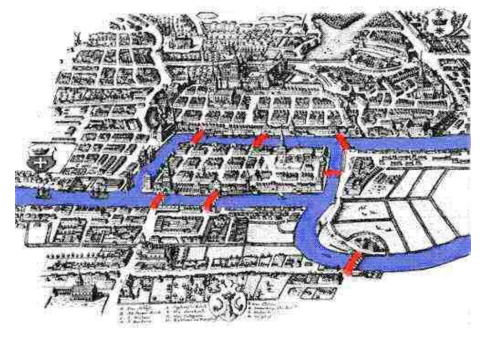
# Graph Algorithms in Bioinformatics

### Outline

- Introduction to Graph Theory
- Eulerian & Hamiltonian Cycle Problems
- Benzer Experiment and Interal Graphs
- DNA Sequencing
- The Shortest Superstring & Traveling Salesman Problems
- Sequencing by Hybridization
- Fragment Assembly and Repeats in DNA
- Fragment Assembly Algorithms

### The Bridge Obsession Problem

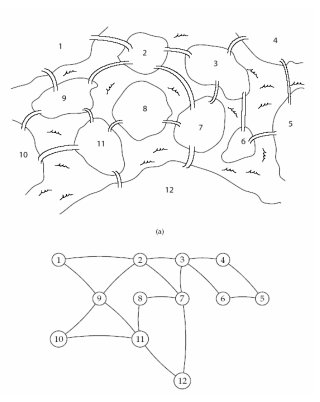
## Find a tour crossing every bridge just once *Leonhard Euler, 1735*



Bridges of Königsberg

### **Eulerian Cycle Problem**

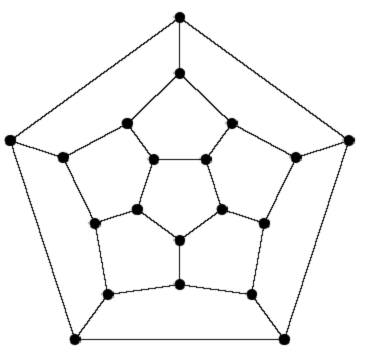
- Find a cycle that visits every edge exactly once
- Linear time



#### More complicated Königsberg

### Hamiltonian Cycle Problem

- Find a cycle that visits every vertex exactly once
- NP complete

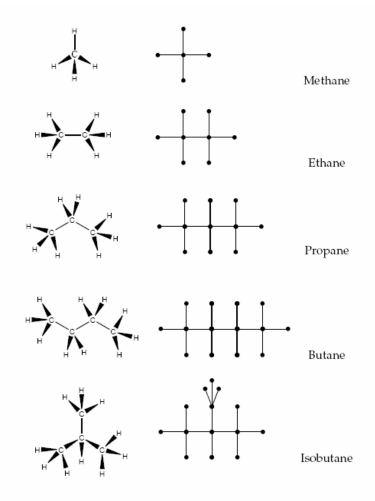


Game invented by Sir William Hamilton in 1857

### Mapping Problems to Graphs

- Arthur Cayley studied chemical structures of hydrocarbons in the mid-1800s
- He used trees

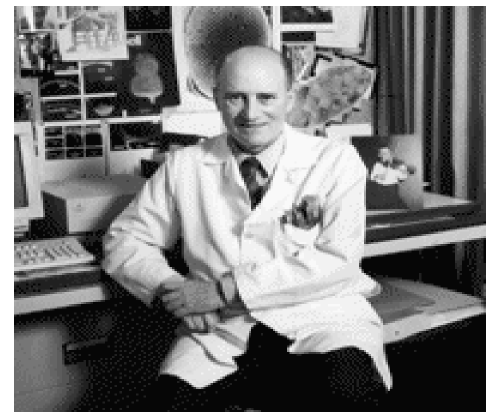
   (acyclic connected graphs) to enumerate structural isomers



### Beginning of Graph Theory in Biology

#### <u>Benzer's work</u>

- Developed deletion mapping
- "Proved" linearity of the gene
- Demonstrated internal structure of the gene



