# Complex-Network Modelling and Inference Lecture 6: Application: Genome Reconstruction

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### Section 1

Genome Reconstruction

#### Genomes

#### **Definition**

A *genome* is the genetic material in an organism.

- Consists of DNA for us (RNA for viruses)
- Made up of sequences of nucleotides
  - ▶ Humans have about 3 billion nucleotides in our genome
- (roughly) DNA has made up of sequences of 4 molecules (the "bases")
  - Adenine
  - Guanine
  - Cytosine
  - ► Thymine

which appear in a double-helix, but in fixed pairs, so we can write a DNA sequence as a sequence of these letters, *e.g.*,

#### *AAGCTTAAGTC*

# Genome sequencing

- We can't just read the sequence
- There are various approaches to reading, but generally
  - they can only see a small part at a time: a "read"
  - we don't know where in the sequence a read comes from
  - if we get lots, then they will have lots of overlaps
- Practicalities
  - real reads contain errors
  - we can't guarantee that reads cover all the genome
  - you don't know which side of the DNA a read comes from

But we will ignore these problems for the moment

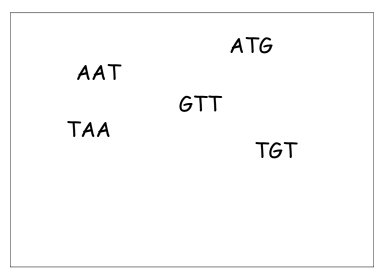
# **Terminolgy**

- A sequence of k symbols is called a k-mer. Also called an n-gram
  - ▶ e.g., in English, 3-mers are sequences like "abc", "rtb", ...
  - ▶ e.g., in binary, 3-mers are sequences like "010", "111", ...
  - ▶ e.g., in DNA, 3-mers are sequences like "ATG", "TTT", ...

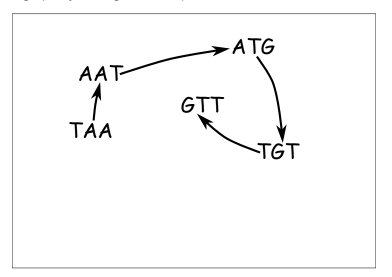
Usually we are interested in sub-sequences from a longer sequence.

- A prefix is the start of a sequence, and a suffix the end
  - lacktriangle when we are dealing with k-mers, we will mean a k-1-prefix or suffix
- Given *n* symbols, how many possible *k*-mers are there?

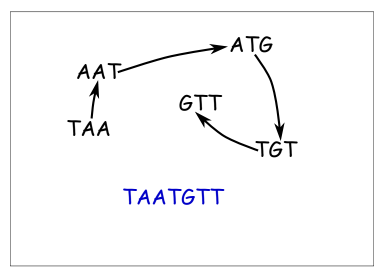
Assume our reads provide all 3-mers from a DNA sequence



Create a graph by linking suffix  $\rightarrow$  prefix



The sequence is just the path through this graph



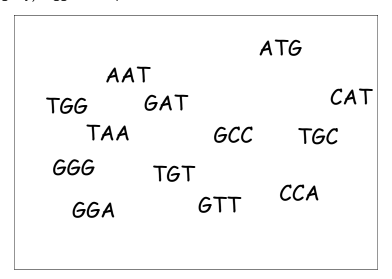
#### Are we done?

#### No!

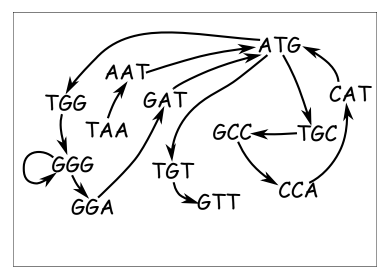
- k-mers are repeated in a long sequence
  - the above assumed that each appeared exactly once in the sequence
  - there can be more than one prefix/suffix match
- Above also assumes
  - we have all k-mers
  - no errors

