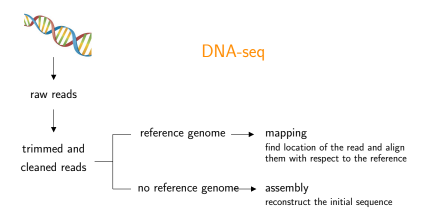


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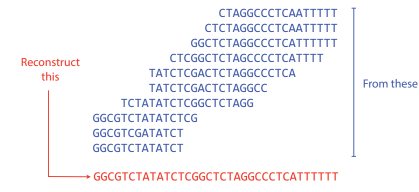
Assembly

Hélène Touzet helene.touzet@univ-lille.fr CNRS, Bonsai, CRIStAL



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The assembly problem



Courtesy of Ben Langmead (Johns Hopkins University)

Why assembling reads?

annotation of genomes

- discovery of new genes
- gene order, structural variants
- noncoding regions
- evolutionnary genomics, phylogenomics
- transcriptome
 - reconstruction of transcripts
 - identification of alternative transcripts

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- metagenomics
 - identification of species

The assembly problem

Reconstruct this CTAGGCCCTCAATTTTT CTCTAGGCCCTCAATTTTT GGCTCTAGGCCCTCATTTTT CTCGGCTCTAGCCCCTCATTTTT TATCTCGACTCTAGGCCCTCA TATCTCGACTCTAGGCC TCTATATCTCGGCTCTAGG GGCGTCTATATCTCG GGCGTCGATATCT GGCGTCTATATCT

From these

<ロト (部) (主) (主) (主)

→ GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTT

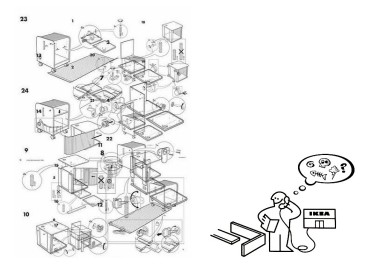
Reconstruct this CTAGGCCCTCAATTTTT GGCGTCTATATCT CTCTAGGCCCTCAATTTTT TCTATATCTCGGCTCTAGG GGCTCTAGGCCCTCATTTTT CTCGGCTCTAGCCCCTCATTTT TATCTCGACTCTAGGCCCTCA GGCGTCGATATCT TATCTCGACTCTAGGCC GGCGTCTATATCTCG

From these

<ロト (部) (主) (主) (主)

→ GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTT





Some assembly is required

How to assemble reads?

Historical perspective

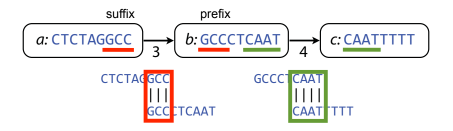


Key features : overlaps between reads

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How to assemble reads?

Historical perspective



Key features : overlaps between reads

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R_1	С	Т	G	А	G	А	А	С	С	Т	G	Т	
R_2	С	С	Т	G	Т	А	А	G	А	Т			
<i>R</i> ₃	С	Т	G	Т	А	С	С	Т					
R_4	G	А	Т	С	Т	G	А						



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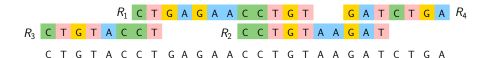


Length of the assembly : 27



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Length of the assembly : 26



Length of the assembly : 27

joining together the reads in decreasing order of the quality of their overlaps

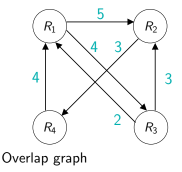


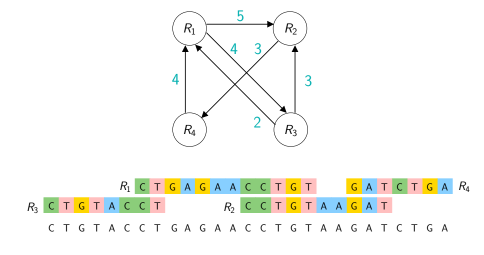
Length of the assembly : 26 trying to maximize the total length of read overlaps



