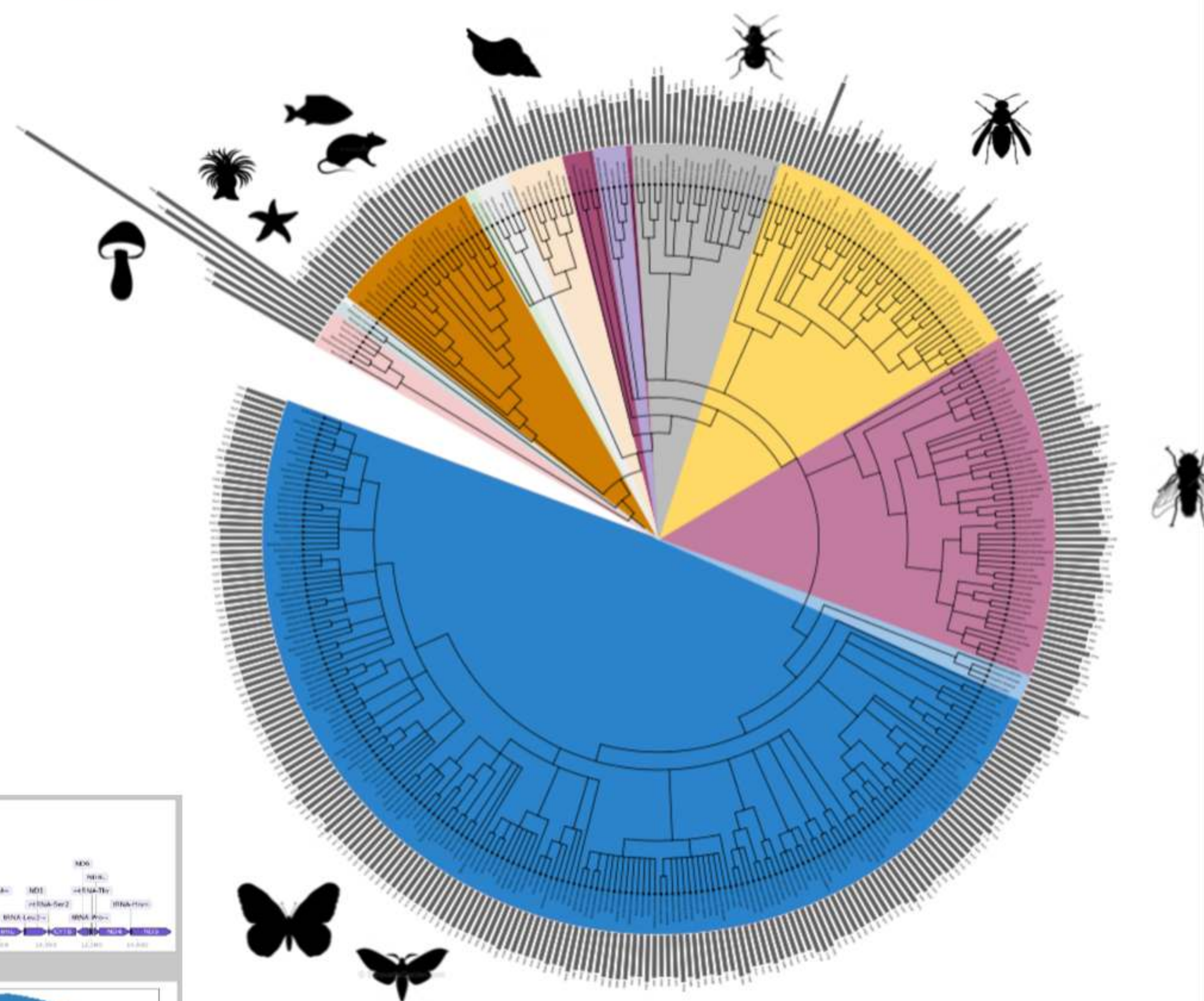
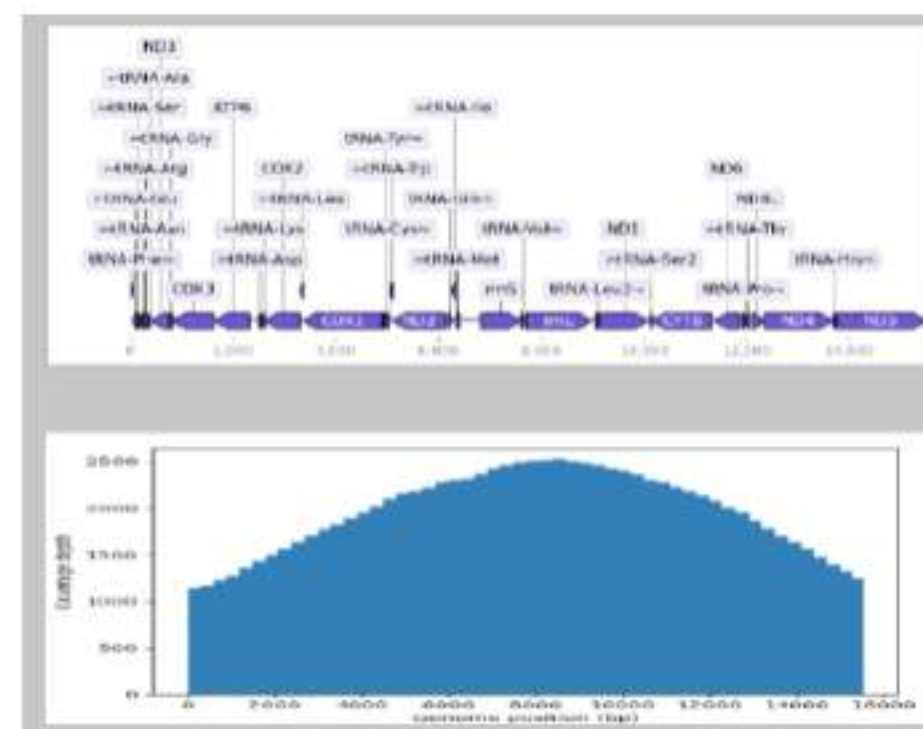
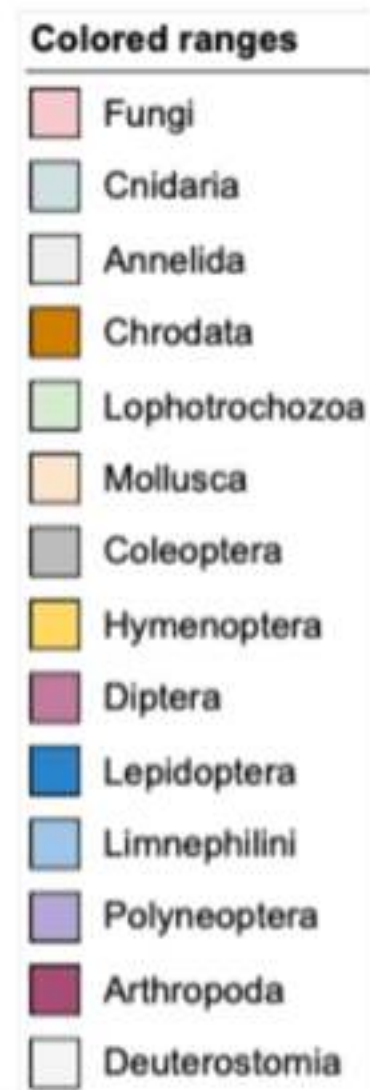