Seymour Benzer, 1950s

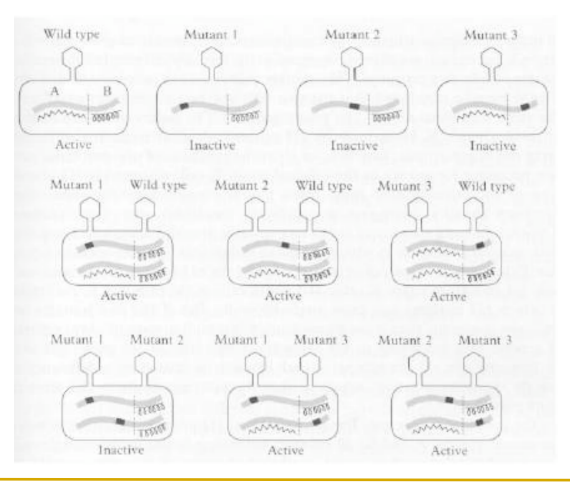
### Viruses Attack Bacteria

- Normally bacteriophage T4 kills bacteria
- However if T4 is mutated (e.g., an important gene is deleted) it gets disable and looses an ability to kill bacteria
- Suppose the bacteria is infected with two different mutants each of which is disabled – would the bacteria still survive?
- Amazingly, a pair of disable viruses can kill a bacteria even if each of them is disabled.
- How can it be explained?

### Benzer's Experiment

- Idea: infect bacteria with pairs of mutant T4 bacteriophage (virus)
- Each T4 mutant has an unknown interval deleted from its genome
- If the two intervals overlap: T4 pair is missing part of its genome and is disabled – bacteria survive
- If the two intervals do not overlap: T4 pair has its entire genome and is enabled – bacteria die

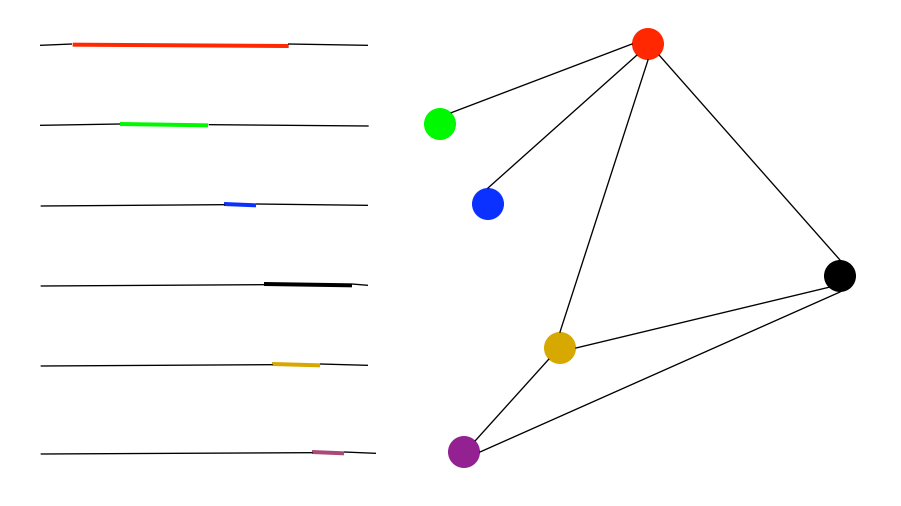
## Complementation between pairs of mutant T4 bacteriophages