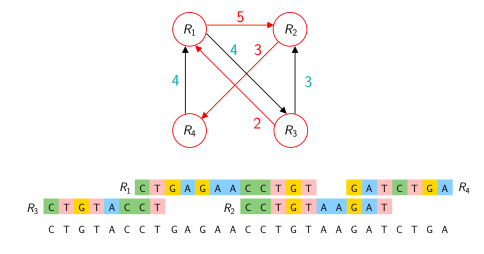
\nearrow	R_1	R_2	R_3	R_4
R_1		5	4	0
R_2	0		0	3
R_3	2	3		0
R_4	4	0	0	

Length of the longest suffix of R_i which is also a prefix of R_j

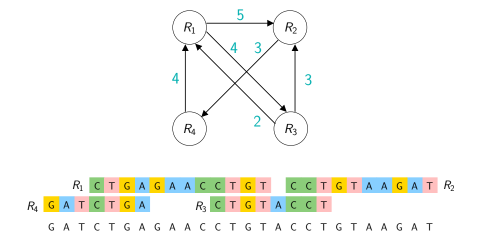




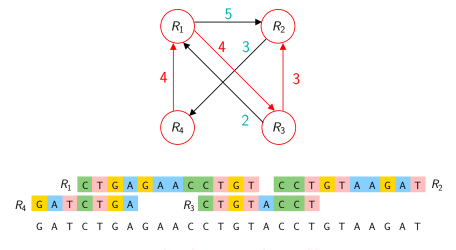
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several paths = several assemblies

Overlap assemblies in real life

- risk of contamination
- existence of sequencing errors
- existence of repeats
- diploid and polyploid genomes
- low coverage or unevent coverage
- + unable to handle the large number of NGS sequencing reads

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Sequencing errors

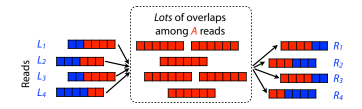
TATCTCGACTCTAGGCC |||||| |||||| TCTATATCTCGGCTCTAGG ↑

• Approximate overlaps : Construction of alignments between reads

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• Assembly : consensus sequence

Repeats



The region A is longer than reads

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Diploidy and polyploidy

Coverage

CTAGGCCCTCAATTTT CTCTAGGCCCTCAATTTTT GGCTCTAGGCCCTCATTTTT CTCGGCTCTAGCCCCTCATTTT TATCTCGACTCTAGGCCCTCA TATCTCGACTCTAGGCC TCTATATCTCGGCTCTAGG GGCGTCTATATCTCG GGCGTCGATATCT GGCGTCTATATCT GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTT

Coverage = 5

CTAGGCCCTCAATTTTT CTCTAGGCCCTCAATTTTT GGCTCTAGGCCCTCATTTTT CTCGGCTCTAGCCCCTCATTTT TATCTCGACTCTAGGCCCTCA TATCTCGACTCTAGGCC 177 bases TCTATATCTCGGCTCTAGG GGCGTCTATATCTCG GGCGTCGATATCT 35 bases GGCGTCTATATCT GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTT

Average coverage = $177 / 35 \approx 5$ -fold

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Overlaps - Historical perspectives

- Sanger sequencing
 - Celera (Myers, 2000) originally developed for the assembly of the human genome
 - SGA (Simpson, Durbin, 2012)
- not suitable for NGS short reads (Illumina)

computationally expensive : construction of the graph, size of the graph, path discovery $% \left({{\left({{{\left({{{\left({{{}}} \right)} \right)}} \right)}_{i}}}} \right)$

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comeback with long reads (Nanopore, Pacbio)

De Bruijn graphs

- introduced in bioinformatics to deal with NGS data
- used by allmost modern short-reads assembly tools
 seminal : Velvet (2008), Abyss (2009), SOAPdenovo2 (2012)
 state-of-the art : SPAdes (2012), MaSuRCA (2013), Megahit (2015)...

