

Session 2.3 – Ensamblado

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Programa Formación Continua, ISCIII

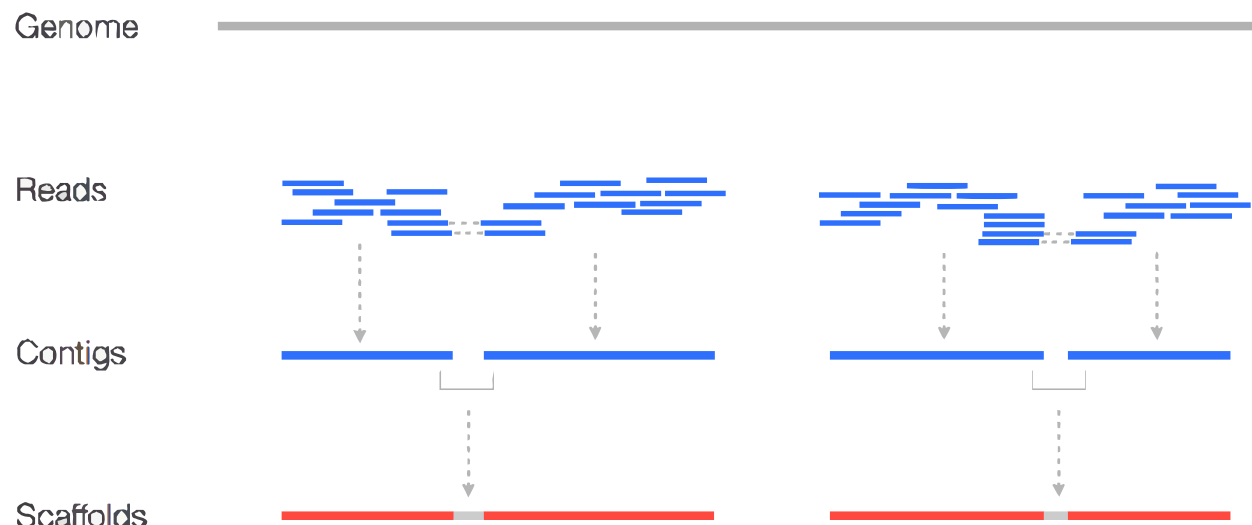
Assembly

Reconstruct the sequence of the original DNA from shorter DNA sequences or small fragments known as reads

- ***De novo***: with no previous knowledge of the genome to be assembled. It overlap the end of the end of each read in order to create a longer sequence.
- ***Assembly with reference***: A similar but not identical genome guides the assembly process. Map reads over supplied genome.

Assembly: contig y scaffold

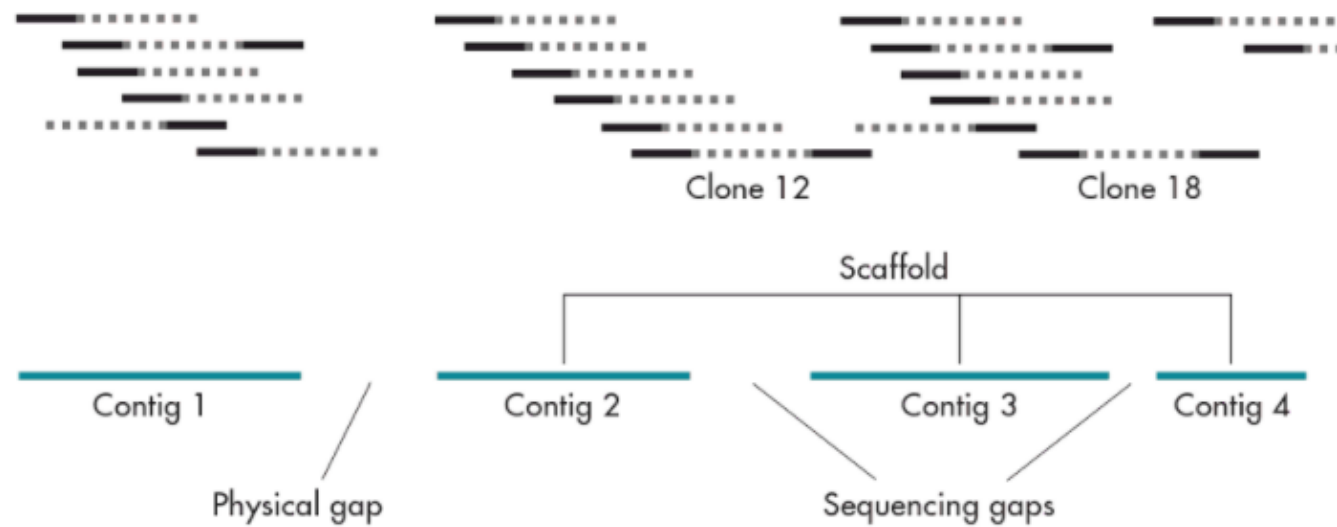
- **Contig:** continuous sequence made up of overlapping shorter sequences
- **Scaffold:** two or more contigs located and rearranged according to spatial information (pair-end, mate pair, reference)



<https://www.biostars.org/p/25322/>

Assembly: gaps

- **Sequencing gaps:** Position and orientation known by spatial information
- **Physical gaps:** No information about adjacent contigs

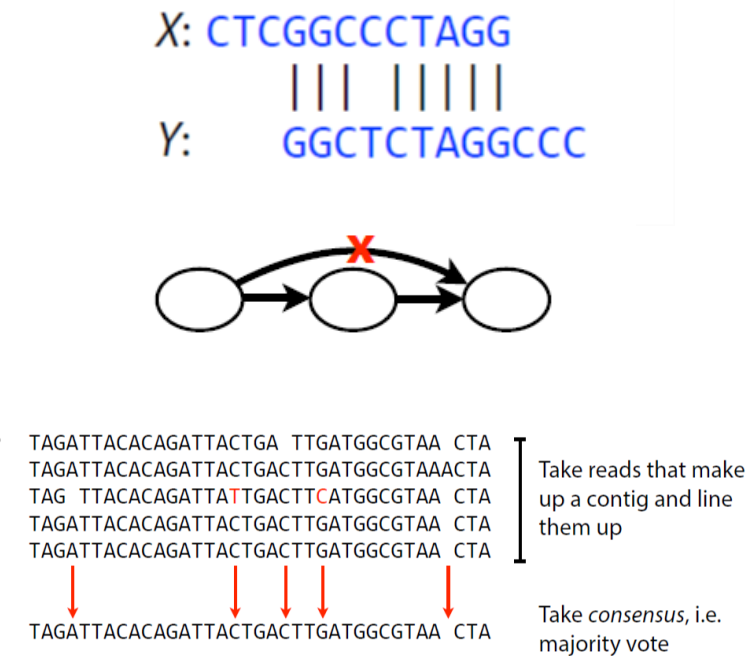


Gene Cloning, Lodge *et al.*

Assembly: Algorithms

- Overlap, Layout, Consensus (OLC - overlap graph):
 - O - first overlaps among all the reads are found
 - L - then it carries out a layout of all the reads and overlaps information on a graph
 - Removes redundant and low quality overlaps
 - C - and finally the consensus sequence is inferred

Ex. Newbler, Mira, Celera Assembler, CAP3, PCAP, Phrap, Phusion.



Assembly: Algorithms

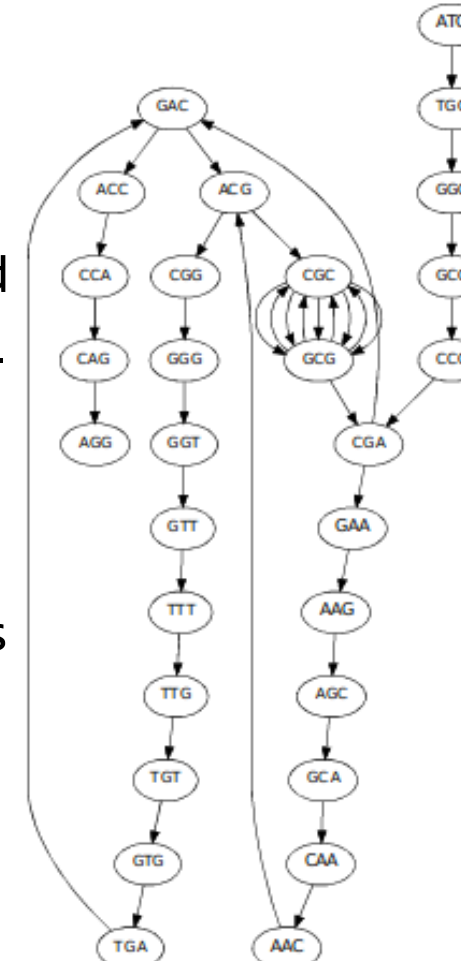
- **De Bruijn Graph (DBG: k-mer graph)**

Chopping reads into much shorter k-mers (fixed length fragments) and then using all the k-mers to form a DBG and infer the contigs.