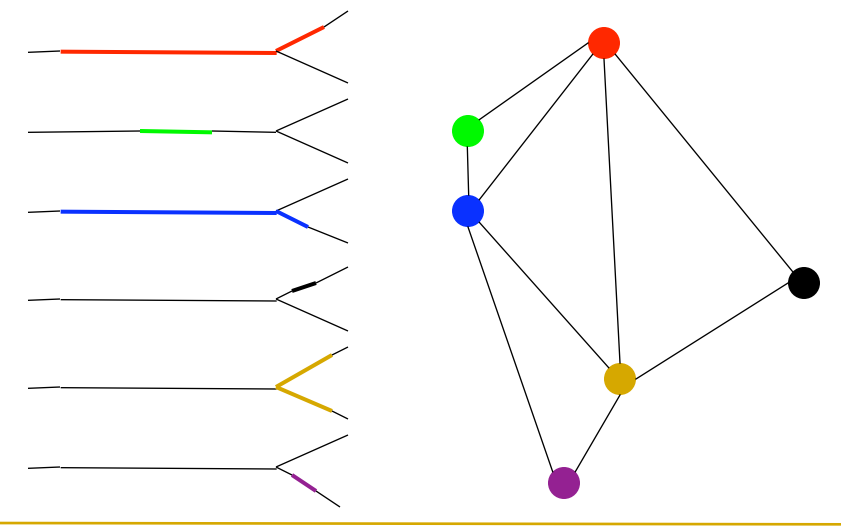
### Benzer's Experiment and Graphs

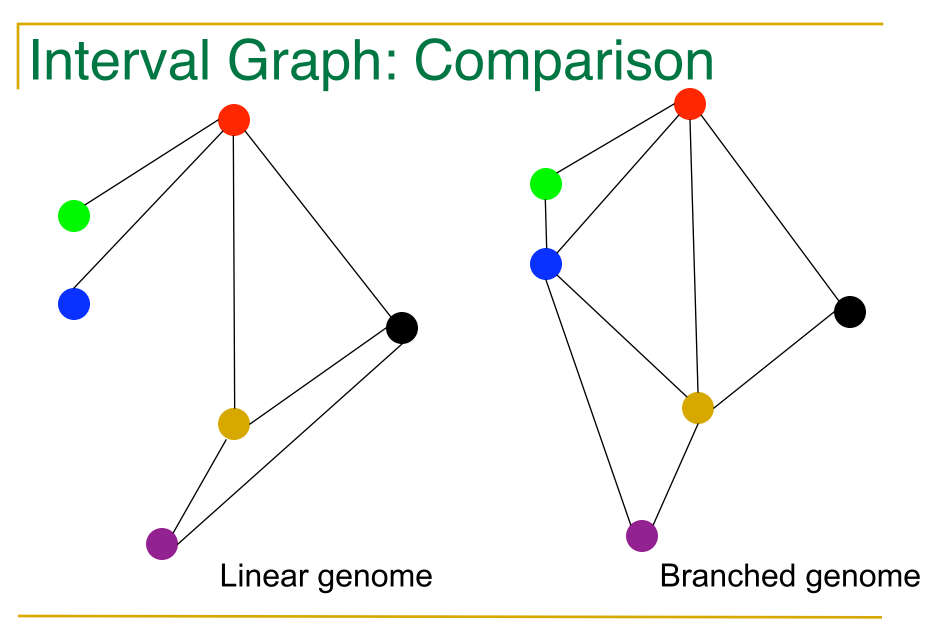
- Construct an interval graph: each T4 mutant is a vertex, place an edge between mutant pairs where bacteria survived (i.e., the deleted intervals in the pair of mutants overlap)
- Interval graph structure reveals whether DNA is linear or branched DNA

### Interval Graph: Linear Genes



### Interval Graph: Branched Genes





### **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).



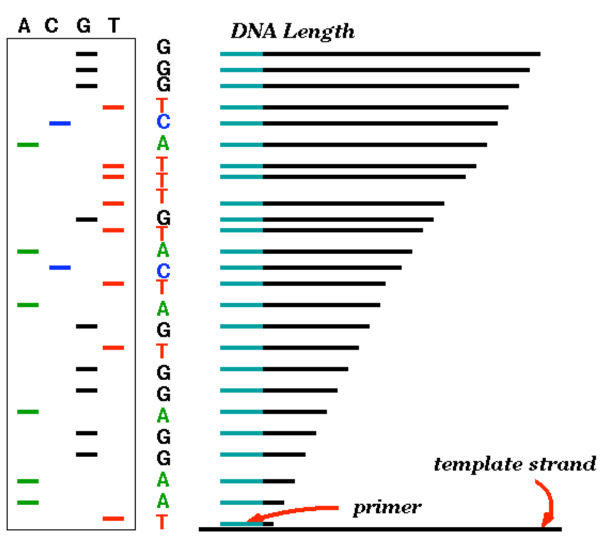
Both methods generate labeled fragments of varying lengths that are further electrophoresed.