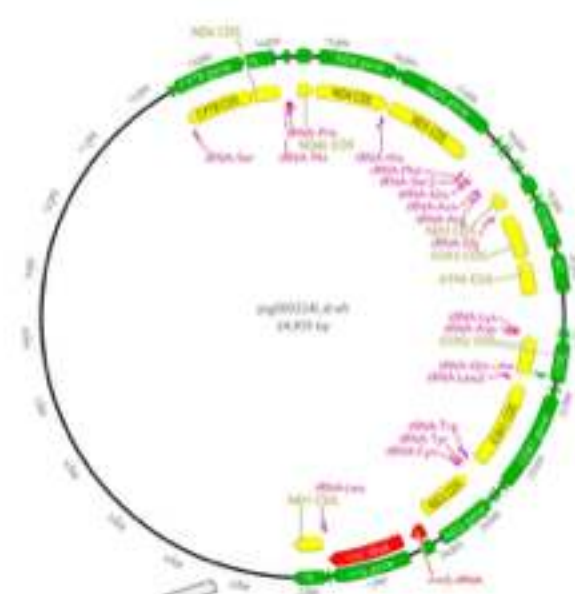
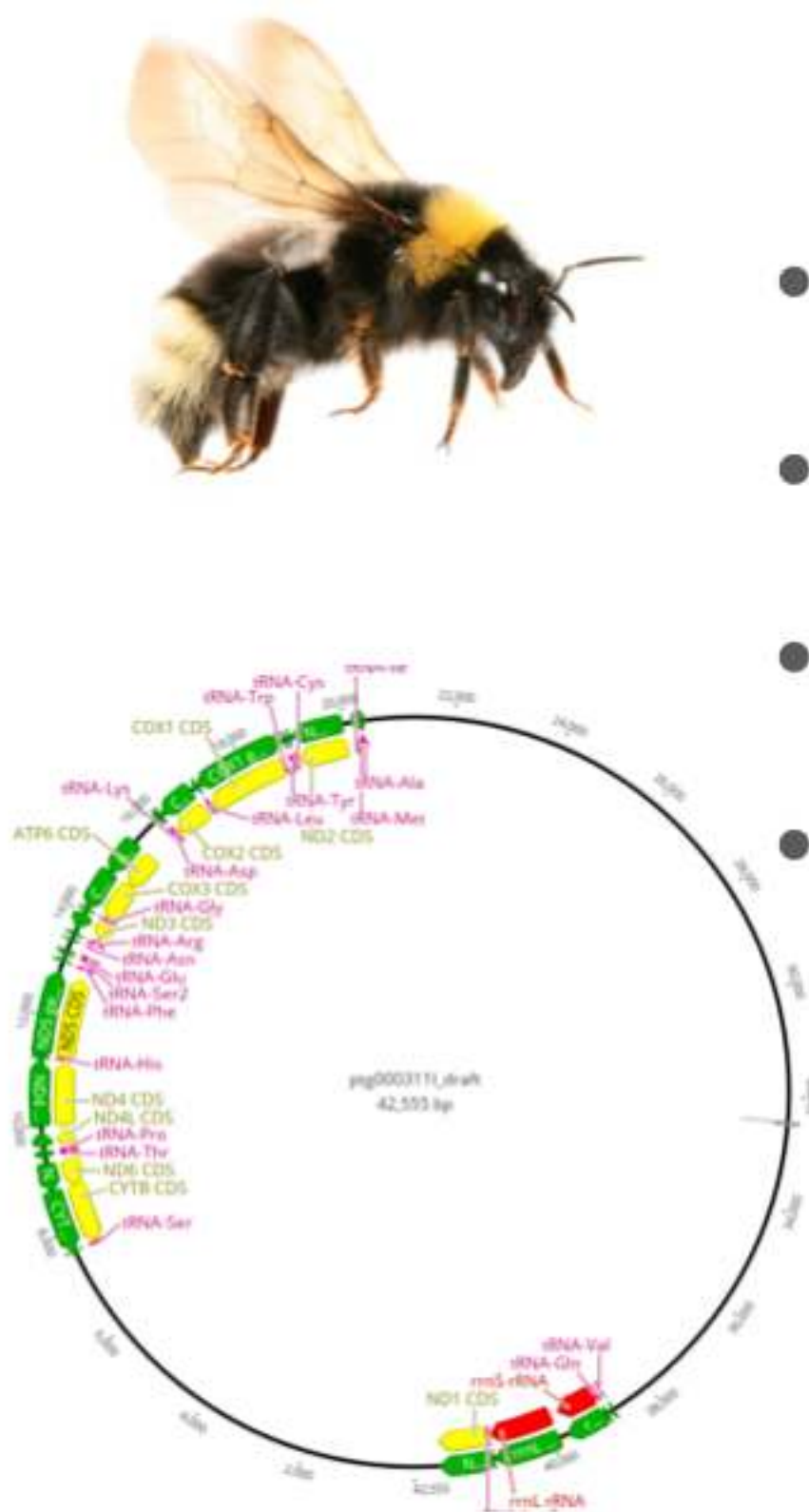