- Nodes in the graph are k-mers
- Edges represent consecutive k-mers (which overlap by k-n symbols)

Ex. SPAdes, ABYSS, Velvet, AllPaths, Soap....

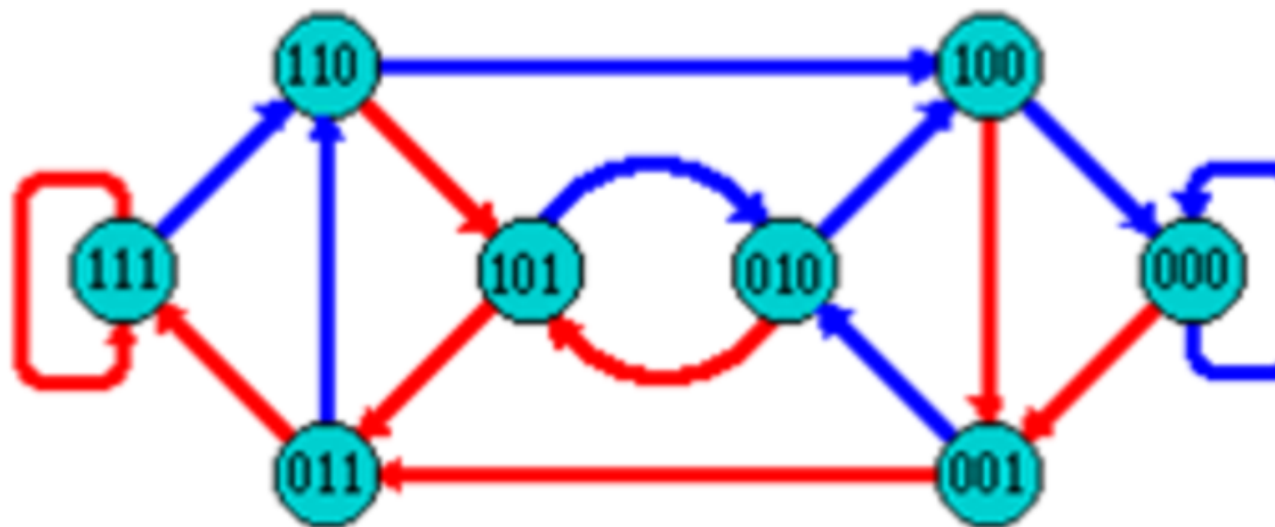
https://medium.com/@han_chen



de Bruijn Graphs

- A directed graph of sequences of symbols
- Nodes in the graph are k-mers
- Edges represent consecutive k-mers (which overlap by k-1 symbols)

Consider the 2 symbol alphabet (0 & 1) de Bruijn Graph for k = 3

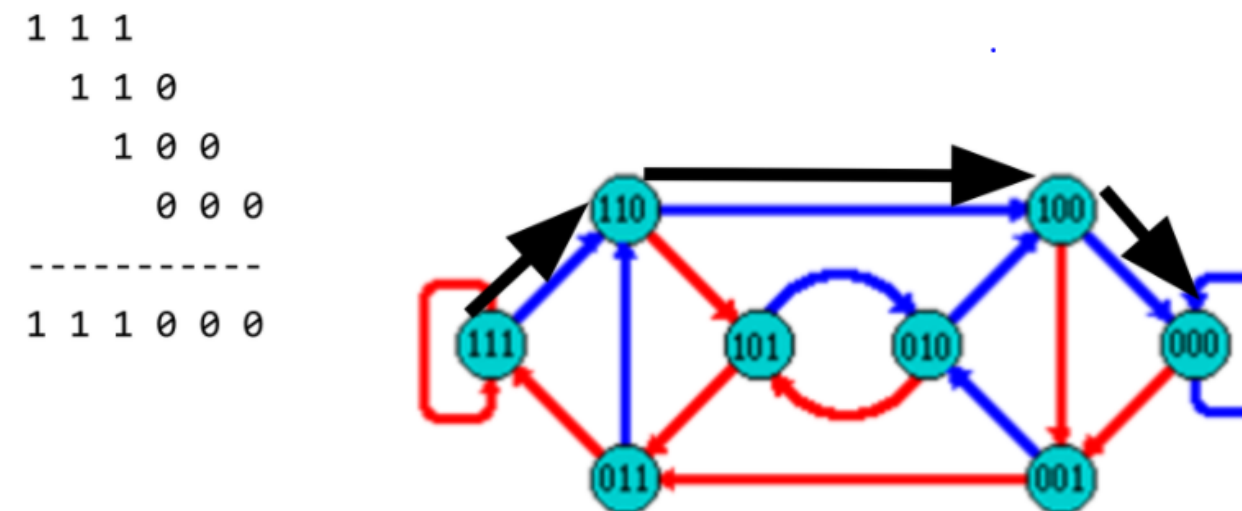


<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

Producing sequences

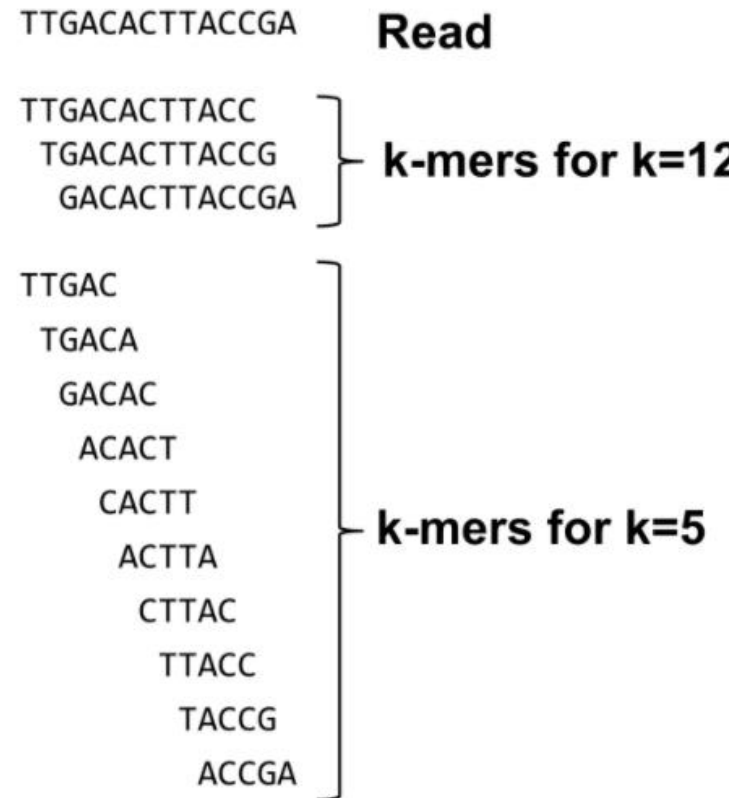
- Sequences of symbols are produced by moving through the graph

e.g. 111000 = 111 -> 110 -> 100 -> 000



<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

What are K-mers?



<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

K-mers de Bruijn graph



Example #1:

HAPPI PINE INESS APPIN

<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

K-mers de Bruijn graph



Example #1:

HAPPI PINE INESS APPIN

All 4-mers:

HAPP PINE INES **APPI**
APPI NESS PPIN

Unique 4-mers:

HAPP **APPI** PINE PPIN INES NESS

<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

K-mers de Bruijn graph



Example #1:

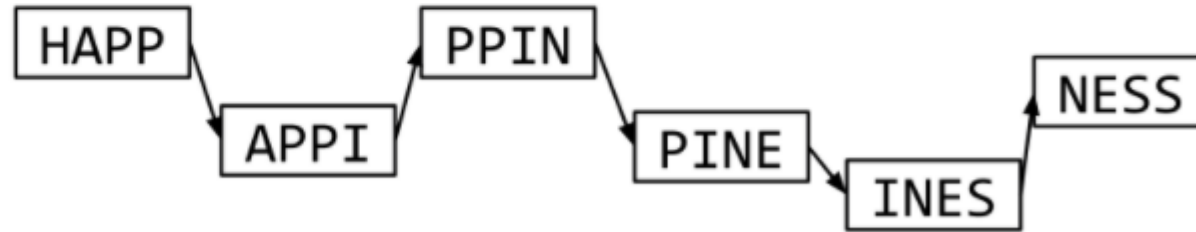
HAPPI PINE INESS APPIN

k = 4 k-mers:

HAPP APPI

PINE PPIN

INES NESS



K-mers de Bruijn graph



Example #1:

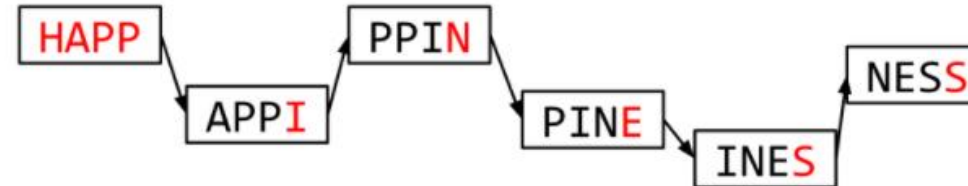
HAPPI PINE INESS APPIN

k = 4 k-mers:

HAPP APPI

PINE PPIN

INES NESS



HAPPINESS

Easy!