### Sanger Method: Generating Read

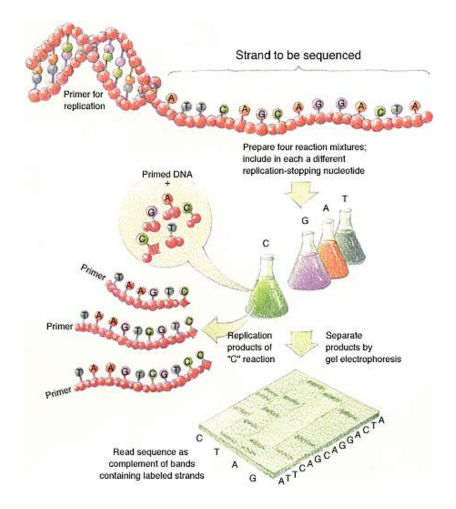


- Start at primer (restriction site)
- <sup>2.</sup> Grow DNA chain
- <sup>3.</sup> Include ddNTPs
- <sup>4.</sup> Stops reaction at all possible points
- <sup>5.</sup> Separate products by length, using gel electrophoresis



### **DNA Sequencing**

- Shear DNA into millions of small fragments
- Read 500 700 nucleotides at a time from the small fragments (Sanger method)



### Fragment Assembly

- <u>Computational Challenge</u>: 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

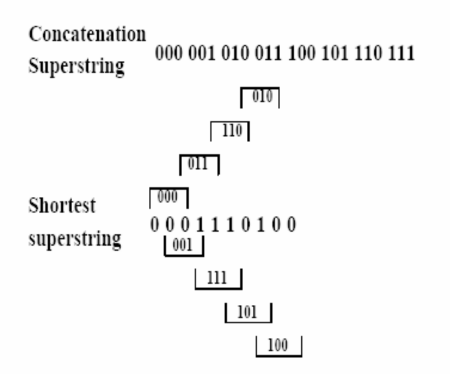
### Shortest Superstring Problem

- <u>Problem</u>: Given a set of strings, find a shortest string that contains all of them
- <u>Input</u>: Strings s<sub>1</sub>, s<sub>2</sub>,..., s<sub>n</sub>
- <u>Output</u>: A string s that contains all strings
   s<sub>1</sub>, s<sub>2</sub>,..., s<sub>n</sub> as substrings, such that the length of s is minimized
- **Complexity:** NP complete
- Note: this formulation does not take into account sequencing errors

#### Shortest Superstring Problem: Example

The Shortest Superstring problem

Set of strings: {000, 001, 010, 011, 100, 101, 110, 111}



### Reducing SSP to TSP

Define overlap (s<sub>i</sub>, s<sub>j</sub>) as the length of the longest prefix of s<sub>i</sub> that matches a suffix of s<sub>i</sub>.

aaaggcatcaaatctaaaggcatcaaa

aaaggcatcaaatctaaaggcatcaaa

What is overlap ( $s_i, s_j$ ) for these strings?

### Reducing SSP to TSP

Define overlap (s<sub>i</sub>, s<sub>j</sub>) as the length of the longest prefix of s<sub>i</sub> that matches a suffix of s<sub>i</sub>.

aaaggcatcaaatctaaaggcatcaaa

aaaggcatcaaatctaaaggcatcaaa

aaaggcatcaaatctaaaggcatcaaa

overlap=12

### Reducing SSP to TSP

Define overlap (s<sub>i</sub>, s<sub>j</sub>) as the length of the longest prefix of s<sub>i</sub> that matches a suffix of s<sub>i</sub>.

aaaggcatcaaatctaaaggcatcaaa

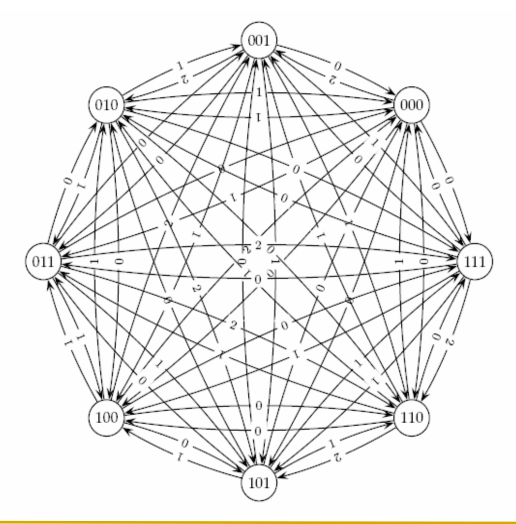
aaaggcatcaaatctaaaggcatcaaa

aaaggcatcaaatctaaaggcatcaaa

- Construct a graph with *n* vertices representing the *n* strings s<sub>1</sub>, s<sub>2</sub>,..., s<sub>n</sub>.
- Insert edges of length overlap (s<sub>i</sub>, s<sub>j</sub>) between vertices s<sub>i</sub> and s<sub>j</sub>.
- Find the shortest path which visits every vertex exactly once. This is the Traveling Salesman Problem (TSP), which is also NP – complete.