MitoHifi



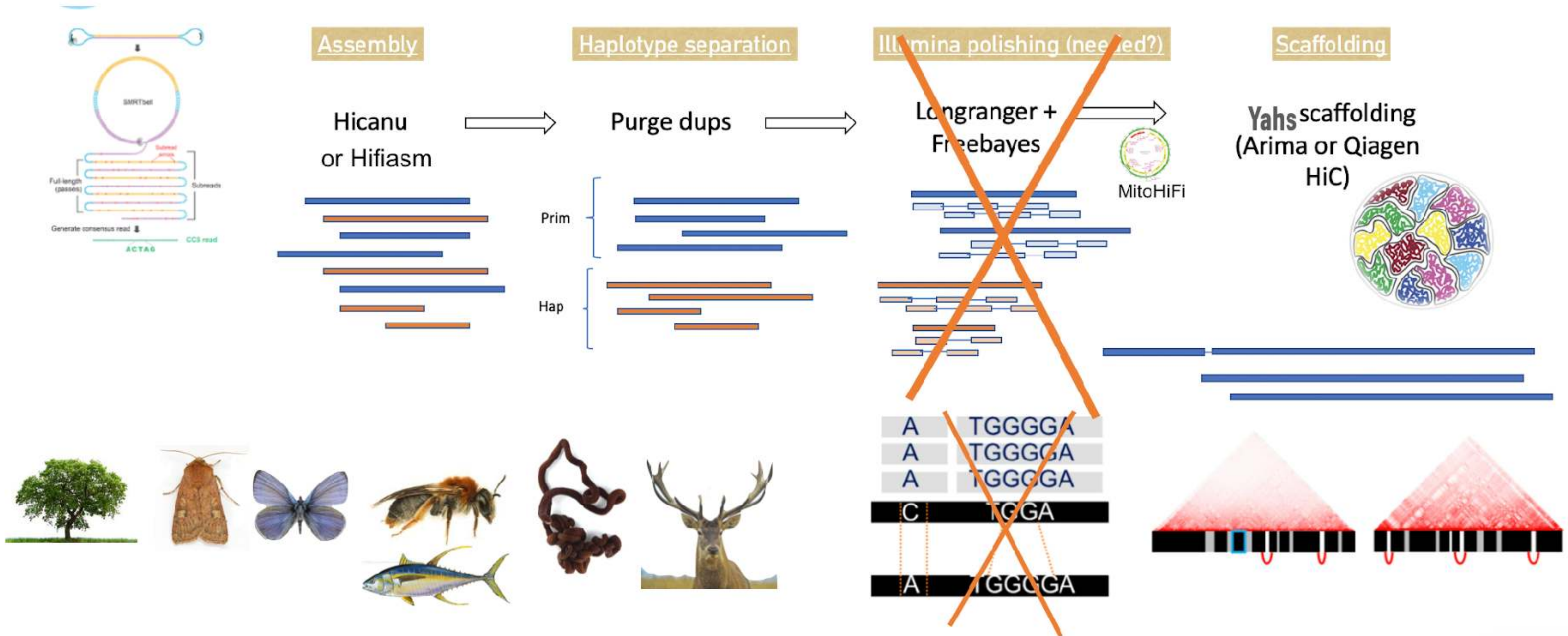
python

- Assemble
- Circularise
- Annotate
- Heteroplasmy



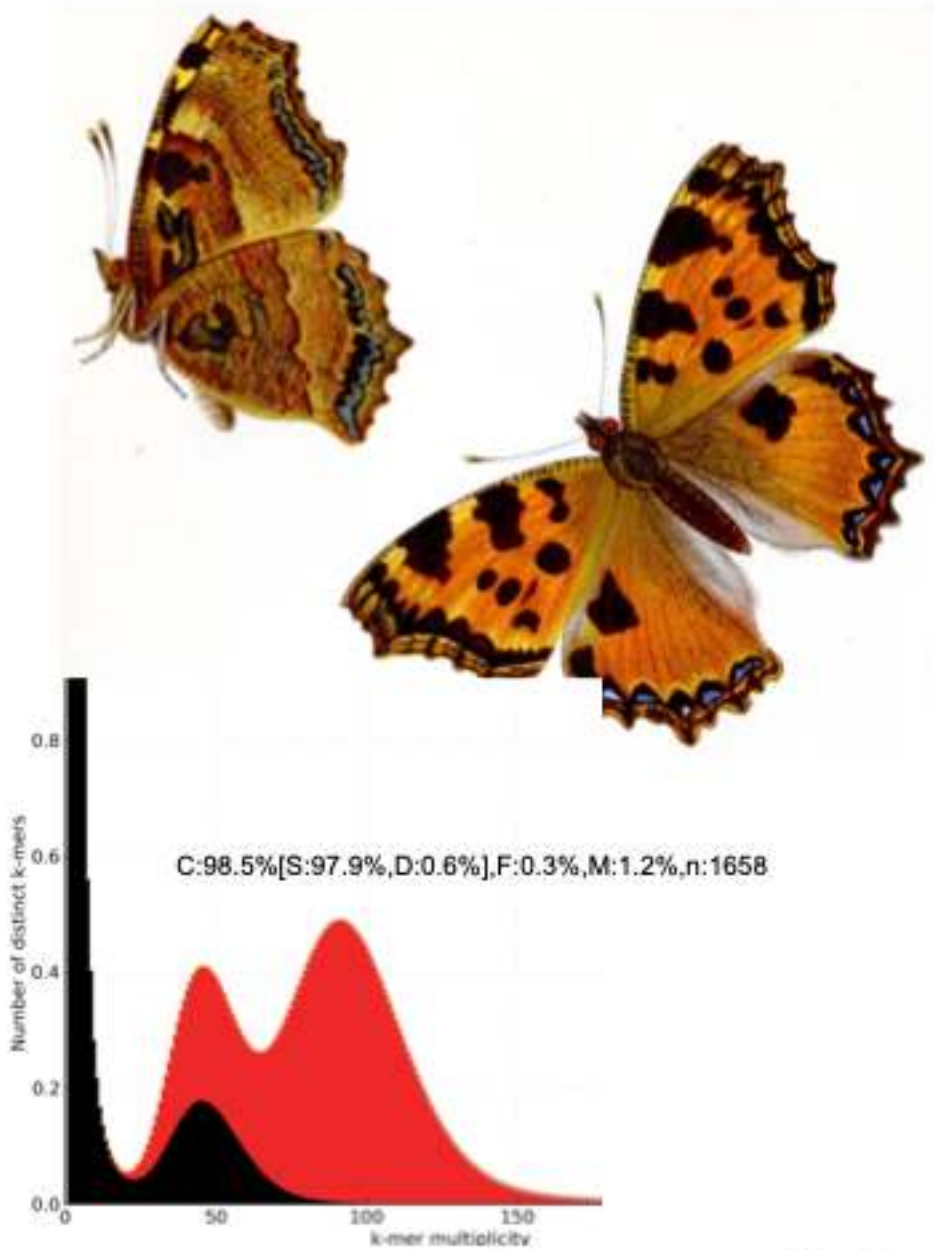
DToL Current Pipeline

- Sequencing technologies: PacBio HiFi + HiC (Arima or Qiagen)