<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

The problem of repeats



Example #2:

MISSIS SSISSI SSIPII

<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

The problem of repeats



Example #2:

MISSIS SSISSSI SSIPPI

All 4-mers (9):

MISS	SSIS	SSIP
ISSI	SISS	SIPP
SSIS	ISSI	IPPI

Unique 4-mers (7):

MISS SSIS SSIP ISSI SISS SIPP IPPI

The problem of repeats

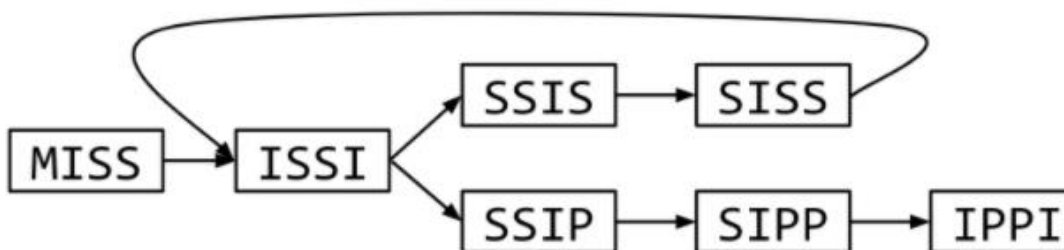


Example #2:

MISSIS SSISSI SSIPPI

All 4-mers:

MISS ISSI SSIS SISS SSIP SIPP IPPI



The problem of repeats

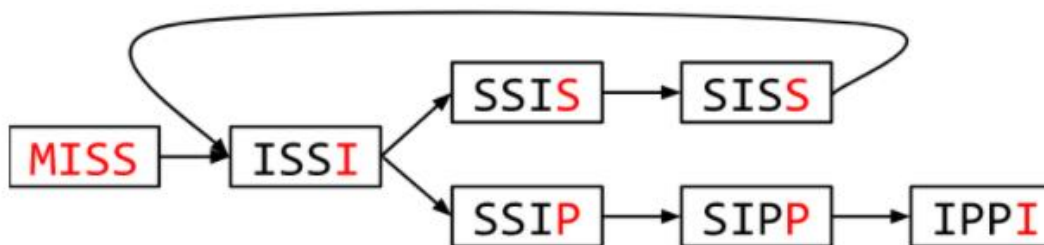


Example #2:

MISSIS SSISSI SSISSI

All 4-mers:

MISS ISSI SSIS SISS SSIP SIPP IPPI



MISSISSIPPI or MISSISSISSISSIPPI or ...

<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

Different k

Example #2a:
MISSIS SSISI SISSIPPI

Different k

Example #2a:

MISSIS SSISSI SSIPPI

All 5-mers (6):

MISSI SSISS SSIPP

ISSIS SISSI SIPPI

Unique 5-mers (6, no duplicates):

MISSI ISSIS SSISS SISSI SSIPP SIPPI

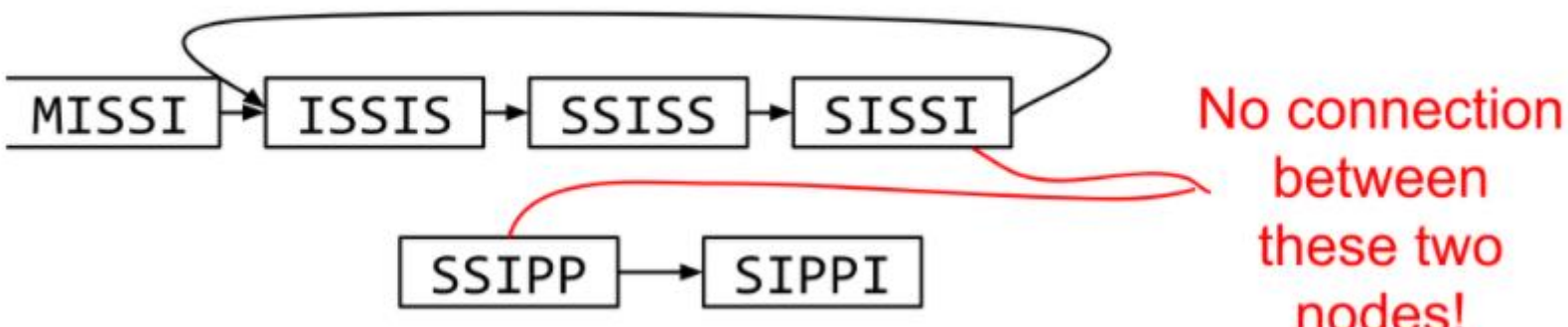
Different k

Example #2a:

MISSIS SSISSI SSIPPI

This time $k = 5$ k-mers:

MISSI ISSIS SSISS SISSI SSIIPP SIPPI



<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

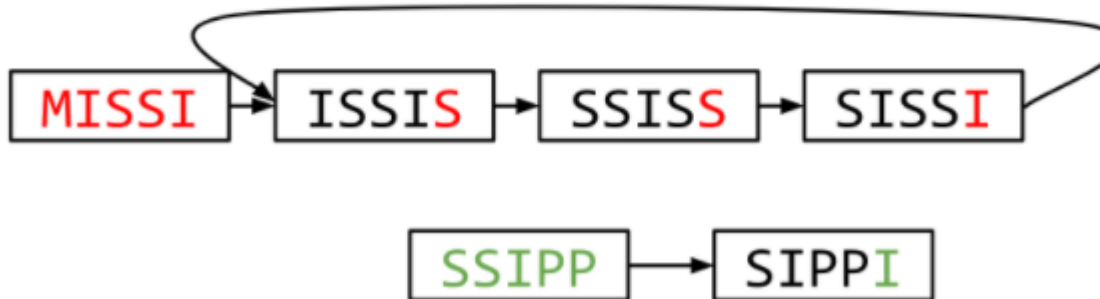
Different k

Example #2a:

MISSIS SSISSI SSISSI

This time $k = 5$ k-mers:

MISSI ISSIS SSISS SISSI SSIIPP SIPPI



MISSISSIS

SSISSIPI

<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

Choose k wisely

- Lower k
 - More connections
 - Less chance of resolving small repeats
 - Higher k-mer coverage
- Higher k
 - Less connections
 - More chance of resolving small repeats
 - Lower k-mer coverage

Optimum value for k will balance these effects.

<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

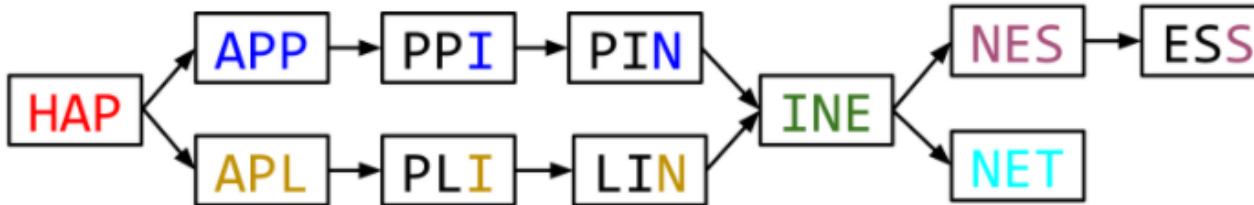
Read errors



Example #3:
HAPPI INESS APLIN PINET

$k = 3$ k-mers:

HAP APP PPI INE NES ESS APL PLI LIN PIN NET



6 contigs: HAP APPIN APLIN INE NESS NET

<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

More coverage

- Errors won't be duplicated in every read
- Most reads will be error free
- We can count the frequency of each k-mer
- Annotate the graph with the frequencies
- Use the frequency data to clean the de Bruijn graph



More coverage depth will help overcome errors!