Genome assembly reborn : recent computational challenges. M. Pop, Briefings in Bioinformatics 2009 https://doi.org/10.1093/bib/bbp026

How to apply de Bruijn graphs to genome assembly. P.E.C. Compeau, P.A. Pevzner, G. Tesler, Nature Biotechnology 2011 doi:10.1038/nbt.2023

Rationale

• The genome can be reconstructed from the k-mers it contains

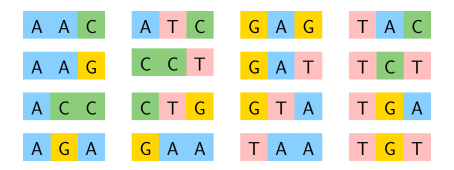
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• Reads are decomposed into k-mers

How many distinct 3-mers are they in

$$R_1$$
CTGAGAACCTGT R_2 CCTGTAAGAT R_3 CTGTACCT R_4 GATCTGA

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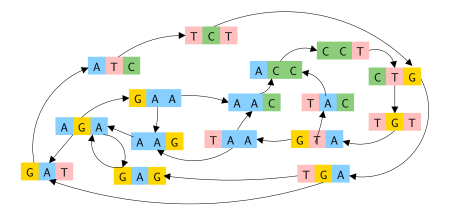
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De Bruijn Graph

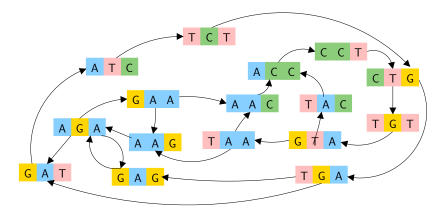
- Nodes : *k*-mers present in the reads
- Arcs : overlaps of length k 1 between k-mers
 Do not depend on the set of reads

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• Easy to construct, low memory footprint Great advantage over overlap graphs



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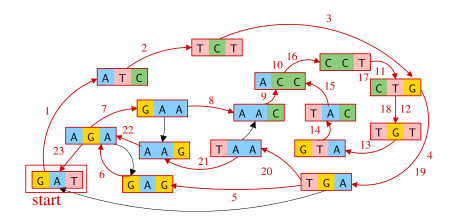


Assembly = path in the graph

Several paths = several assemblies

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De Bruijn Graphs in practice - choice of kLength of k-mers

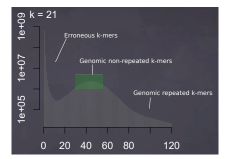
- small *k* :
 - pro : more non-erroneous *k*-mers
 - cons : less signal, more random overlaps, repeat collapsing
- large k :
 - pro : higher signal, less random overlaps, less repeat collapsing

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- cons : more erroneous k-mers
- generally $k \ge 20$ (may be longer for large genomes)
- higher sequencing coverage means larger k values can be used
- multi-k assembly $(k = 21 \rightarrow k = 55 \rightarrow k = 72)$ IDBA, SPAdes, Megahit

In this lecture : SPAdes

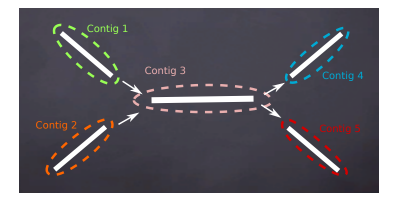
De Bruijn Graphs in practice - cleaning k-mers



Courtesy of Rayan Chikhi (Institut Pasteur) horizontal axis : number of occurrences vertical axis : number of *k*-mers

- k-mers with low frequency are likely to contain sequencing errors
- remove *k*-mers with too few occurrences before the construction of the graph
- k-mer counting : Jellyfish, Kmergenie, DSK

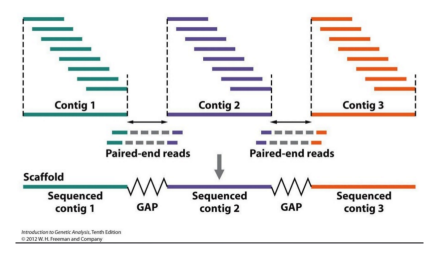
De Bruijn Graphs in practice - contigs and scaffolds



