### 582670 Algorithms for Bioinformatics

Lecture 5: Graph Algorithms and DNA Sequencing

26.9.2013

# **DNA Sequencing: History**

### Sanger method (1977):

Labeled ddNTPs terminate DNA copying at random points.

#### Gilbert method (1977):

► Chemical method to cleave DNA at specific points (G, G+A, T+C, C).



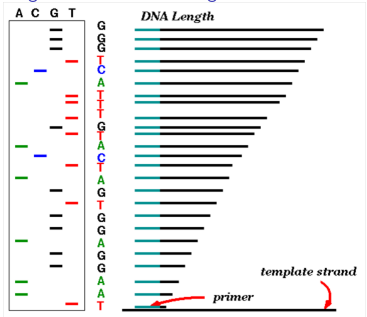
► Both methods generate labeled fragments of varying lengths that are further measured by electrophoresis.



### Sanger Method: Generating a Read

- 1. Divide sample into four.
- 2. Each sample will have available all normal nucleotides and modified nucleotides of one type (A, C, G or T) that will terminate DNA strand elongation.
- 3. Start at primer (restriction site).
- 4. Grow DNA chain.
- 5. In each sample the reaction will stop at all points ending with the modified nucleotide.
- 6. Separate products by length using gel electrophoresis.

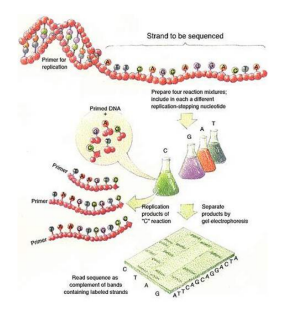
# Sanger Method: Generating a Read



### **DNA Sequencing**

- Shear DNA into millions of small fragments.
- Read 500-700

   nucleotides at a time from the small fragments (Sanger method)



### Fragment Assembly

- ► Computational Challenge: assemble individual short fragments (reads) into a single genomic sequence ("superstring")
- ► Until late 1990s the shotgun fragment assembly of human genome was viewed as intractable problem
  - Now there exists "complete" sequences of human genomes of several individuals
- ► For small and "easy" genomes, such as bacterial genomes, fragment assembly is tractable with many software tools
- ▶ Remains to be difficult problem for more complex genomes

### Shortest Superstring Problem

- Problem: Given a set of strings, find a shortes string that contains all of them
- ▶ Input: Strings  $S = \{s_1, s_2, \dots, s_n\}$
- ▶ Output: A string s that contains all string  $s_1, s_2, \ldots s_n$  as substrings, such that the length of s is minimized
- Complexity: NP-hard
- Recall:
  - Greedy approximation algorithm at the study group
  - Extension to approximate case in the exercises

### Overlaps and prefixes

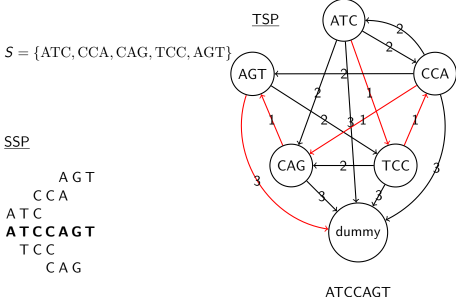
Define  $overlap(s_i, s_j)$  as the longest prefix of  $s_j$  that matches a suffix of  $s_i$   $overlap(s_i, s_j)$  aaaggcatcaatctaaaggcatcaaa aaaggcatcaaatctaaaggcatcaaa  $prefix(s_i, s_j)$ 

▶ Define  $prefix(s_i, s_j)$  as the part of  $s_i$  after its longest overlap with  $s_j$  is removed.

### SSP as a Graph Problem

- Construct a prefix graph with
  - ▶ *n* vertices representing the *n* strings  $s_1, s_2, ..., s_n$  and
  - edges of length  $|prefix(s_i, s_j)|$  between vertices  $s_i$  and  $s_j$
- Add a dummy vertex d to prefix graph with edges of length  $|s_i|$  between each  $s_i$  and d.
- Find the shortest path which visits every vertex exactly once.
- ► This is the Asymmetric Travelling Salesman Problem (ATSP), which is also NP-complete

# SSP to TSP: An example



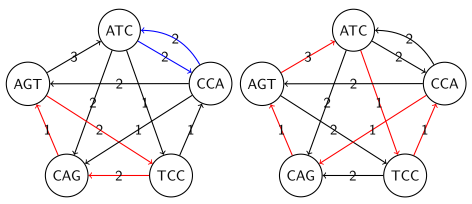
(note: only subset of edges shown) 10 / 32

### Shortest superstring: 4-approximation

- ► There are logarithm-factor approximation algorithms for ATSP, but the prefix graph instances admit constant factor approximations algorithms:
  - ▶ Resulting superstring is at most *c* times longer than the optimal OPT, for some constant *c*.
- 4-approximation algorithm:
  - Construct the prefix graph corresponding to strings in S
  - ► Find a *minimum weight cycle cover* on the prefix graph
  - ▶ Read the superstring defined by the cycle cover
  - Proof of approximation ratio in a study group.