<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

Algorithms: DBG

- Why choosing DBG:
 - Sequencing bias
 - Sequence errors
 - Sequence length
- DBG Flaws:
 - Millions of pieces
 - Much, much shorter than the genome
 - Lots of them look similar
 - Missing pieces
 - Some parts can't be sequenced easily
 - Dirty Pieces - Multiplex
 - Lots of errors in reads
 - Repeats
 - If they are longer than the read length
 - Causes nodes to be shared, locality confusion

<https://galaxyproject.github.io/training-material/topics/assembly/tutorials/debruijn-graph-assembly/slides.html#23>

SPAdes

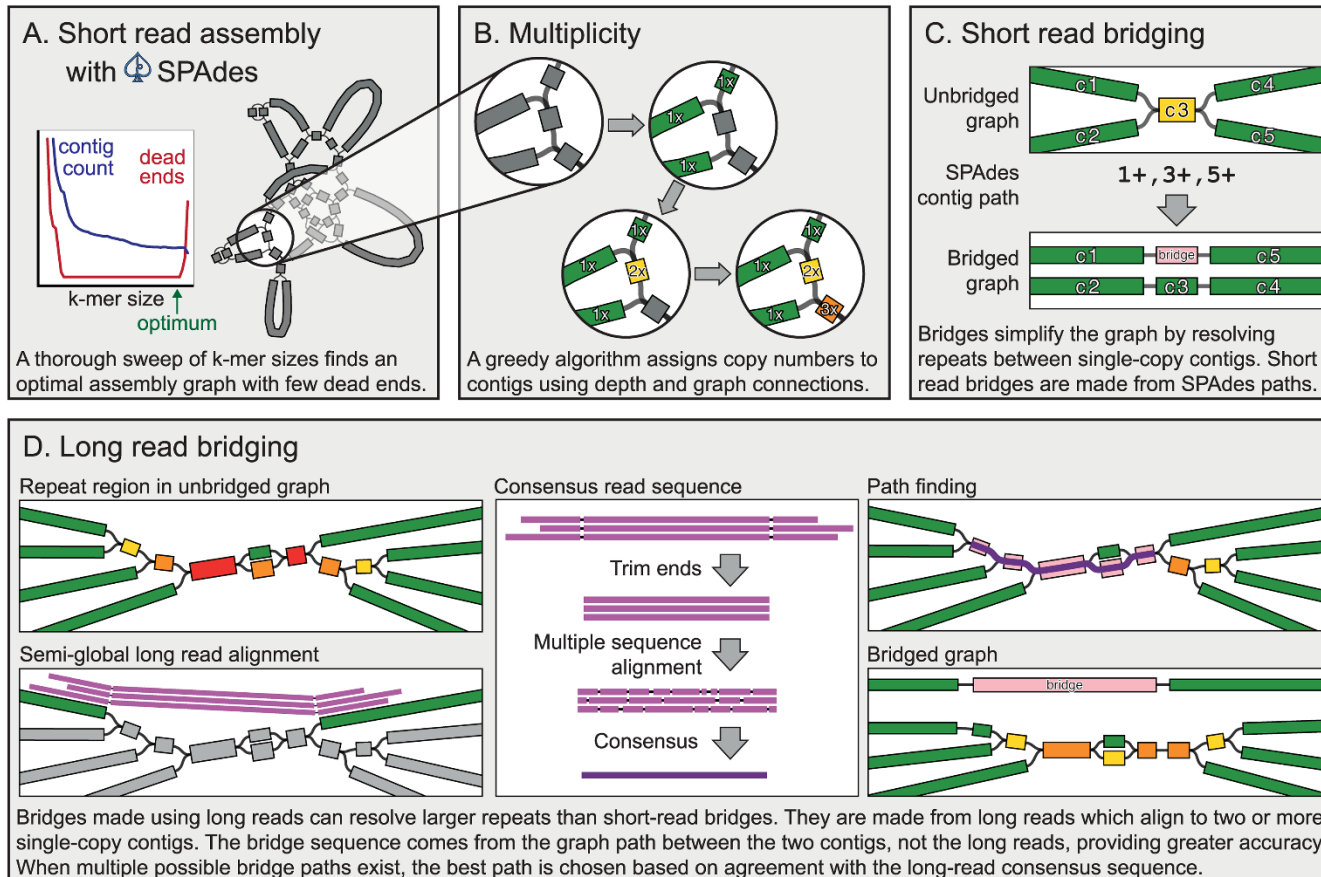


- de Bruijn graph assembler by Pavel Pevzner's group out of St. Petersburg
- Uses multiple k-mers to build the graph
 - Graph has connectivity **and** specificity
 - Usually use a low, medium and high k-mer size together.
- Performs error correction on the reads first
- Maps reads back to the contigs and scaffolds as a check
- Under active development
- Much slower than Velvet
- Should be used in preference to Velvet now.

Assembly: Scaffolding

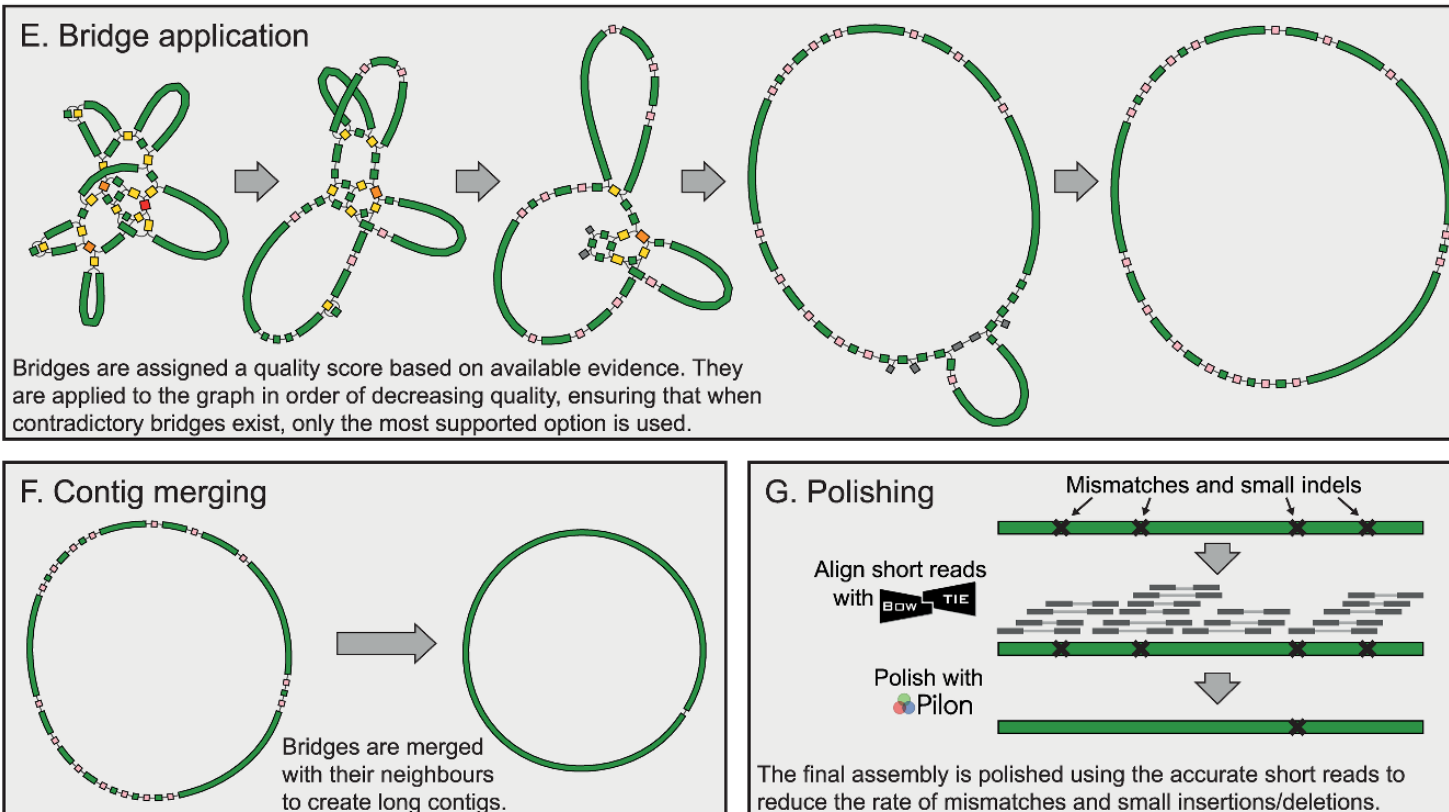
- **From draft:**
 - Order contigs** (Nucmer, if there is reference it can be used to align and guide)
 - Fill the GAPs** (GapFiller, fill sequencing gap (not physical gap))
 - Solve repeated sequence ambiguities** (Expander)
 - Resequence** with different library:
 - Longer fragments and/or distance
- **Tools for assembly improvement**
 - SSPACE (Scaffolding) REAPR (evaluate scaffolding, breaking incorrect scaffolds)
- **Assembly visualizing**
 - Artemis, ACT (compare two or more sequences), Icarus (Quast)