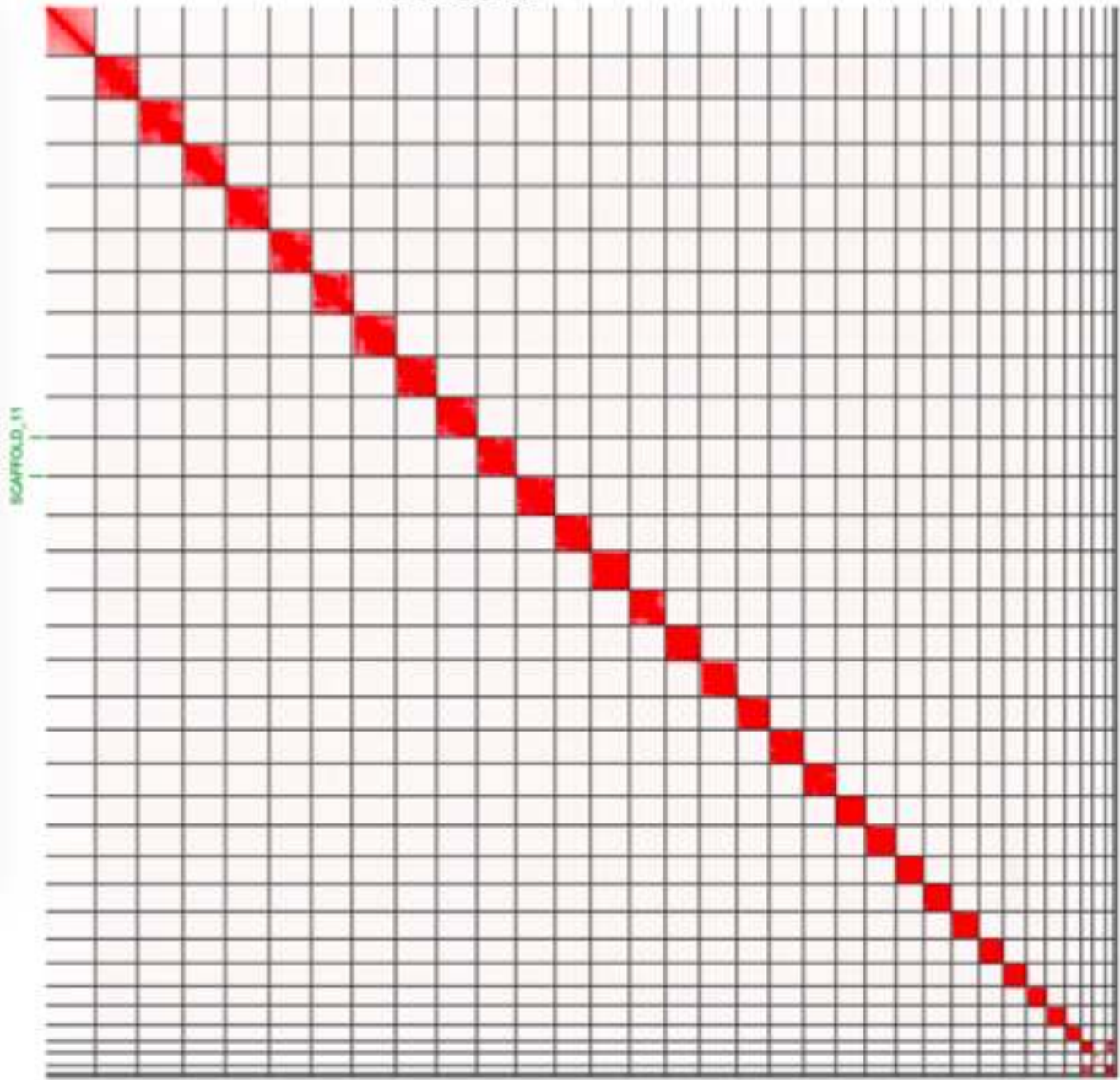


OUT OF THE BOX

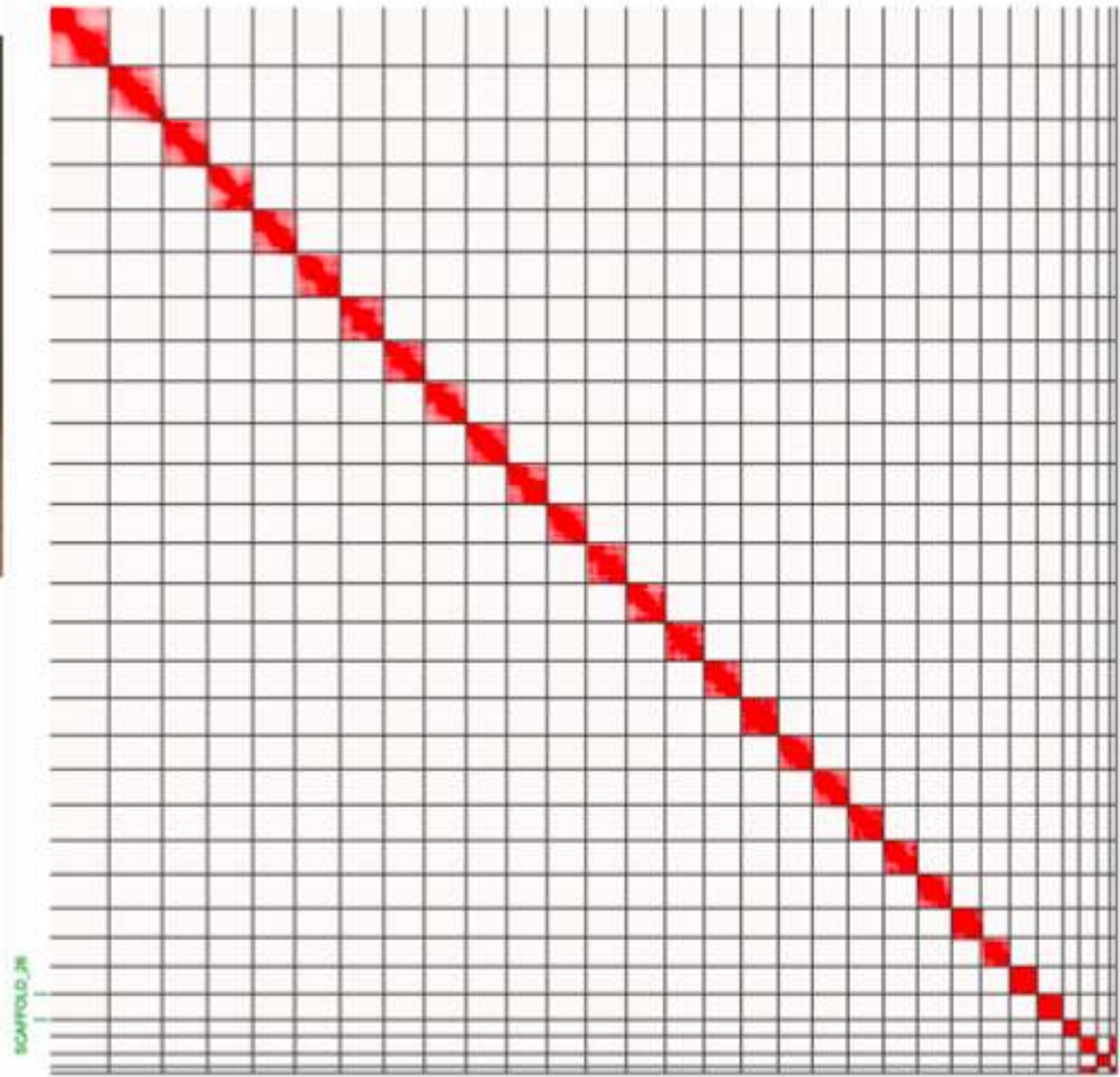
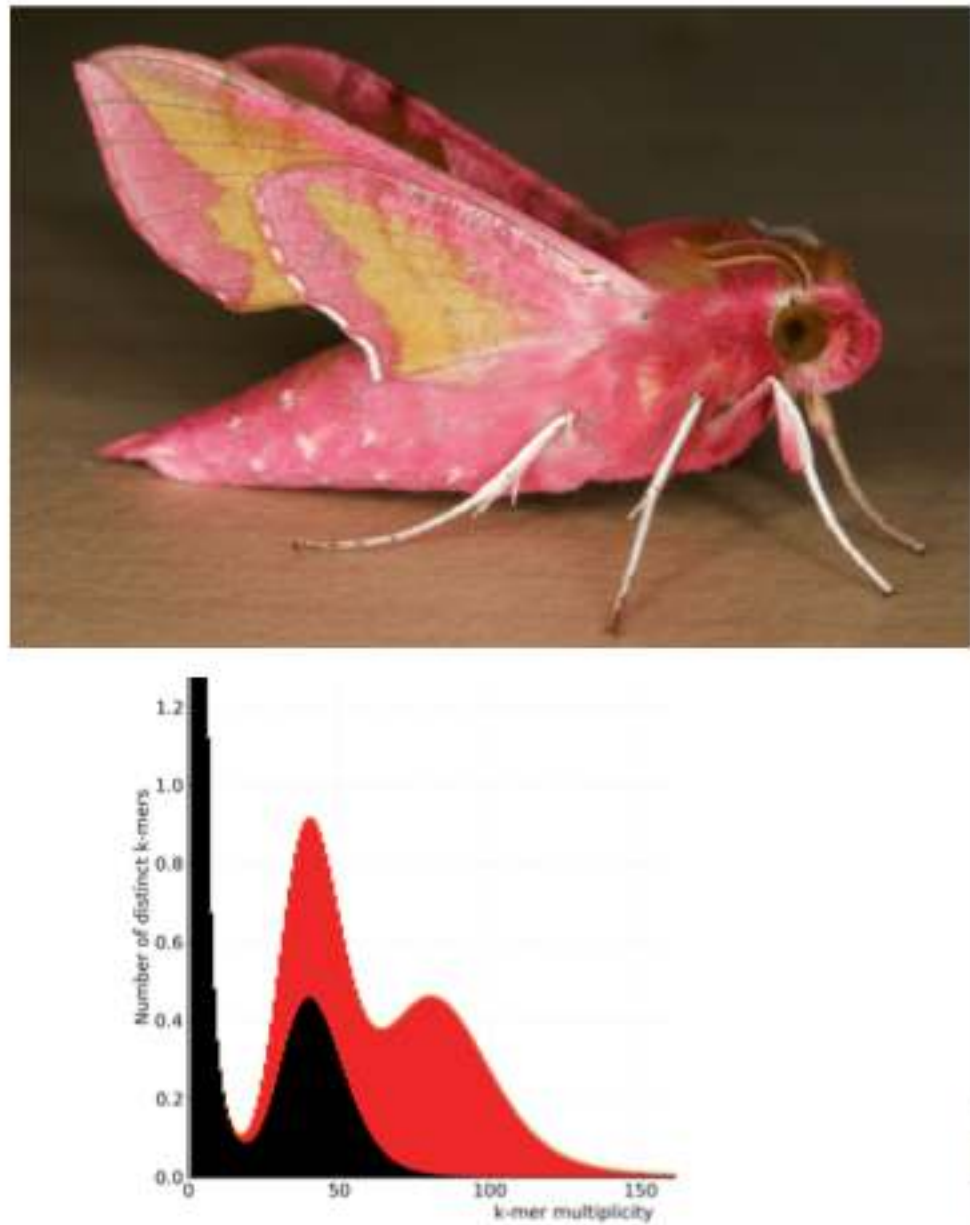
ilNymPoly1



