# Day 4: Genome Assembly

**MMB-114** 

## Schedule

Day 1: Basics of UNIX and working with the command line

Day 2: Handling of Nanopore/Illumina data

Day 3: Check-up

Day 4: Genome assembly

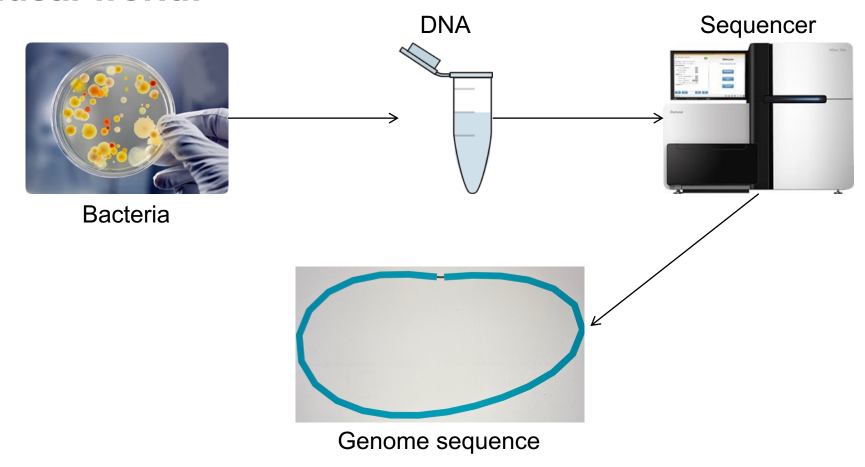
Day 5: Genome annotation

Day 6: Metabolic pathway analysis

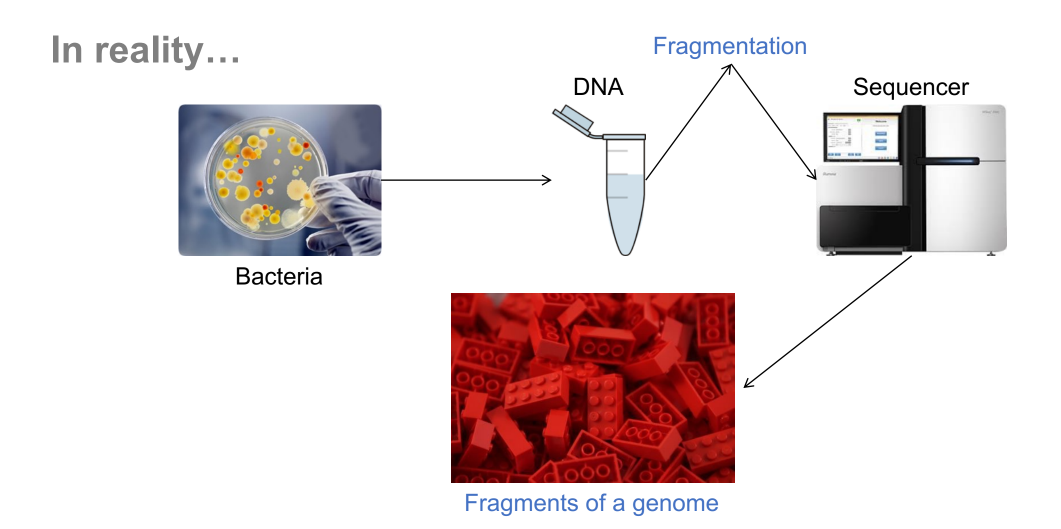


## De novo genome assembly

#### In an ideal world:



# De novo genome assembly



## Genome assembly

 Reconstruct the original genome from long/short sequence reads

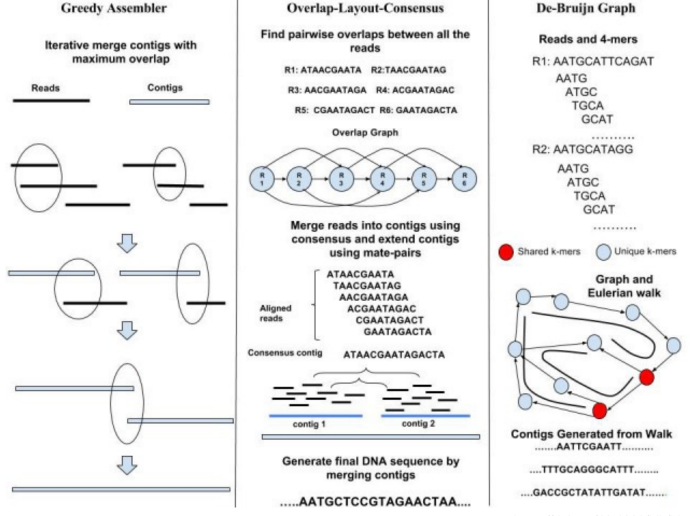
#### **Short reads:**

- In an ideal world: sequences = one complete genome
- In reality: sequences = multiple contigs
  - Contiguous, unambiguous stretches of sequences