Unicycler



<https://journals.plos.org/ploscompbiol/article?id=10.1371/journal.pcbi.1005595>

Unicycler



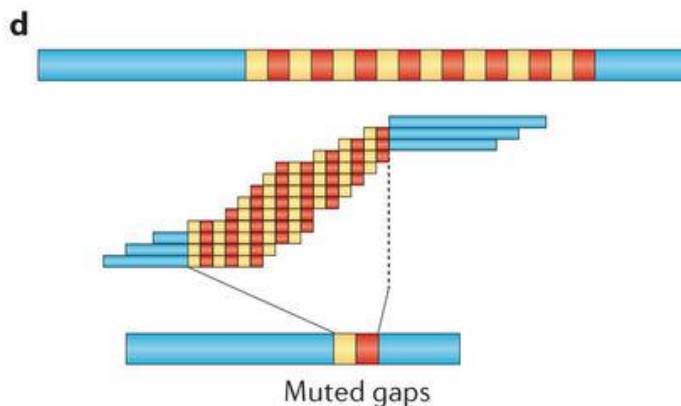
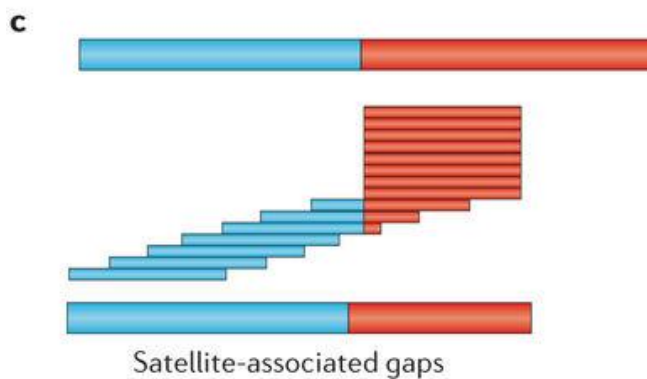
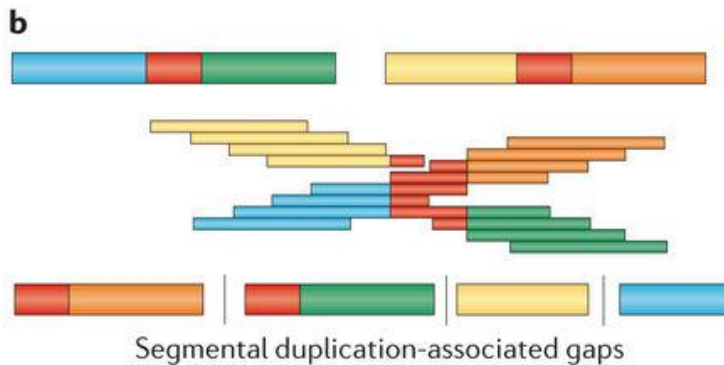
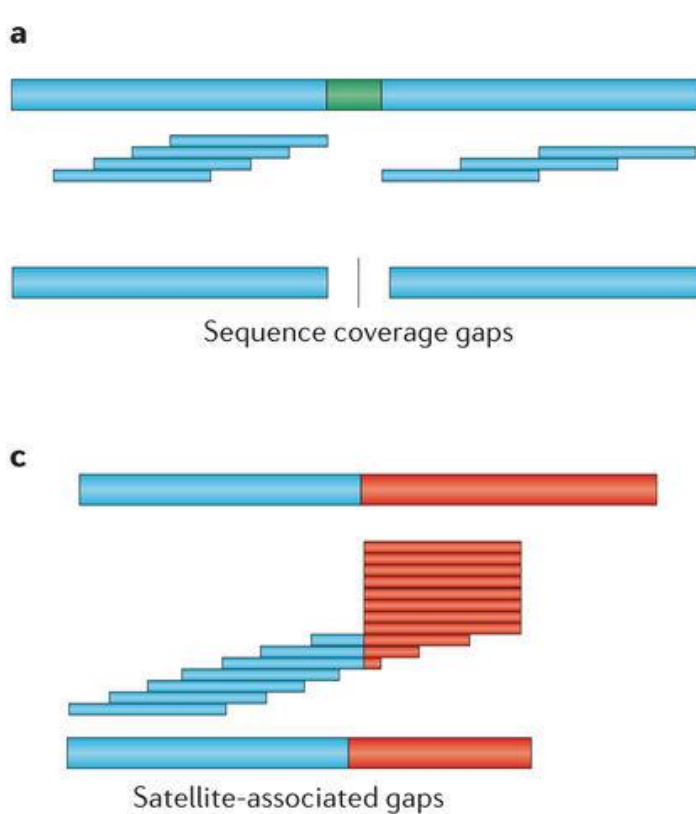
<https://journals.plos.org/ploscompbiol/article?id=10.1371/journal.pcbi.1005595>

A move back to OLC

- New long read technologies
 - PacBio and MinIon
- Assemblers: HGap, CANU
 - Use overlap, layout consensus approach
- CANU can perform hybrid assemblies with long and short reads



Ensamblado: Errores

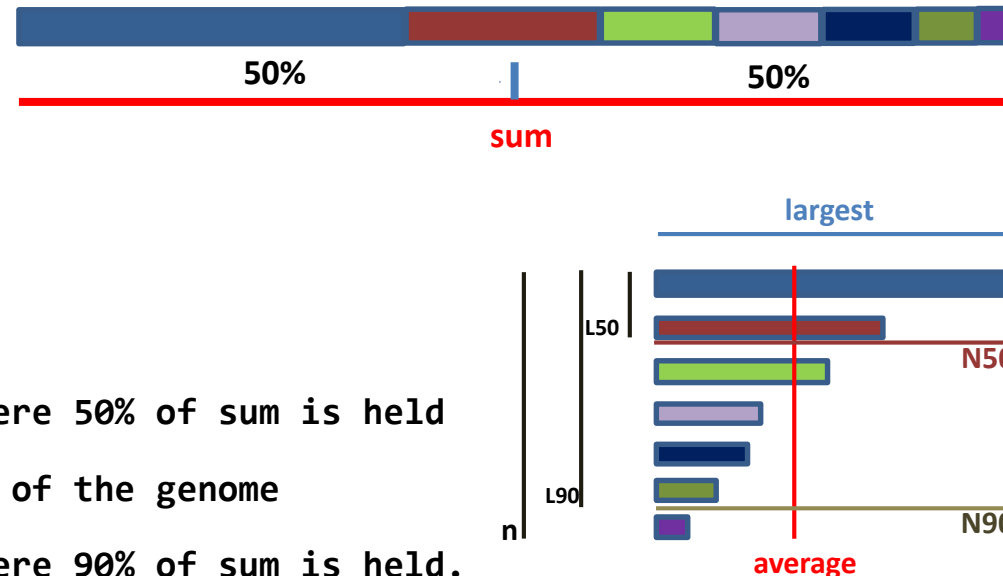


- **A. Gaps – región del genoma sin secuenciar**
- **B. Duplicaciones de gran tamaño**
 - Quimeras
- **Regiones repetidas colapsadas**
 - **C. Terminales**
 - **D. Intersticiales**

Genetic variation and the de novo assembly of human genomes
Chaisson *et al.* 2015

Assembly: Metrics

- **sum** = total bases number
- **n** = contigs number
- **average** = average contig length
- **largest** = largest contig
- **N50** = length of the shortest contig where 50% of sum is held
- **L50** = number of contigs which have 50% of the genome
- **N90** = length of the shortest contig where 90% of sum is held.
- **L90** = number of contigs which have 90% of the genome