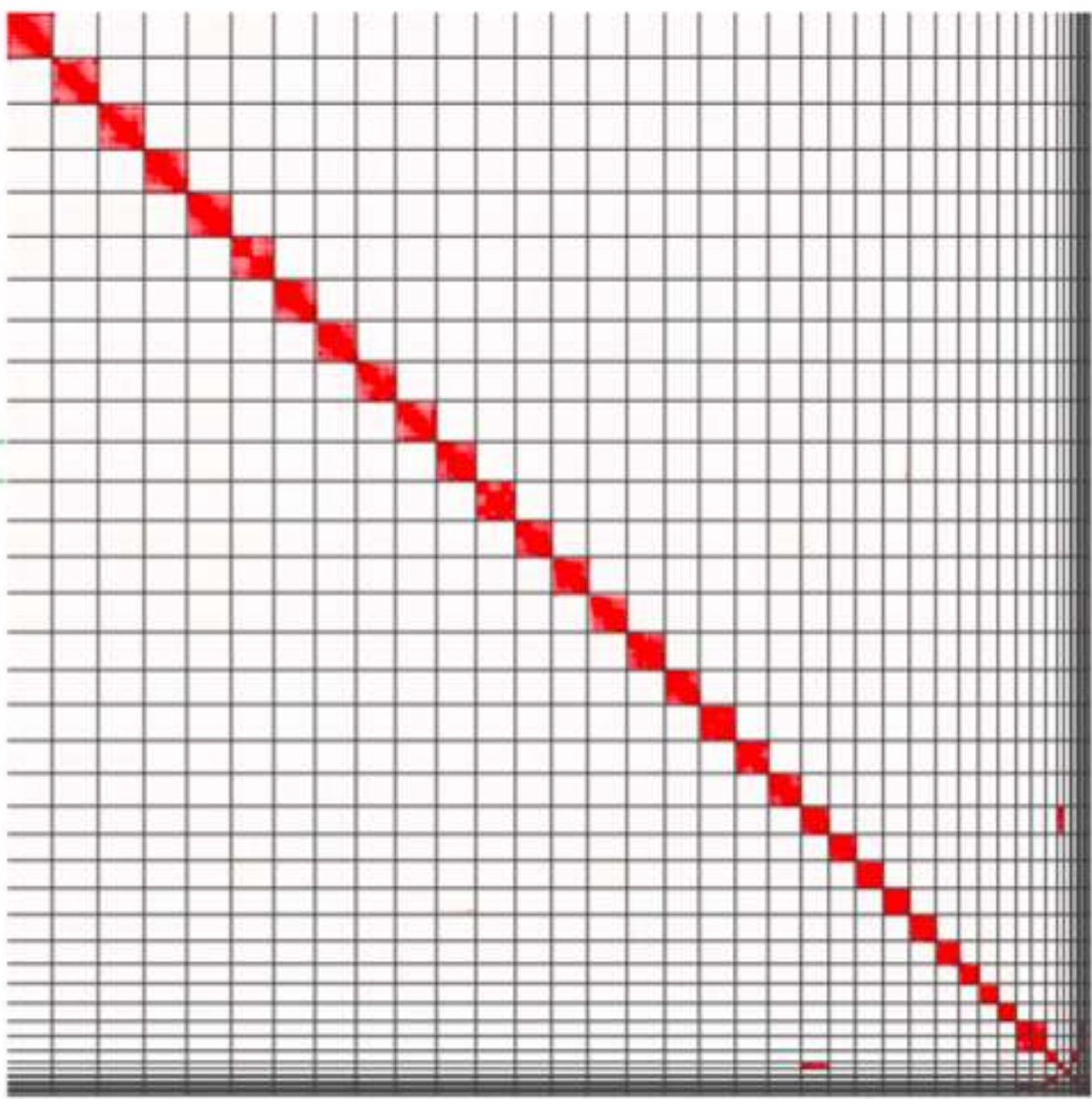
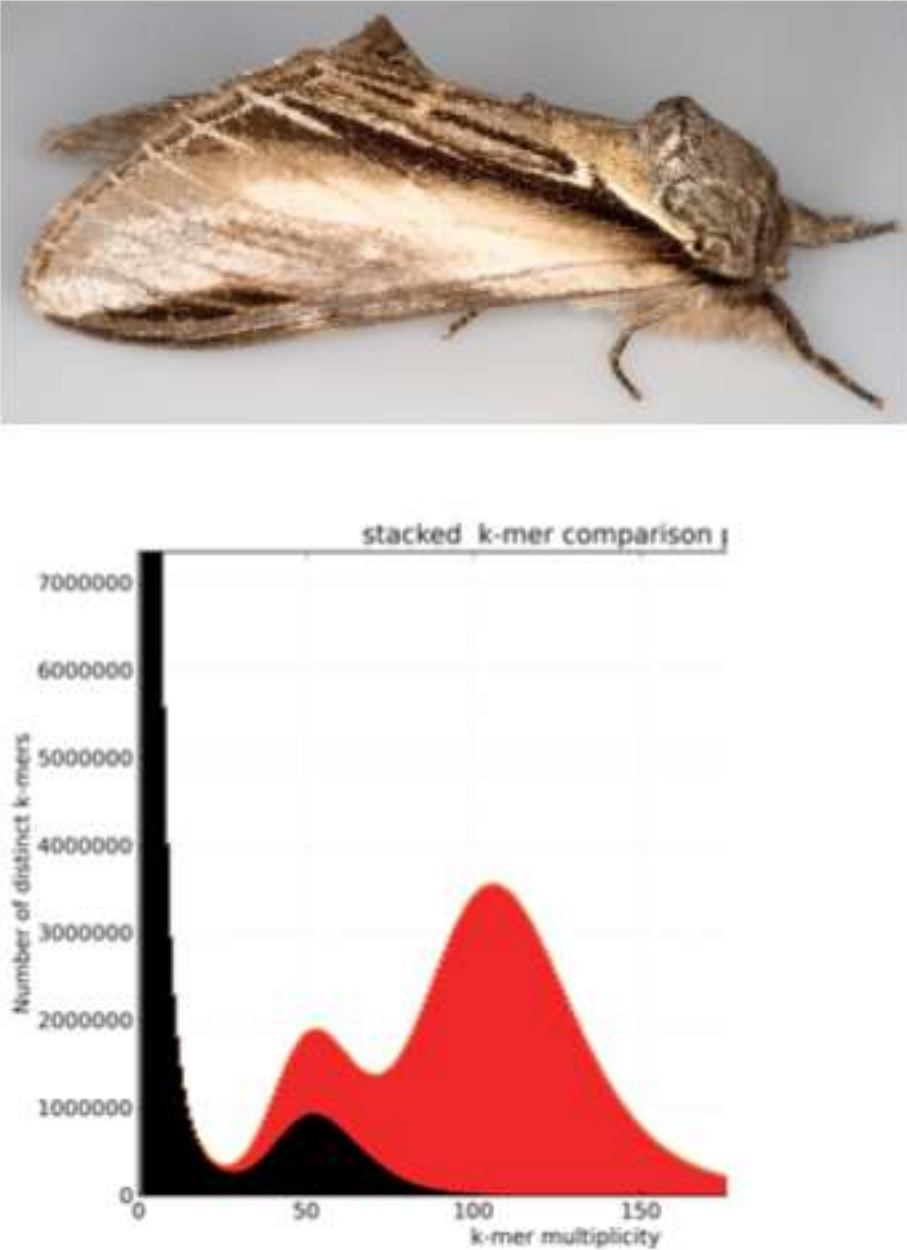
Hifiasm