## Long reads:

Sometimes we reach the ideal world

# **Assembly strategies**



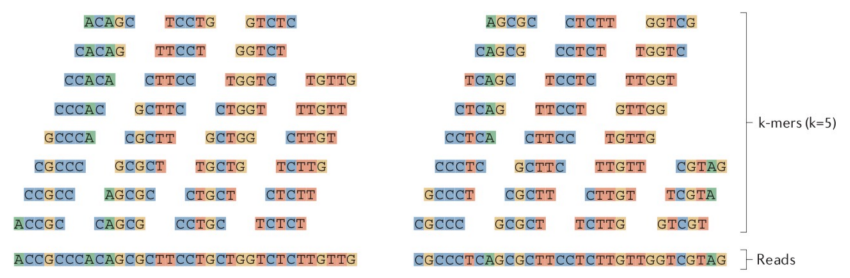
MMB-114 | Antti Karkman

## **Assemblers**

- Bioinformatic tools that combine short sequencing reads into longer contigs
- Spades: kmer-based assembler for short reads (accepts also long reads). De Bruijn graphs.
- Flye: uses repeat graphs. Can tolerate the higher noise of single-molecule sequencing reads.

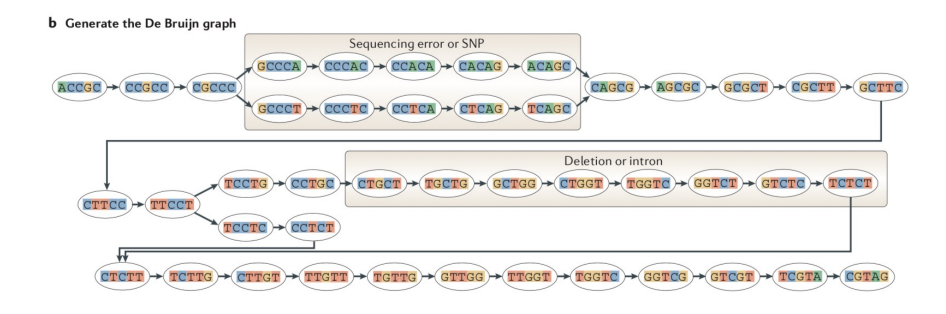
# De Bruijn graph

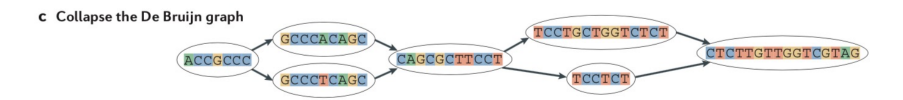
- In real life 10<sup>6</sup> sequences cannot be compared with each other
  - $10^6 \times 10^6 = 10^{12}$  comparisons
- Sequences are reduced to k-mers
  - Smaller subsets of length k