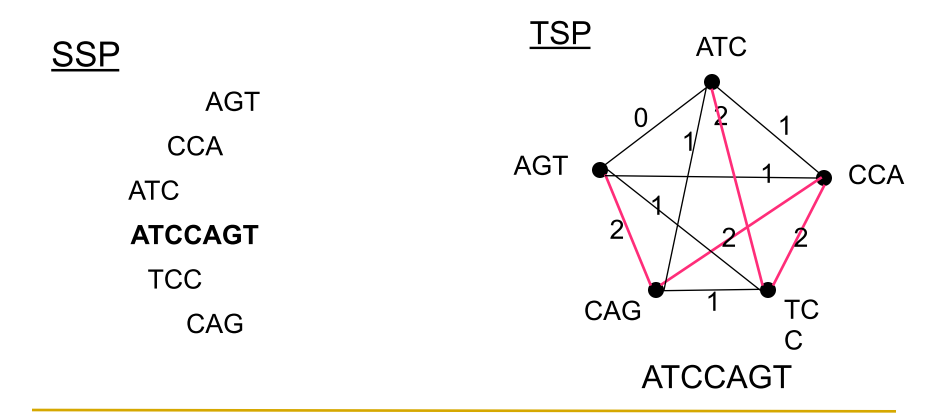
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### Reducing SSP to TSP (cont'd)



### SSP to TSP: An Example

#### S = { ATC, CCA, CAG, TCC, AGT }



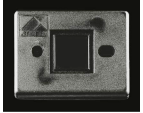
#### Sequencing by Hybridization (SBH): History

• 1988: SBH suggested as an an alternative sequencing method. Nobody believed it will ever work

• **1991:** Light directed polymer synthesis developed by Steve Fodor and colleagues.

> 500.000 features per chip (2002)







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

First microarray prototype (1989)

• **1994:** Affymetrix develops first 64-kb DNA microarray

### How SBH Works

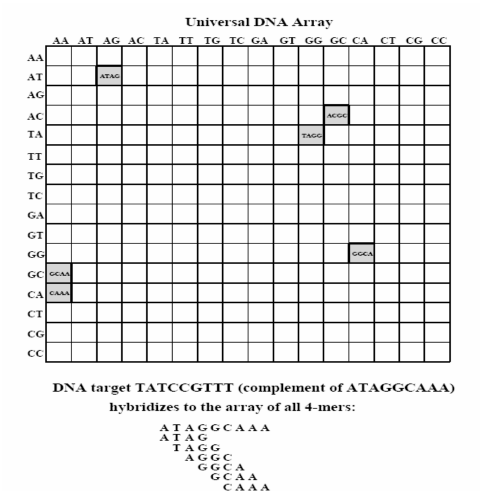
- Attach all possible DNA probes of length / to a flat surface, each probe at a distinct and known location. This set of probes is called the DNA array.
- Apply a solution containing fluorescently labeled DNA fragment to the array.
- The DNA fragment hybridizes with those probes that are complementary to substrings of length / of the fragment.

### How SBH Works (cont'd)

 Using a spectroscopic detector, determine which probes hybridize to the DNA fragment to obtain the *I*-mer composition of the target DNA fragment.

 Apply the combinatorial algorithm (below) to reconstruct the sequence of the target DNA fragment from the *I* – mer composition.

### Hybridization on DNA Array



### *I*-mer composition

- Spectrum (s, I) unordered multiset of all possible (n l + 1) l-mers in a string s of length n
- The order of individual elements in Spectrum (s, l) does not matter
- For s = TATGGTGC all of the following are equivalent representations of Spectrum (s, 3):

{TAT, ATG, TGG, GGT, GTG, TGC} {ATG, GGT, GTG, TAT, TGC, TGG} {TGG, TGC, TAT, GTG, GGT, ATG}

### *I*-mer composition

- Spectrum (s, I) unordered multiset of all possible (n l + 1) l-mers in a string s of length n
- The order of individual elements in Spectrum (s, l) ) does not matter
- For s = TATGGTGC all of the following are equivalent representations of Spectrum (s, 3): {TAT, ATG, TGG, GGT, GTG, TGC} {ATG, GGT, GTG, TAT, TGC, TGG} {TGG, TGC, TAT, GTG, GGT, ATG}
- We usually choose the lexicographically maximal representation as the canonical one.