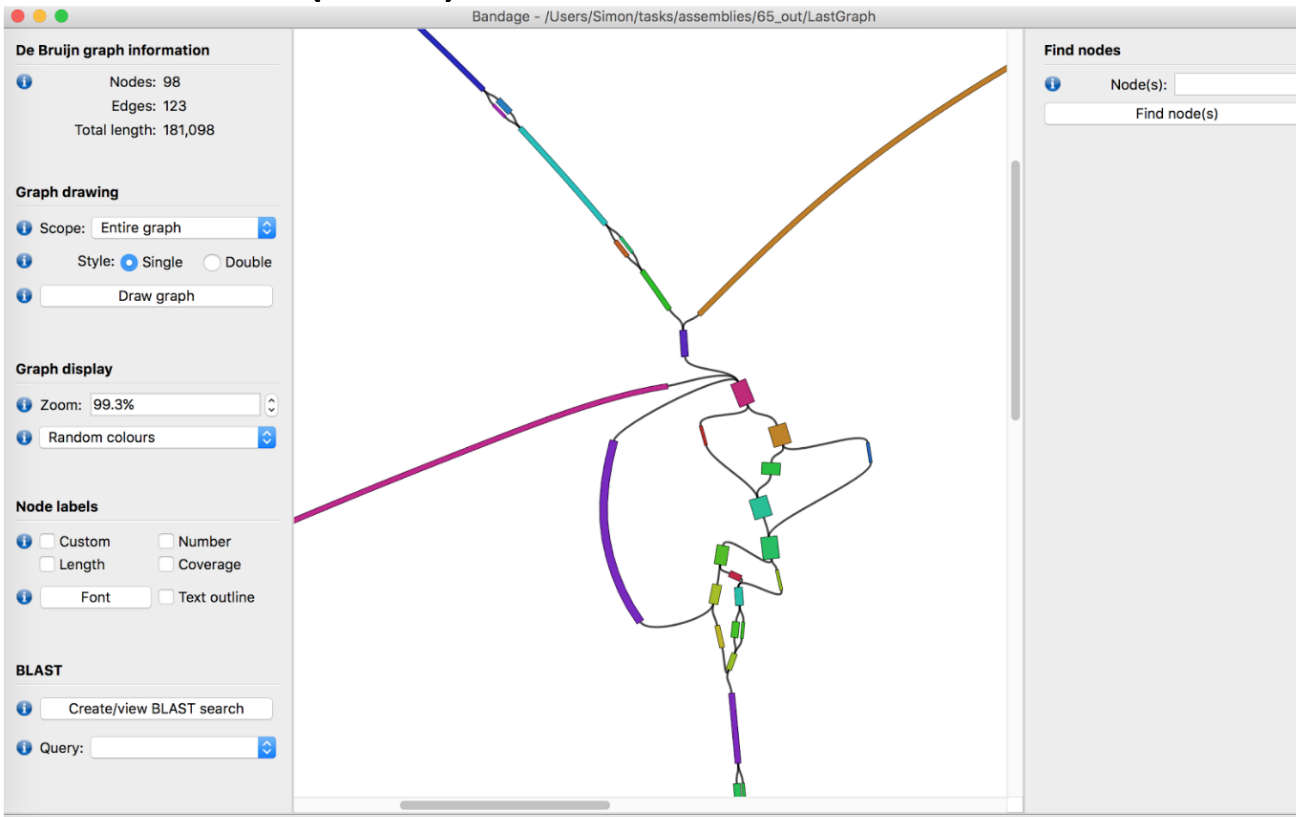


Assembly: Evaluation

- Software that evaluate different algorithms & parameters
iMetAMOS, Koren et al., *BMC Bioinformatics* 2014, 15:126
GAGE-B, Magoc et al., *Bioinformatics* 2013, 29(14):1718-25
- **Graph evaluation:** Bandage, Wick R.R., Schultz M.B., Zobel J. & Holt K.E. (2015)
- **Assembly evaluation:** Quast, Gurevich et al., *Bioinformatics* 2013, 29:8
- **Metrics for a good assembly:**
Large N50
Sum closest to expected
Low n
Low L50

Assembly: Evaluation - Bandage

- Graph evaluation: Bandage, Wick R.R., Schultz M.B., Zobel J. & Holt K.E. (2015)



Assembly: Evaluation - Quast

- Assembly evaluation: Quast, Gurevich *et al.*, *Bioinformatics* 2013, 29:8

	RA_L2073_paired_assembly	RA_L2391_paired_assembly	RA_L2677_paired_assembly	RA_L2978_paired_assembly	RA_L2281_paired_assembly	RA_L2450_paired_assembly	RA_L2701_paired_assembly
Genome statistics							
Genome fraction (%)	81.079	88.828	84.92	90.172	85.733	88.172	92.463
Duplication ratio	1	1	1.001	1.001	1.001	1	1
# genomic features	1736 + 824 part	2113 + 600 part	1881 + 768 part	2157 + 611 part	1992 + 637 part	2073 + 643 part	2368 + 412 part
Largest alignment	16 612	33 033	21 336	25 068	29 638	30 305	40 471
Total aligned length	2 405 510	2 635 297	2 519 300	2 675 166	2 543 440	2 615 874	2 743 222
NGA50	3176	6162	4234	5948	5104	5358	9519
LGA50	267	151	219	153	166	166	96
Misassemblies							
# misassemblies	23	1	14	2	17	12	4
Misassembled contigs length	84 193	9611	45 868	6390	111 490	72 879	37 962
Mismatches							
# mismatches per 100 kbp	17	18.78	15	16.71	341.39	15.75	13.49
# indels per 100 kbp	1.21	1.25	1.87	1.94	7.27	1.45	0.87
# N's per 100 kbp	0	0	0	0	0	0	0
Statistics without reference							
# contigs	748	546	684	569	569	584	392
Largest contig	16 612	33 033	21 336	25 068	30 915	30 305	40 471
Total length	2 440 656	2 676 227	2 562 578	2 714 287	2 629 607	2 618 624	2 787 129
Total length (>= 1000 bp)	2 439 127	2 676 227	2 559 569	2 714 287	2 628 029	2 615 105	2 785 415
Total length (>= 10000 bp)	257 236	739 181	320 638	811 392	700 516	658 319	1 419 641
Total length (>= 50000 bp)	0	0	0	0	0	0	0
Extended report							

Assembly: Evaluation - Quast

- Assembly evaluation: Quast, Gurevich et al., *Bioinformatics 2013, 29:8*



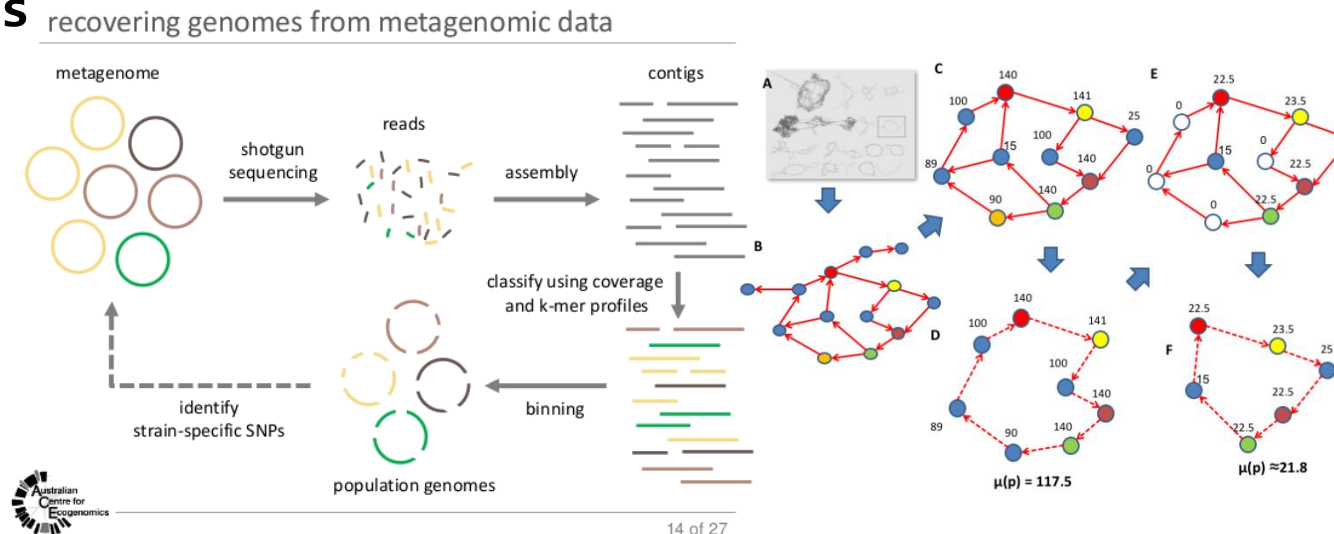
Assembly: Assemblers

Name	Type	Technologies	Author	Presented /Last updated	Licence*	Homepage
DNASTAR Lasergene Genomics Suite	(large) genomes, exomes, transcriptomes, metagenomes, ESTs	Illumina, ABI SOLiD, Roche 454, Ion Torrent, Solexa, Sanger	DNASTAR	2007 / 2016	C	link
Newbler	genomes, ESTs	454, Sanger	454/Roche	2004/2012	C	link
Canu	Small and large, haploid/diploid genomes	PacBio/Oxford Nanopore reads	Koren et al. ^[8]	2001 / 2018	OS	link
SPAdes	(small) genomes, single-cell	Illumina, Solexa, Sanger, 454, Ion Torrent, PacBio, Oxford Nanopore	Bankevich, A et al.	2012 / 2017	OS	link
Velvet	(small) genomes	Sanger, 454, Solexa, SOLiD	Zerbino, D. et al.	2007 / 2011	OS	link