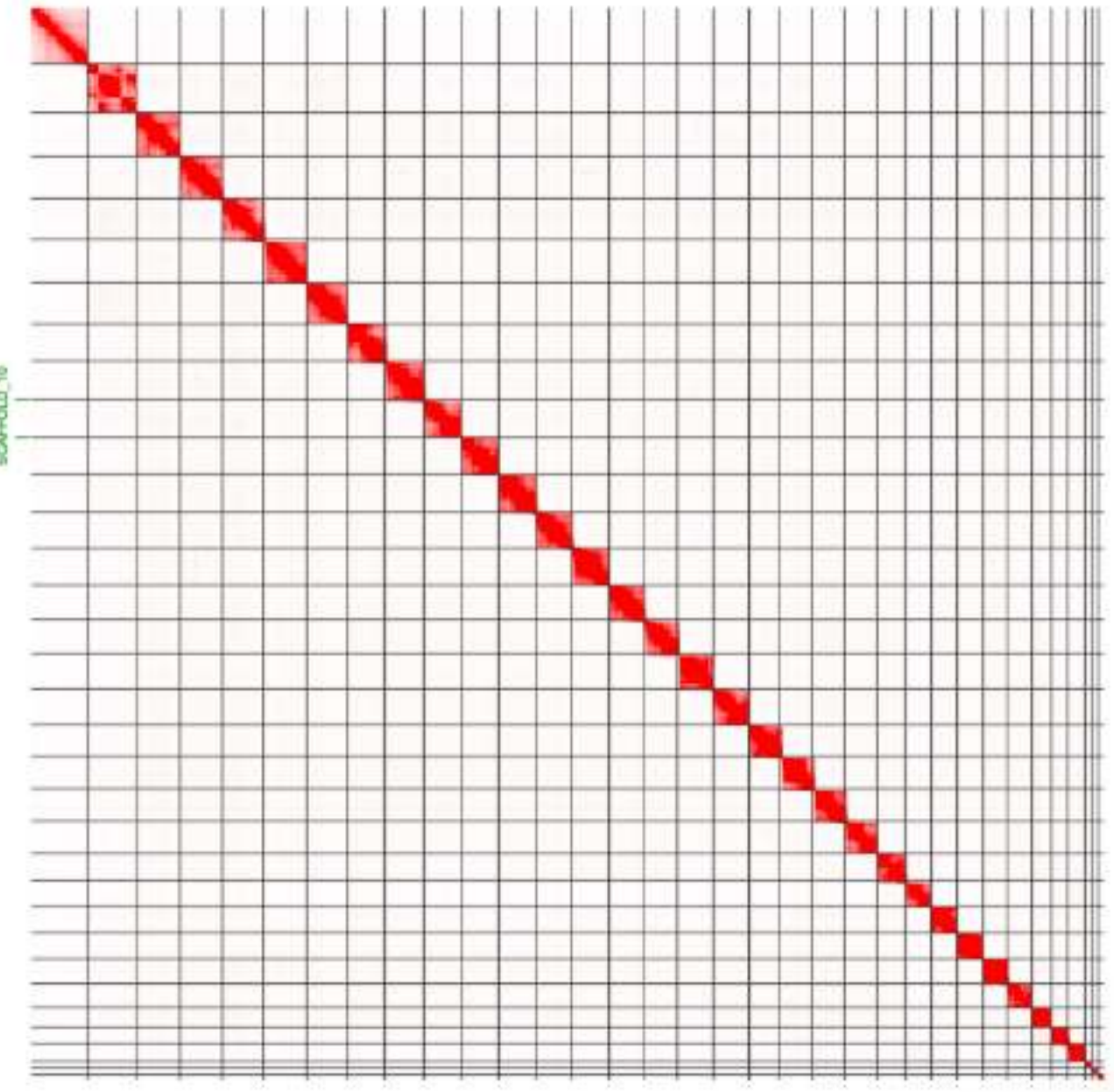
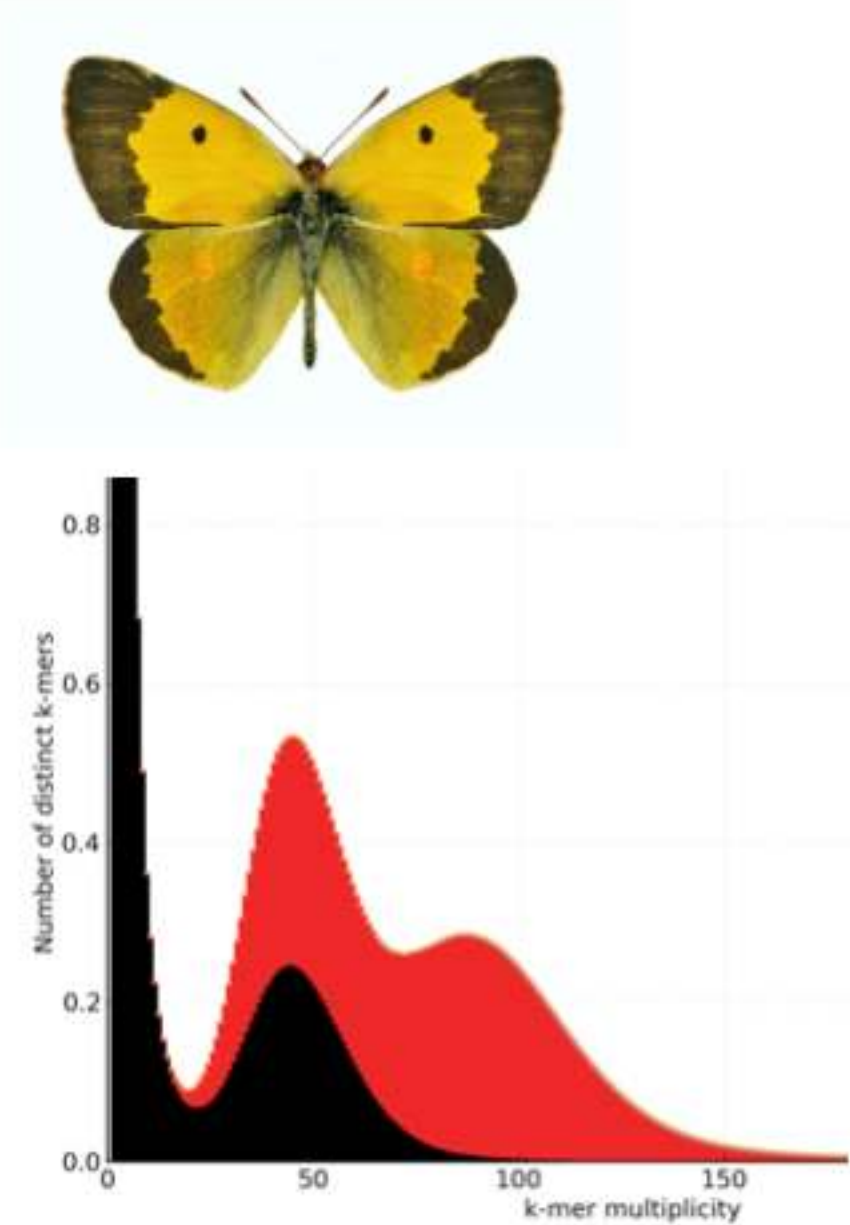
ilDeiPorc1 - *Deilephila porcellus*



Pheosia Tremula (ilPheTrem1)