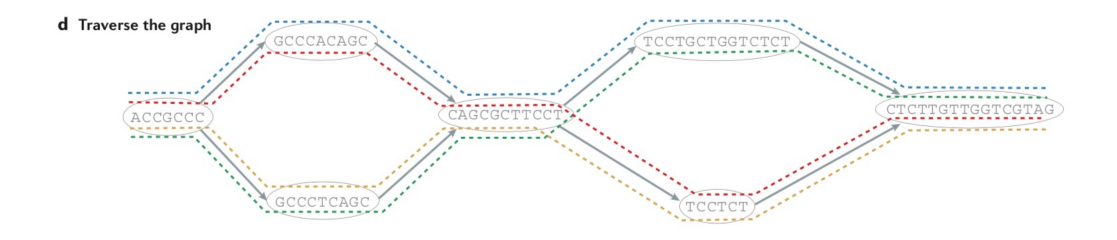
https://doi.org/10.1038/nrg3068

# De Bruijn graph

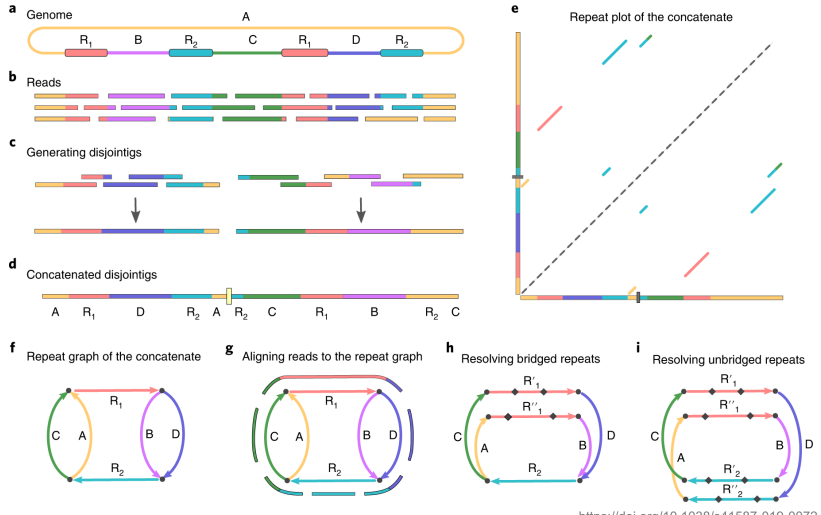




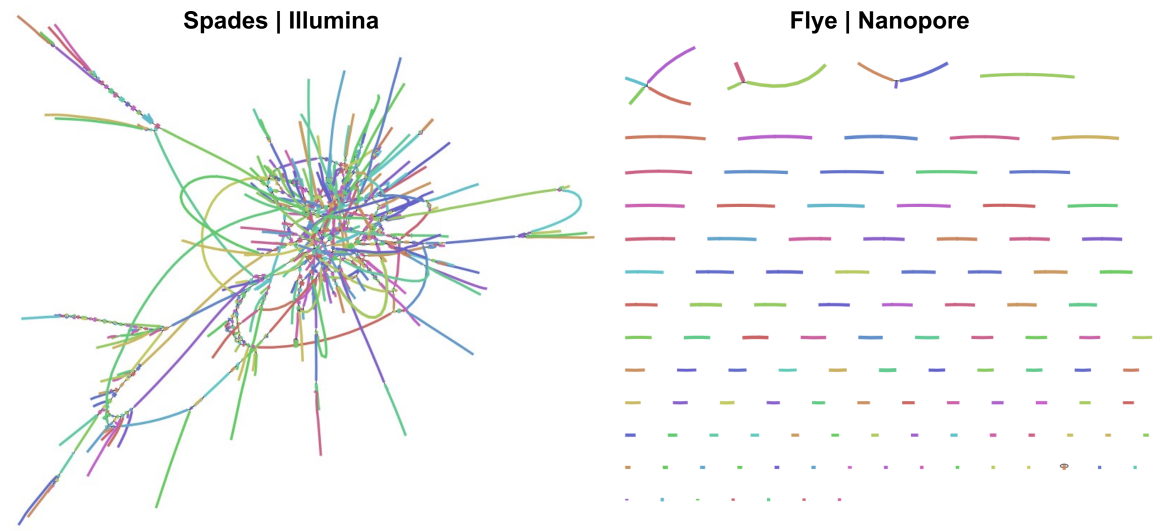
# De Bruijn graph



# Repeat graphs



## Real world assemblies

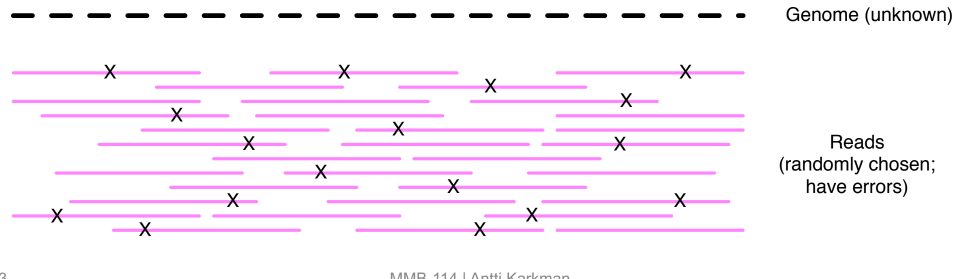


# What makes assembly tricky?

- Many pieces (computationally-intensive)
- Errors in sequence (which one is correct?)
- Missing fragments
- Repetitive fragments
- Multiple copies (rRNA gene as an example)
- Circular genome (no starting point)
- Choice of k-mer
  - Too small → misassembly (anything can assemble)
  - Too long → no assembly

# Sequencing coverage

- Coverage describes the number of time that each base of the genome is present in the reads
- Assemblers expect equal and high enough coverage (>50x) to work optimally in genome assembly.



# Estimated average coverage of your genome

- Your genome size: X bp
- Received sequence data: Y bp

Coverage = Y / X

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# **Assembly quality**

## What is a good assembly?

- Good coverage throughout the contigs
- Correct size (for bacteria, not 400 kb or 14 Mb)
- As few contigs as possible
- Similar GC across contigs

## **Additional notes about Puhti**

## Batch jobs:

sbatch

scancel

squeue

seff

#### batch\_job.sh

```
#!/bin/bash -1
#SBATCH --job-name spades
#SBATCH --output spades out %j.txt
#SBATCH --error spades err %j.txt
#SBATCH --time 1:00:00
#SBATCH --nodes 1
#SBATCH --ntasks-per-node 1
#SBATCH --cpus-per-task 4
#SBATCH --mem 5000
#SBATCH --account project XXX
module load spades
spades.py -1 Data/MMB-114 trimmed 1.fastq.gz \
          -2 Data/MMB-114 trimmed 2.fastq.gz \
          -o SPADES \
          -t 4 \
          --careful
```

sbatch batch\_job.sh

# **Assembly outputs**

## Flye

- assembly.fasta assembled contigs
- assembly graph.gfa assembly graph
- assembly info.txt information about each contig

#### **Spades**

- contigs.fasta assembled contigs
- assembly graph.fastg assembly graph

# Let's assemble your genomes

Go to Github and follow the instructions:

https://github.com/karkman/MMB-114 Genomics

(Day 3: Genome assembly)

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