*Licences: OS = Open Source; C = Commercial; C / NC-A = Commercial but free for non-commercial and academics

Assembly: Specials assemblers

- **Diploid genomes**
- **Metagenomics**
- **Plasmids**
- **Transcriptome**



Hybrid genome assembly – short and long reads

1



Long reads: 100s-1000s nucleotides long (from 3rd gen technology)

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

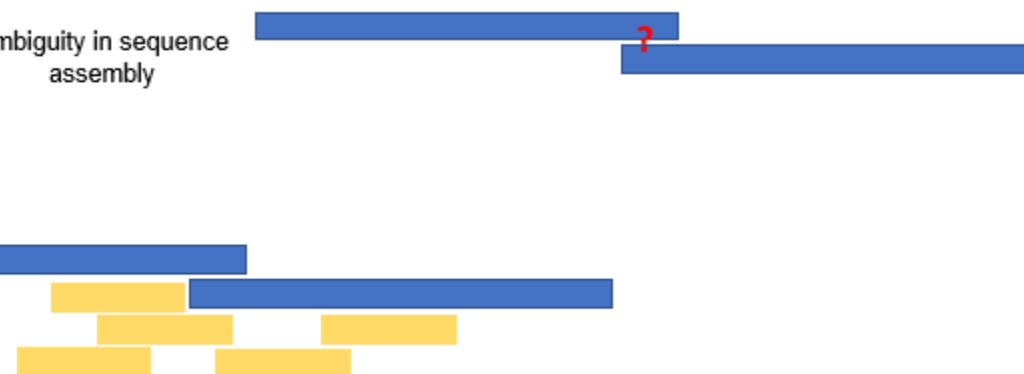
2



Ambiguity in sequence assembly

3

Hybrid assembly helps resolve ambiguities with higher coverage and differing read lengths



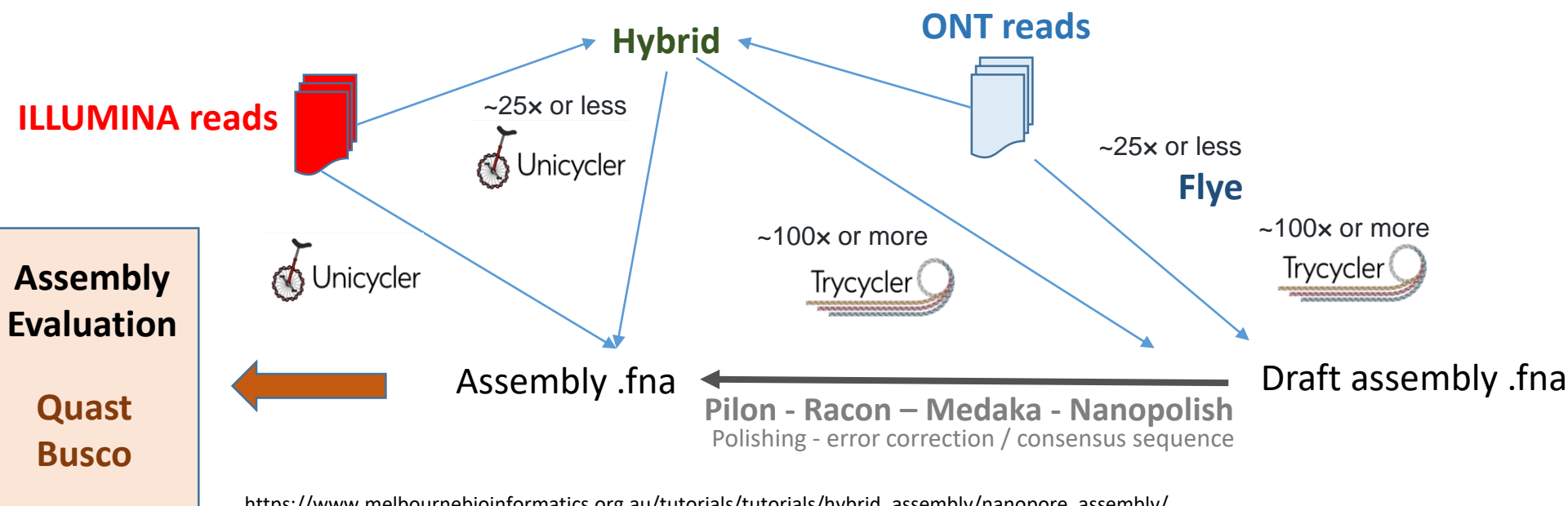
Hybrid genome assembly - nanopore and illumina

Short reads (ILLUMINA) + Long reads (ONT) → deNovo assembly (De novo assembly is the process of assembling a genome from scratch using only the sequenced reads as input - no reference genome is used.) → **high-quality assembly**

ONT: >40.000b, higher error rate – **genome structure**

ILLUMINA: 300b, lower error rate – **high base-level accuracy**

Higher COST

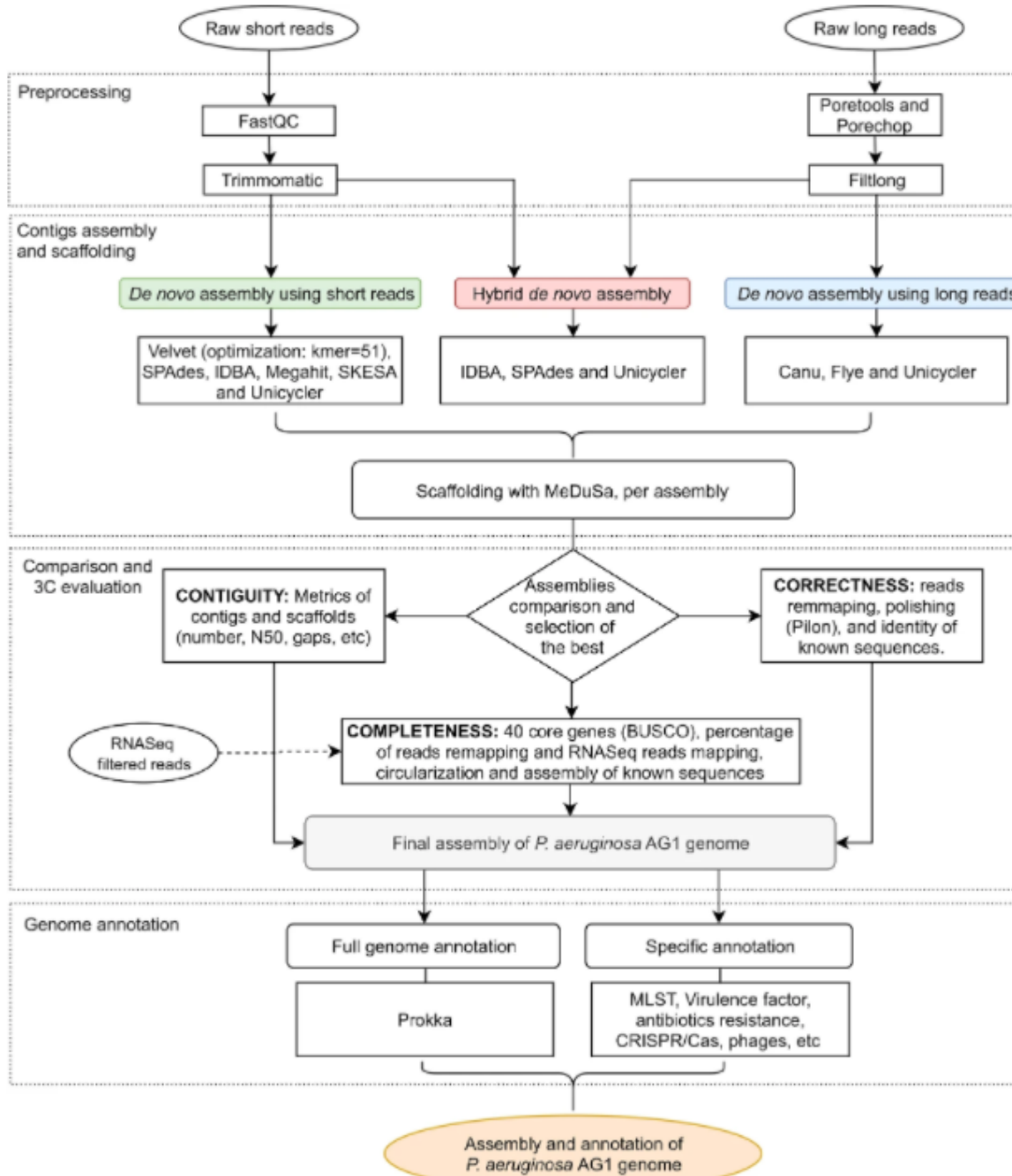


https://www.melbournebioinformatics.org.au/tutorials/tutorials/hybrid_assembly/nanopore_assembly/