ilColCroc2 - *Colias crocea*



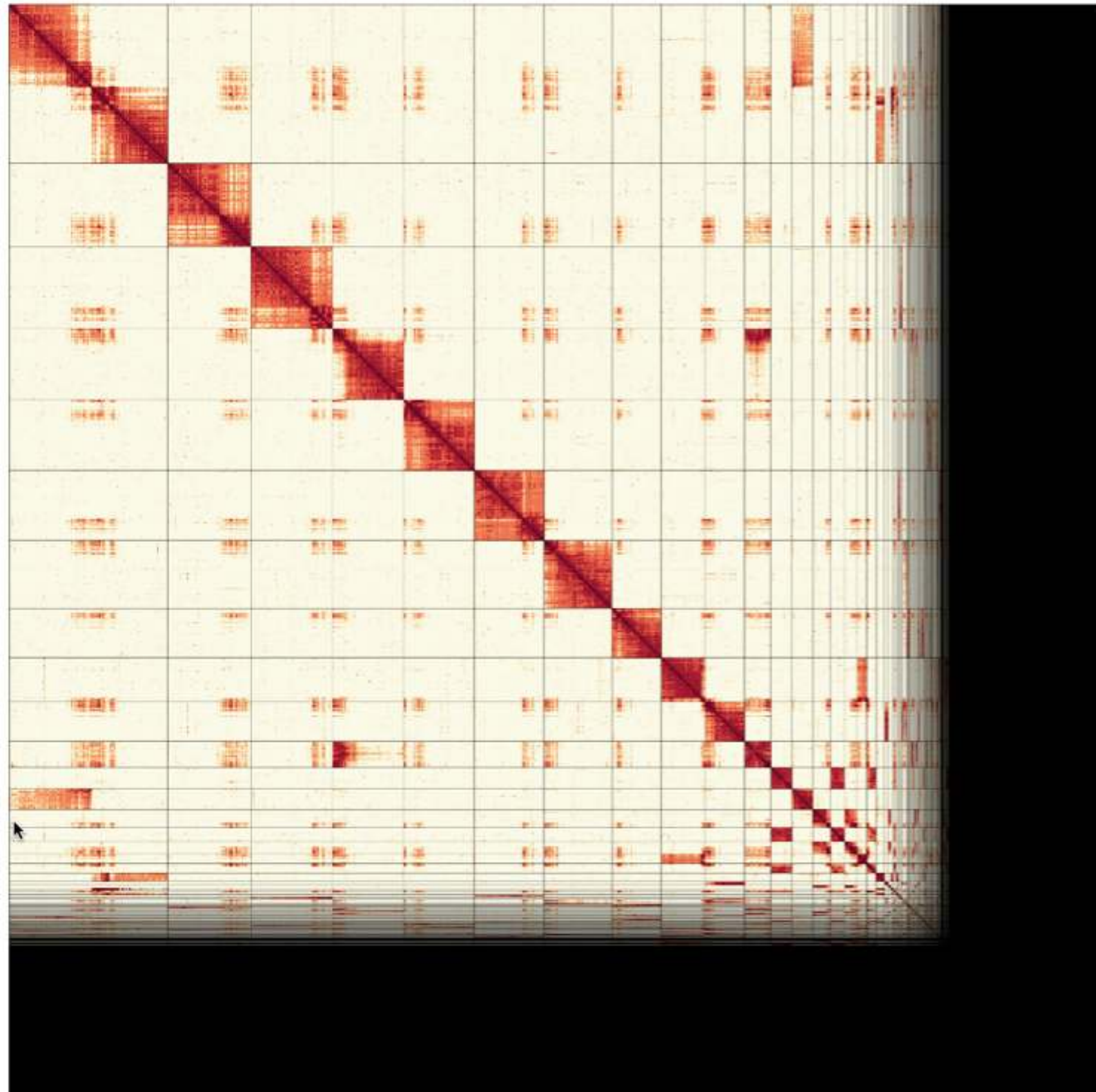
BUT! MANUAL CURATION IS ESSENTIAL



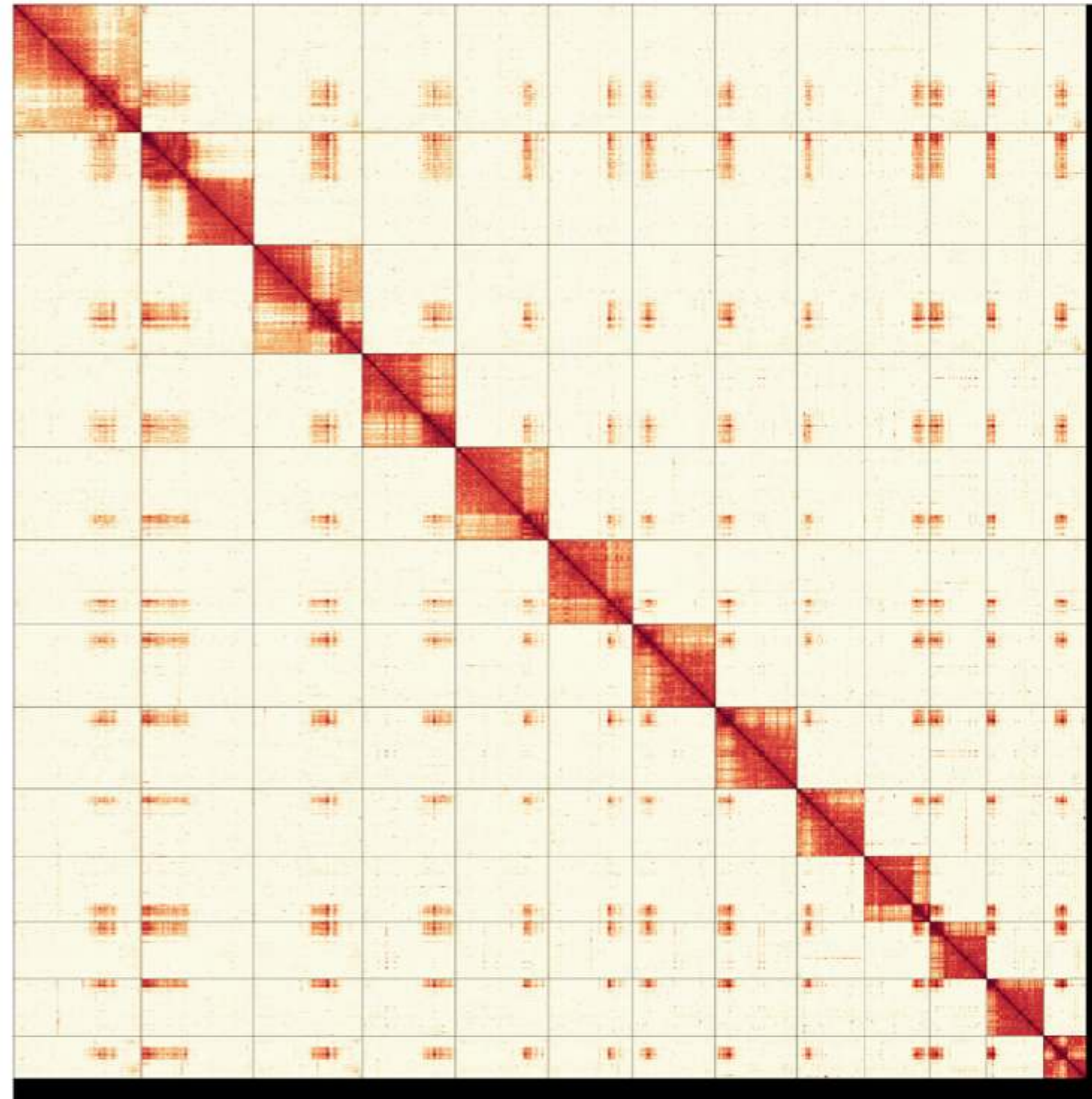
Jo Wood's GRIT
curation team

iyOphLute1_1 -> 372 breaks, 1,377 joins, 2 haplotig removals

BEFORE CURATION



AFTER CURATION

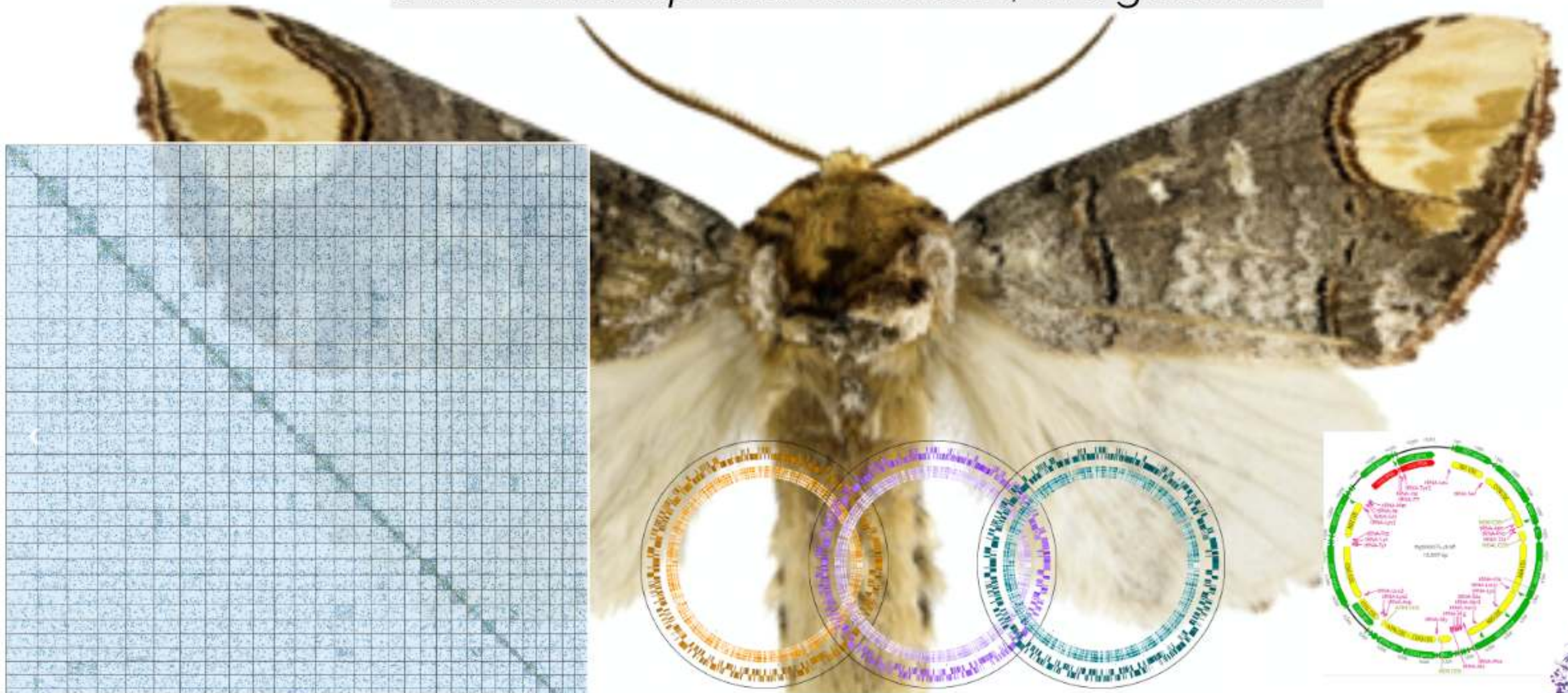


Ophion luteus

Alan Tracey

Assembling genomes of target *and* microbiome

Phalera bucephala: one moth, five genomes



31 nuclear chromosomes

Wolbachia A, B1, B2

mitochondrion

Emmelien Vancaester, Claudia Weber, Marcela Uliano, Alan Tracey, James Torrance, Jonathan Wood