Different sequences – the same spectrum

Different sequences may have the same spectrum:

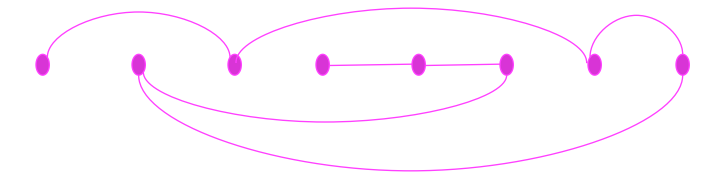
Spectrum(GTATCT,2)= Spectrum(GTCTAT,2)= {AT, CT, GT, TA, TC}

### The SBH Problem

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

### SBH: Hamiltonian Path Approach

#### S = { ATG AGG TGC TCC GTC GGT GCA CAG } ATG AGG TGC TCC GTC GGT GCA CAG



#### ATGCAGGTCC

Path visited every VERTEX once

### SBH: Hamiltonian Path Approach

A more complicated graph:

 $S = \{ATG TGG TGC GTG GGC GCA GCG CGT\}$ 

### SBH: Hamiltonian Path Approach

 $S = \{ATG TGG TGC GTG GGC GCA GCG\}$ CGT }

Path 1:

ATGCGTGGCA

Path 2:

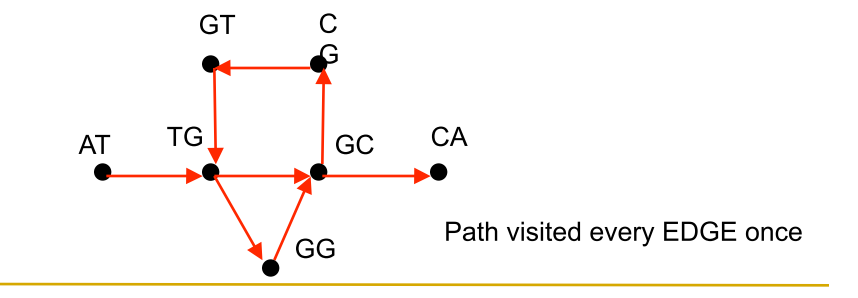
ATGGCGTGCA

#### SBH: Eulerian Path Approach

S = { ATG, TGC, GTG, GGC, GCA, GCG, CGT }

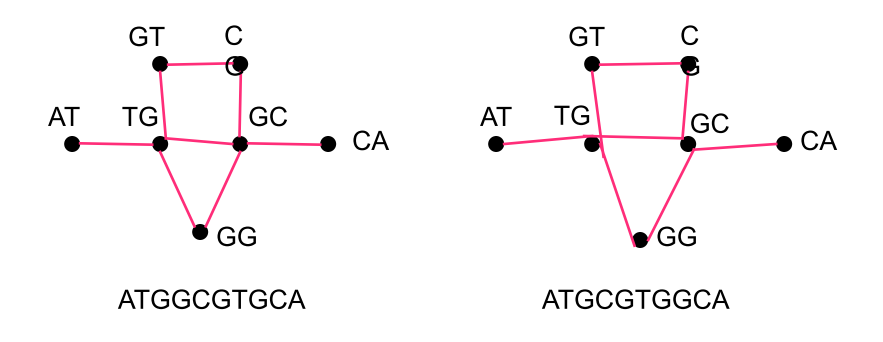
Vertices correspond to  $(I - 1) - mers : \{AT, TG, GC, GG, GT, CA, CG\}$ 

Edges correspond to *I* – mers from S



#### SBH: Eulerian Path Approach

S = { AT, TG, GC, GG, GT, CA, CG } corresponds to two different paths:



#### **Euler Theorem**

 A graph is balanced if for every vertex the number of incoming edges equals to the number of outgoing edges:

*in(v)=out(v)* 

 Theorem: A connected graph is Eulerian if and only if each of its vertices is balanced.