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Contigs =*simple* paths in the graph



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 $\label{eq:scaffold} \begin{array}{l} {\sf Scaffold} = {\sf link} \mbox{ between contigs using paired-end reads} \\ {\sf Error-prone} \end{array}$

Short read assembly is still difficult

even with De Bruijn graphs

- risk of contamination
- existence of sequencing errors solved
- existence of repeats
- diploid and polyploid genomes
- low coverage or unevent coverage

+ unable to handle the large number of NGS sequencing reads $\ensuremath{\textbf{solved}}$

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Short read assembly is still difficult

- library design
 - longest read lengths
 - coverage \geq 50x, x ploidy number
 - for 1 bacterial genome, no point going above 200x
 - BROAD recipe : several mate pairs libraries of increasing size
- assembler
 - SPAdes for small genomes
 - unclear for large genomes
 - try at least two assemblers, try different parameters
 - high computational requirements overall
- an assembly is not the absolute truth, it is a mostly complete, generally fragmented and mostly accurate hypothesis

How to compare/analyse assemblies?

- no trivial ranking between assemblies
- no simple criteria
- assembly with high coverage and short contigs / assembly with low coverage and long contigs

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Quast

Quality Assessment Tool for Genome Assemblies

- provides a large number of statistics and metrics : contigs, missamblies, functional elements
- works both with and without a reference genome
- accepts multiple assemblies, thus is suitable for comparison

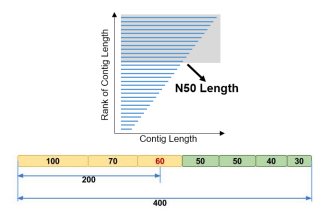
QUAST : quality assessment tool for genome assemblies. Bioinformatics 2013 https://doi.org/10.1093/bioinformatics/btt086

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Contigs

- number of contigs
- length of the largest contig
- total number of bases in the assembly (sum of contig lengths)

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- N50 : contig length N for which 50% of all bases in the sequences are in a contig of length L ≥ N
- *NG*50 : contig length such that using equal or longer length contigs produces 50% of the expected length of the reference genome

Miassemblies (requires a reference genome)

- missassembly breakpoints : position in the contig where the left flanking sequence aligns over 1 kb away from the right flanking sequence on the reference, or they overlap >1 kb, or align on opposite strands or different chromosomes :
- metrics : total number of missambly breakpoints, number of contigs that contain misassembly breakpoints, number of bases contained in all contigs that have one or more misassemblies (Mummer)
- number of unaligned contigs : contigs that have no alignment to the reference sequence
- number of ambiguously mapped contigs : contigs that have multiple alignments to the reference genome

Functional elements

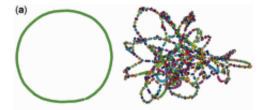
- genome fraction (%) : number of aligned bases in the reference, divided by the genome size.
- duplication ratio : number of aligned bases in the assembly divided by the number of aligned bases in the reference
- number of mismatches and number of indels per 100 kb
- number of genes based on a user-provided annotated list of gene positions in the reference genome
- number of predicted genes in the assembly (GeneMark.hmm for prokaryotes and GlimmerHMM for eukaryotes)

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Bandage

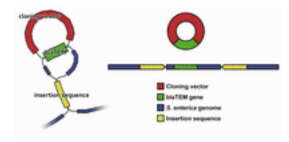
Bioinformatics Application for Navigating De novo Assembly Graphs Easily

- interactive visualization of the assembly graph (such as de Bruijn graph)
- enables BLAST searches



Left : ideal bacterial assembly with one single contig Right : poor assembly with many short contigs

Bandage : interactive visualization of de novo genome assemblies, Bioinformatics 2015 https://doi.org/10.1093/bioinformatics/btv383



Left : Repeated sequences : single nodes with multiple inputs and outputs Right, underlying gene structure deduced from Bandage visualization

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16S rRNA regions of a bacterial genome assembly graph (found in the graph with Blast)

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