<https://github.com/rrwick/Unicycler>

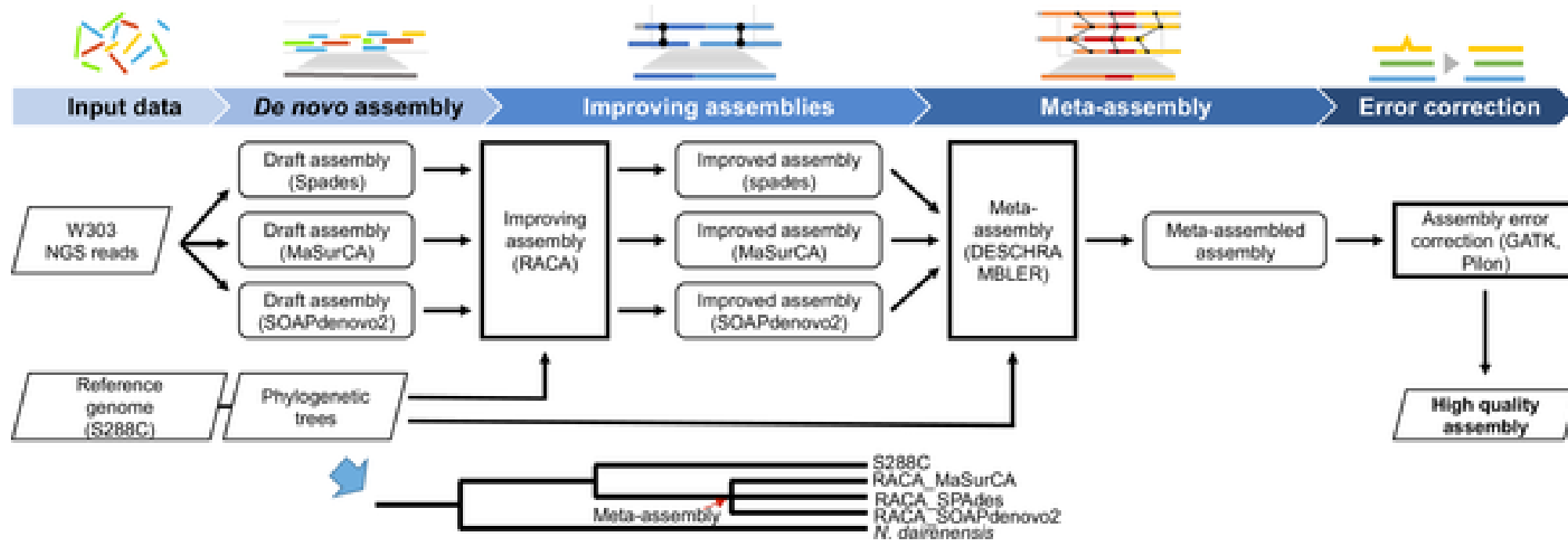
Secuenciación de genomas bacterianos: herramientas y aplicaciones

<https://denbi-nanopore-training-course.readthedocs.io/en/latest/index.html>



Molina-Mora et al.,
Scientific Reports 2020

Fig 1. Data flow chart of the integrative meta-assembly pipeline (IMAP).



Song G, Lee J, Kim J, Kang S, Lee H, et al. (2019) Integrative Meta-Assembly Pipeline (IMAP): Chromosome-level genome assembler combining multiple de novo assemblies. PLOS ONE 14(8): e0221858. <https://doi.org/10.1371/journal.pone.0221858>

Table 1. Assembly evaluation metrics and results.

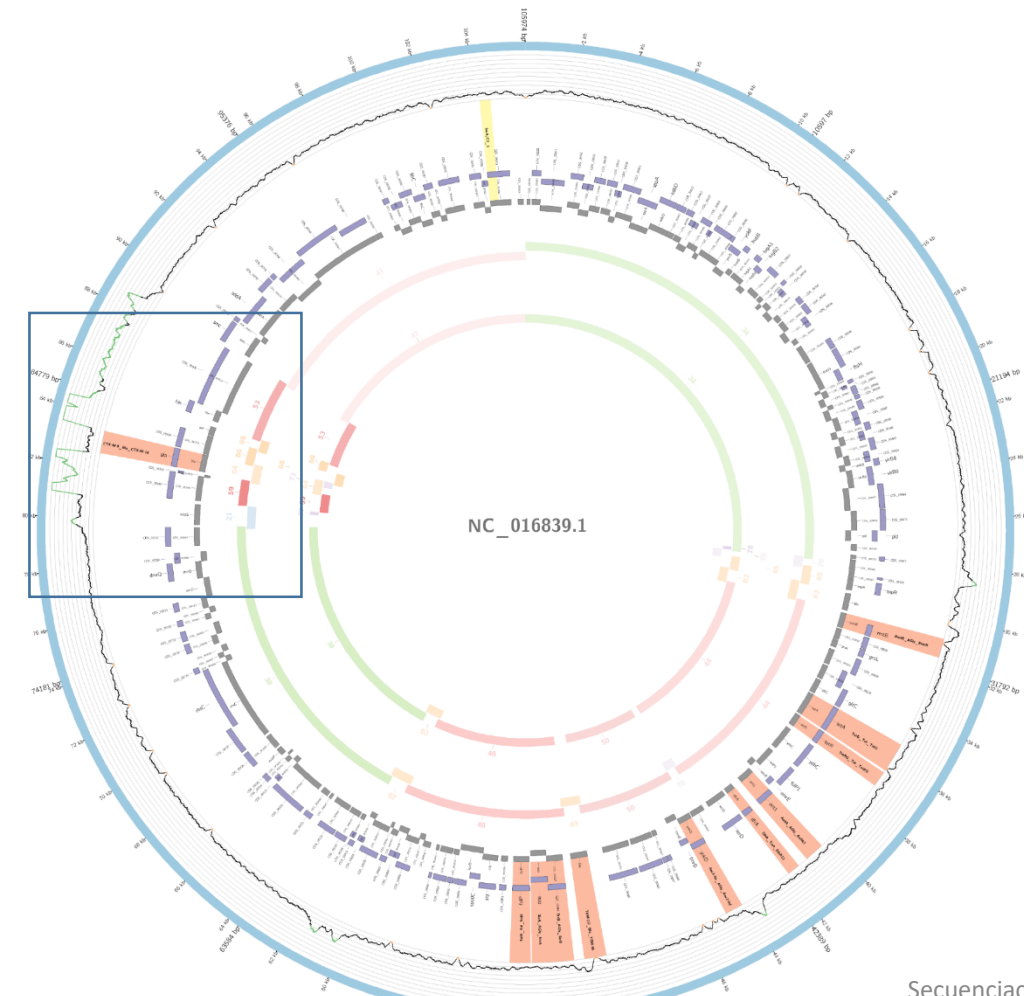
Dataset (W303)		MIN (bp)	MAX (bp)	N50 (bp)	Total length (bp)	Mapped reads (%)	Proper pairs (%)
<i>De novo</i> assembly	SPAdes	80	515,973	187,035	13,901,101	99.40	96.24
	MaSuRCa	300	784,921	273,283	11,838,299	82.86	97.88
	SOAPdenovo2	200	61,911	13,286	11,749,637	50.60	88.48
RACA assembly	RACA-SPAdes	80	1,058,428	716,084	13,905,771	99.40	96.24
	RACA-MaSuRCa	300	1,436,612	706,991	11,842,202	82.86	97.88
	RACA-SOAPdenovo2	200	1,076,849	69,631	11,772,637	50.60	88.49
Meta assembly	Meta	80	1,448,740	702,641	13,773,679	98.56	96.19
Final assembly	Corrected assembly	80	1,450,556	705,629	13,847,490	98.57	97.10
PacBio	PacBio	3,688	1,575,129	929,095	12,433,409	99.15	98.73

<https://doi.org/10.1371/journal.pone.0221858.t001>

Song G, Lee J, Kim J, Kang S, Lee H, et al. (2019) Integrative Meta-Assembly Pipeline (IMAP): Chromosome-level genome assembler combining multiple de novo assemblies. PLOS ONE 14(8): e0221858. <https://doi.org/10.1371/journal.pone.0221858>

<https://journals.plos.org/plosone/article?id=10.1371/journal.pone.0221858>

PlasmidID



Secuenciación de genomas bacterianos: herramientas y aplicaciones

Thanks for your attention!

Questions ?