#### Euler Theorem: Proof

• Eulerian  $\rightarrow$  balanced

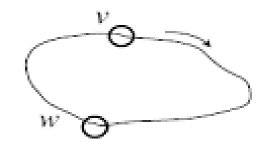
for every edge entering *v* (incoming edge) there exists an edge leaving *v* (outgoing edge). Therefore

*in(v)=out(v)* 

Balanced → Eulerian
 ???

#### Algorithm for Constructing an Eulerian Cycle

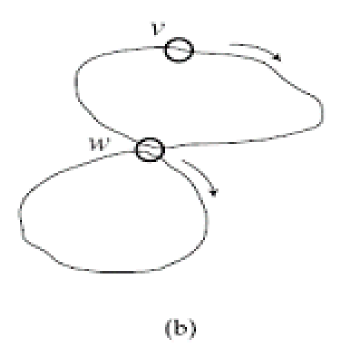
а. Start with an arbitrary vertex v and form an arbitrary cycle with unused edges until a dead end is reached. Since the graph is Eulerian this dead end is necessarily the starting point, i.e., vertex v.



(a)

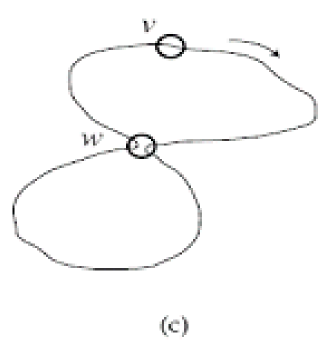
#### Algorithm for Constructing an Eulerian Cycle (cont'd)

If cycle from (a) above is b. not an Eulerian cycle, it must contain a vertex w, which has untraversed edges. Perform step (a) again, using vertex w as the starting point. Once again, we will end up in the starting vertex w.



Algorithm for Constructing an Eulerian Cycle (cont'd)

Combine the cycles
 from (a) and (b) into
 a single cycle and
 iterate step (b).

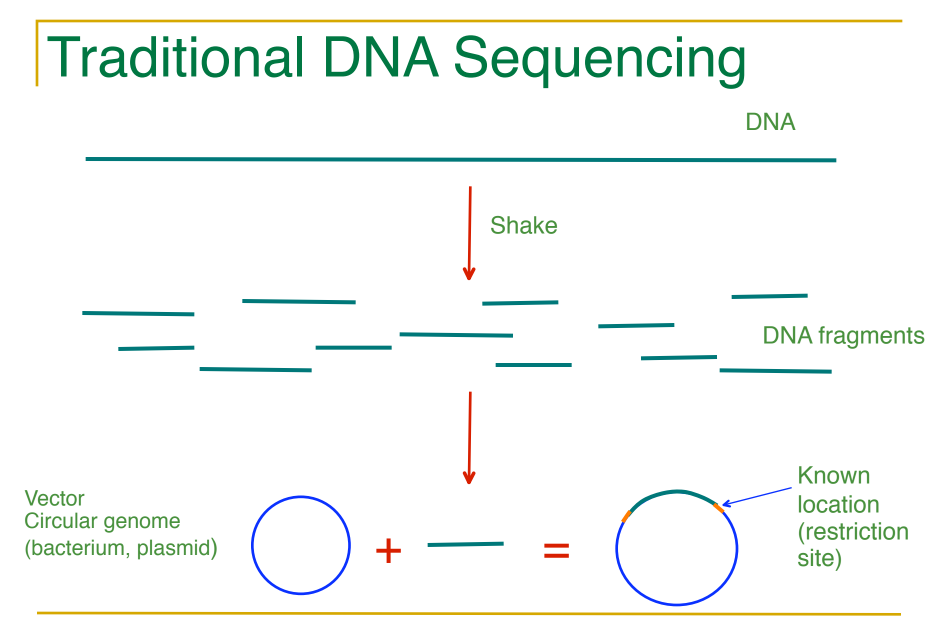


#### Euler Theorem: Extension

 Theorem: A connected graph has an Eulerian path if and only if it contains at most two semi-balanced vertices and all other vertices are balanced.