but for the moment, lets ignore these issues, as the first one is big enough

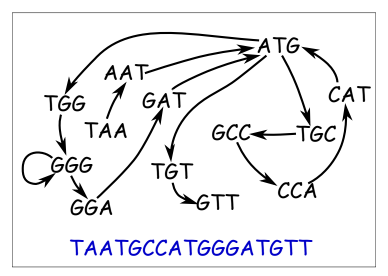
A (slightly) bigger example



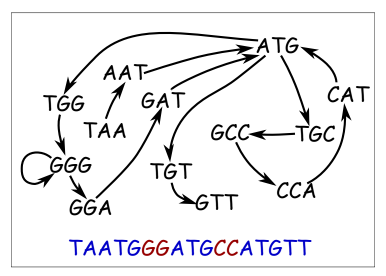
Harder to spot the "walk" (it isn't cycle-free, so not a "path")



Here is one possibility



But there is ambiguity



#### A how to

- If we knew multiplicity of ATG we would create 3 nodes.
  - this simplifies some problems
    - ★ e.g., we know we don't loop through GGG
  - it makes the graph quite a bit more complicated
    - 2 extra nodes
    - lots of extra links
  - it doesn't resolve all ambiguities
    - ★ only way to avoid is to have longer sequences
- Given this new graph the problem becomes one of finding a Hamiltonian path

# Hamiltonian paths and cycles

#### Definition

A *Hamiltonian path* is a path that visits each node exactly once. A *Hamiltonian cycle* visits each node exactly once, and then returns to the start.

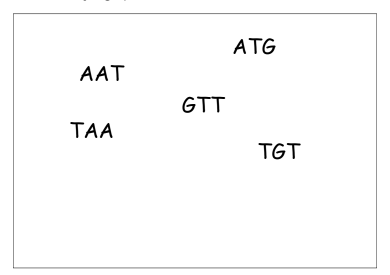
The problem of finding a Hamiltonian path is NP-complete

- this is a hard class of problems (computationally)
- there are no known polynomial-time algorithms
  - but it is easy to check a given cycle is correct
- this is the problem early genome sequencers attempted, but you can't (practically) solve big NP-complete problems
- so they came up with an alternative graph: the de Brujin graph

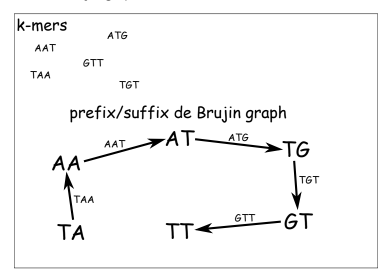
## de Brujin graph

- Often one set of data can be represented by multiple different graphs
- This is a classic example
  - the obvious graph is not the right one to work with
- de Bruijin graph
  - the nodes are prefixes and suffixes
  - ▶ the edges are the *k*-mers
    - ★ they link their prefix to their suffix

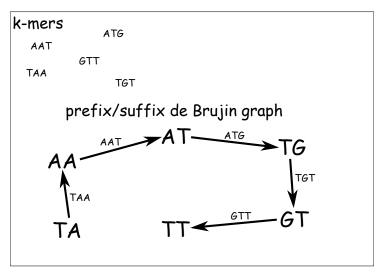
Construct a de Brujin graph



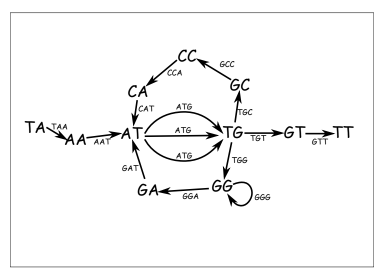
#### Construct a de Brujin graph



Now we just look for an Eulerian path in this graph



de Brujin graph of second example



### Eulerian paths

- The sequence on the de Brujin graph can be found by taking finding an Eulerian path
  - a path that goes along each edge exactly once
  - need to include multiplicity of edges in construction
- Eulerian paths are easy to construct
- We need a minor adaptation here for directed graphs

#### **Definition**

A digraph is *balanced* if the in-degree of each node is the same as its out degree.

#### Theorem

An Eulerian cycle exists on a digraph if and only if it is balanced and strongly connected.

I leave the proof as an exercise, as well as the extension to an Eulerian path.

# Further reading I