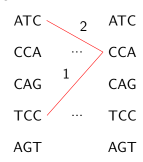
## Cycle cover

- ▶ A cycle cover is a set of disjoint cycles covering all vertices.
- ▶ ATSP tour is a special case: cycle cover with exactly one cycle.



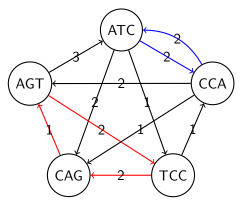
### Minimum weight cycle cover

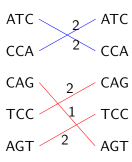
- Minimum weight cycle cover is polynomial time solvable!
- ▶ Reduction to *minimum weight perfect mathing on a bipartite graph*:
  - ▶ Bipartite graph: vertices can be divided into two sets so that all edges have one endpoint in one set and the other endpoint in the other set
  - ▶ Perfect matching: a set of disjoint edges that covers all vertices
  - Create two vertices u<sub>i</sub> and v<sub>i</sub> for each string s<sub>i</sub> to a graph H
- Add edge  $(u_i, v_j)$  with weight  $|prefix(s_i, s_i)|$  for  $i \neq j$
- Each cycle cover in prefix graph corresponds to a minimum weight perfect matching on H and vice versa.



## Minimum weight perfect matching

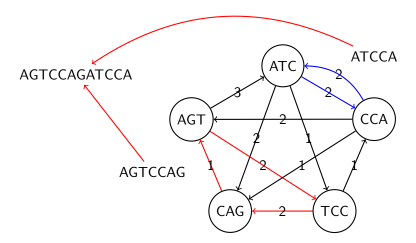
► Classical non-trivial graph problem with polynomial time solutions.





### Reading superstring from cycle cover

- ► For each cycle
  - concatenate prefixes corresponding to weight starting from any vertex
  - append the overlap of last and first vertex
- ► Concatenate the string read from each cycle



# Sequencing by Hybridization (SBH): History

- ▶ 1988: SBH suggested as an alternative sequencing method. Nobody believed it will ever work.
- ▶ 1991: Light directed polymer synthesis developed by Steve Fodor and colleagues.
- 1994: Affymetrix develops first 64-kb DNA microarray.

First microarray prototype (1989)

First commercial DNA microarray prototype with 16,000 features (1994)

500,000 features per chip (2002)



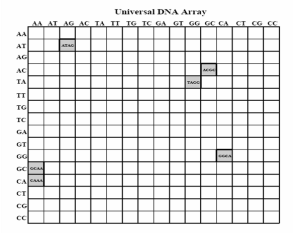




#### How SBH works

- ► Attach all possible DNA probes of length ℓ to a flat surface, each probe at a distinct and known location. This set of probes is called the DNA microarray.
- Apply a solution containing fluorescently labeled DNA fragment to the array.
- ► The DNA fragment hybridizes with those probes that are complementary to substrings of length ℓ of the fragment.
- Using a spectroscopic detector, determine which probes hybridize to the DNA fragment to obtain the ℓ-mer composition of the DNA fragment.
- ▶ Reconstruct the sequence of the DNA fragment from the ℓ-mer composition.

### Hybridization on DNA Array



DNA target TATCCGTTT (complement of ATAGGCAAA) hybridizes to the array of all 4-mers:

> ATAGGCAAA ATAG TAGG AGGC GGCA GCAA

### $\ell$ -mer composition

- ▶ Spectrum( $s, \ell$ ) is a multiset of all possible  $(n \ell + 1) \ell$ -mers in a string s of length n.
- ▶ E.g. for s = TATGGTGC, *Spectrum*(s, 3):

$$S = \{TAT, ATG, TGG, GGT, GTG, TGC\}$$

Different sequences may have the same spectrum:

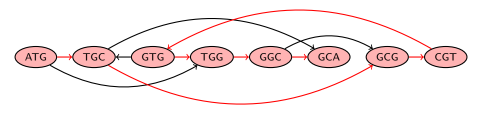
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Spectrum(GTATCT, 2) = Spectrum(GTCTAT, 2) = \{AT, CT, GT, TA, TC\}
```

### The SBH Problem

- ► <u>Goal</u>: Reconstruct a string from its *ℓ*-mer composition
- ▶ Input: A set S, representing all  $\ell$ -mers from an (unknown) string s
- ▶ Output: A string s such that  $Spectrum(s, \ell) = S$

### SBH: Hamiltonian Path Approach

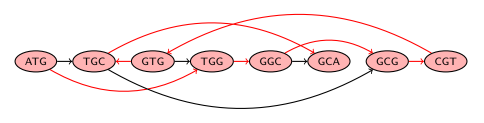
- Construct a graph
  - One vertex for each ℓ-mer in the input spectrum
  - ▶ Draw an edge between two vertices if the  $\ell$ -mers overlap by  $\ell-1$  nucleotides
- Find a path that visits each vertex once.
- ▶ Example:  $S = \{ATG, TGC, GTG, TGG, GGC, GCA, GCG, CGT\}$



**ATGCGTGGCA** 

### SBH: Hamiltonian Path Approach

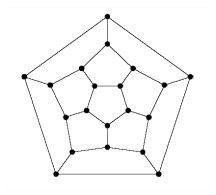
► Another path for: S = {ATG, TGC, GTG, TGG, GGC, GCA, GCG, CGT}



**ATGGCGTGCA** 

### Hamiltonian Cycle Problem

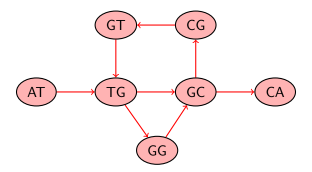
- Find a cycle that visit every vertex exactly once.
- ▶ NP-complete



Game invented by Sir William Hamilton in 1857

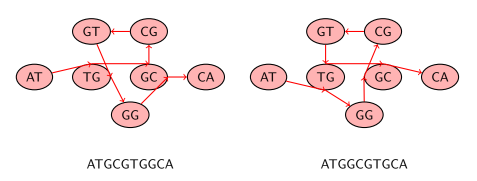
### SBH: Eulerian Path Approach

- Construct a graph
  - ▶ A vertex for each  $(\ell 1)$ -mer
  - ▶ An edge between two vertices corresponds to an  $\ell$ -mer from S
  - Find a path that visits each edge once.
  - ▶ Example:  $S = \{ATG, TGC, GTG, TGG, GGC, GCA, GCG, CGT\}$



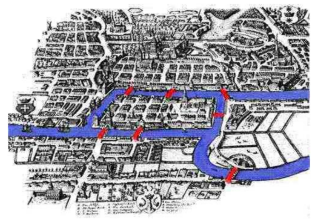
### SBH: Eulerian Path Approach

► *S* = {ATG, TGC, GTG, TGG, GGC, GCA, GCG, CGT} corresponds to two different paths:



### The Bridge Obsession Problem

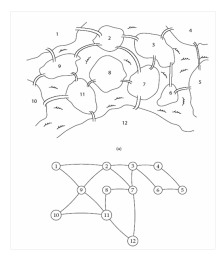
Find a tour crossing every bridge just once Leonhard Euler, 1735



Bridges of Königsberg

### Eulerian Cycle Problem

- ► Find a cycle that visits every edge exactly once
- Linear time



More complicated Königsberg

#### **Euler Theorems**

▶ A graph is *balanced* if for every vertex the number of incoming edges equals the number of ougoing edges:

$$in(v) = out(v)$$

each of its vertices is balanced.

▶ **Theorem**: A connected graph has an *Eulerian cycle* if and only if

- A vertex is semi-balanced if in(v) = out(v) + 1 or in(v) = out(v) 1
- ▶ A graph is *balanced* is for every vertex the number of incoming edges equals the number of ougoing edges:
- ▶ Theorem: A connected graph has an Eulerian path if and only if it contains a vertex v with in(v) = out(v) 1, a vertex w with in(w) = out(w) + 1 and all other vertices are balanced.

### Some Difficulties with SBH

- ► In practice, ℓ-mer composition can never be measured with 100% accuracy
  - ▶ With inaccurate data, the computational problem is again NP-hard.
    - Find minimum completion (insertion/deletion of edges and vertices) of the graph so that it becomes Eulerian
    - Jacek Błazewicz and Marta Kasprzak: Complexity of DNA sequencing by hyridization. *Theoretical Computer Science*, 290(3):1459–1473, 2003.
- Microarray technology has found other uses:
  - Widely used in expression analysis and SNP analysis
- ▶ Virtual ℓ-mer compositions are used in many fragment assembly tools, leading to heuristics exploiting the Eulerian path approach.

### Study Group 1: Lastnames A-K

- ▶ Read pages 284–290 from Jones and Pevzner.
  - ► The peptide sequencing problem
- ▶ At study group draw an example spectrum graph.

### Study Group 2: <u>Last</u>names L-San

- ► Read pages 61–64 from Vazirani: Approximation Algorithms, Springer, 2001.
  - Analysis of the 4-approximation algorithm for Shortest Superstring Problem.
  - Copies distributed at lecture. Ask lecturer for a pdf if you were not present.
- ▶ At study group explain visually the proofs of Lemmas 7.2. and 7.3. Explain how Lemma 7.3 leads to the proof of Theorem 7.4.

### Study Group 3: <u>Last</u>names Sor-Z

- ▶ Read pages 272–275 from Jones and Pevzner.
  - Eulerian cycles and paths.
- ▶ At study group explain the algorithm for finding a Eulerian cycle using an example. How can the algorithm be modified for finding a Eulerian path?