October 2 - 6
SAVE THE DATE

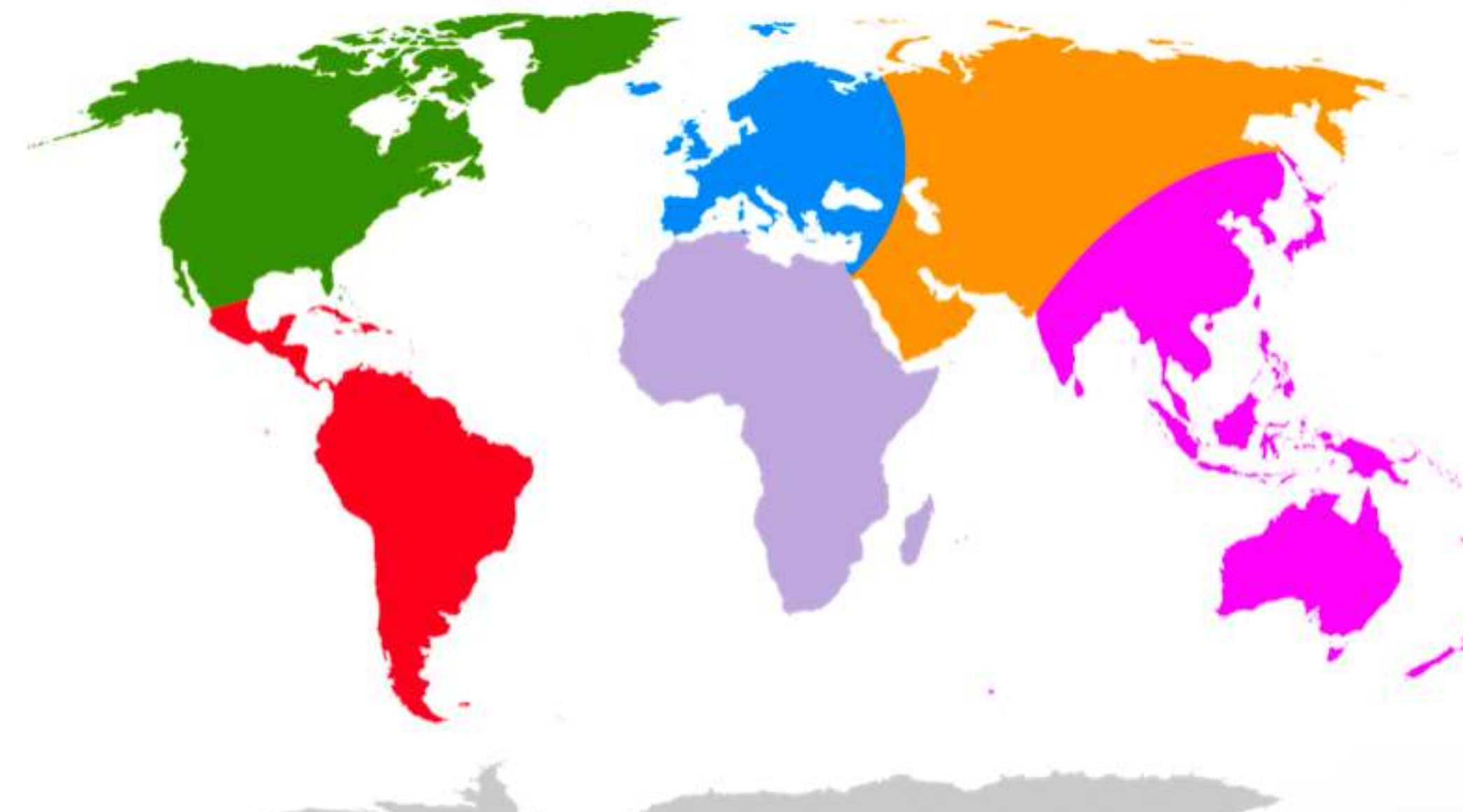


Biodiversity Genomics 2023

Regional Meetings, Scientific topics, JEDI sessions, keynote speakers.

Free to attend!

@BiodivGenomics



Obrigada! Thank you!



Bill Baker
Ester Gaya
Paul Kersey
Ilia Leitch
Greg Palmer



Royal
Botanic Garden
Edinburgh

David Bell
David Long
Laura Forrest
Mary Gibby
Michelle Hart
Neil Bell
Pete Hollingsworth
Rebecca Yahr



Nova
Mieszkowska
Willie Wilson
Michael Cunliffe
John Bishop
Helen Jenkins
Robert Mrowicki
Padrick Adkins
Joanna Harley



Alex Twyford



Neil Hall
Iain Macaulay
Karim Gharbi
Jim Lipscombe
David Swarbreck
Ross Lowe
Rob Davey
Felix Shaw
Sally Warring
Jamie McGowan
Alice Minotto
Seanna McTaggart



Ian Barnes
Gavin Broad
Jonathan Gabriel
Charlotte Barclay
Andrew Briscoe
Mark Carine
Matt Clark
Gerry Hey
Lauren Hughes
Tim Littlewood
Jacqueline MacKenzie-Dodds
Raju Misra
Ben Price
Chris Raper
Fred Rumsey
John Tweddle
Heather Allen
Darren Chooneea
Lyndall Pereira da Conceicao
Laura Sivess
Olga Sivell



University of Oxford and Wytham Woods

Peter Holland
Owen Lewis
Tom Richards
Liam Crowley
Amber Harper
Elisabet Alacid Fernandez
Estelle Kilias
Nigel Fisher
František Sládeček
Lauren Sumner-Rooney
Doug Boyes (CEH)
Alistair McGregor (Brookes Univ)
Karl Wotton (Exeter Univ)



Richard Durbin
Shane McCarthy
Iliana Bista

EMBL-EBI

Paul Flicek
Suran Jayatilaka
Fergal Martin
David Thybert
Jeena Rajan
Kevin Howe
Guy Cochrane
Peter Harrison
Leanne Haggerty
Jamie Allen
Carlos Garcia Giron
Matthieu Muffato



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institute

Tree of Life

Alan Tracey
Amit Vishwakarma
Andrew Varley
Chloe Leech
Damon Lee Pointon
Emmelen Vancaester
Graeme Oatley
James Torrance
Joanna Collins
Jonathan Wood
Katie Woodcock
Kenneth Haug
Kerstin Howe
Ksenia
Krasheninnikova
Maja Todorovic
Manuela Kieninger
Mara Lawniczak
Marcela Uliano da Silva
Mark Blaxter
Matt Berriman
Michelle Strickland
Nancy Holroyd
Nick Salmon
Radka Platte
Raquel Amaral
Robbie Heathcote
Sarah Pelan
Sophie Potter
Victoria Wright
William Chow
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James Uphill
John Tushabe
Karen Oliver
Michelle Smith
Robin Moll
Tracey Chillongworth

Collaborators

Jonas Korch et al
Pacific Biosciences
Dan Turner et al
Oxford Nanopore

Team301

Chris Laumer
Claudia Weber
Emmelein Vancaester
Erna King
Lewis Stevens
Max Brown
Pablo Gonzalez
Rich Challis



Obrigada! Thank you!



Jayathilaka
Martin
Thybert
Rajan
Howe
ochrane
Harrison
e Haggerty
Allen
Garcia Giron
eu Muffato

Matt Berriman
Michelle Strickland
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Scientific Operations

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Oxford Nanopore



.....		
Phred Quality Score	Probability of incorrect base call	Base call accuracy
30	1 in 1000	99.9%
40	1 in 10,000	99.99%
50	1 in 100,000	99.999%