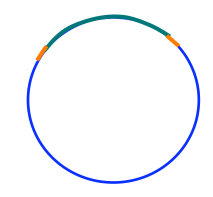
#### Some Difficulties with SBH

- Fidelity of Hybridization: difficult to detect differences between probes hybridized with perfect matches and 1 or 2 mismatches
- Array Size: Effect of low fidelity can be decreased with longer *I*-mers, but array size increases exponentially in *I*. Array size is limited with current technology.
- Practicality: SBH is still impractical. As DNA microarray technology improves, SBH may become practical in the future
- Practicality again: Although SBH is still impractical, it spearheaded expression analysis and SNP analysis techniques



## **Different Types of Vectors**

VECTOR	<u>Size of insert (bp)</u>
Plasmid	2,000 - 10,000
Cosmid	40,000
BAC (Bacterial Artificial Chromosome)	70,000 - 300,000
YAC (Yeast Artificial Chromosome)	> 300,000 Not used much recently



## **Electrophoresis Diagrams**

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# Challenging to Read Answer

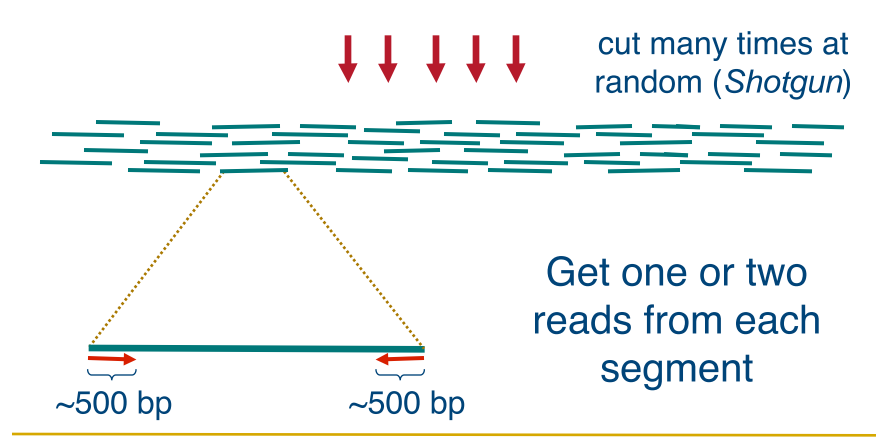
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# Reading an Electropherogram

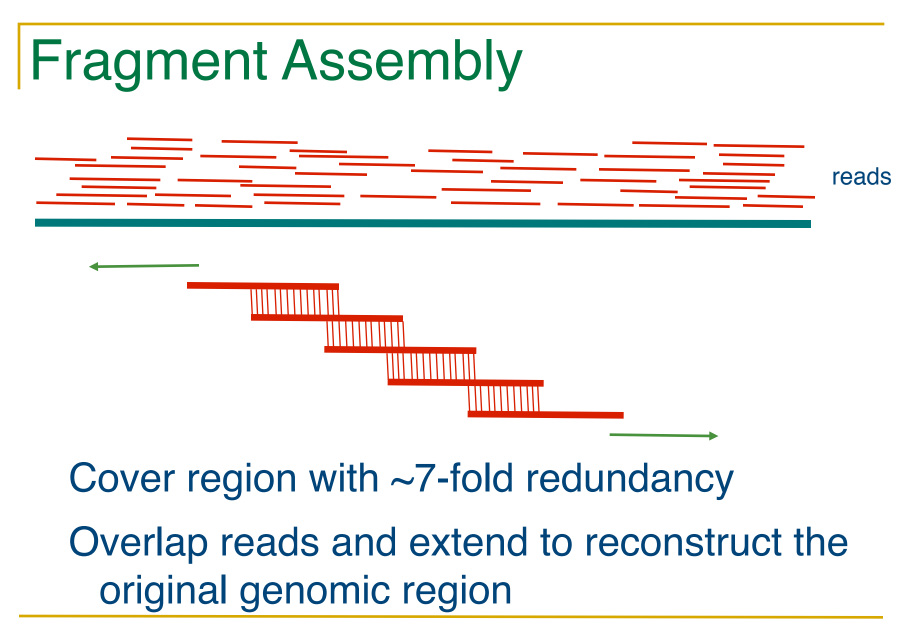
- Filtering
- Smoothening
- Correction for length compressions
- A method for calling the nucleotides PHRED

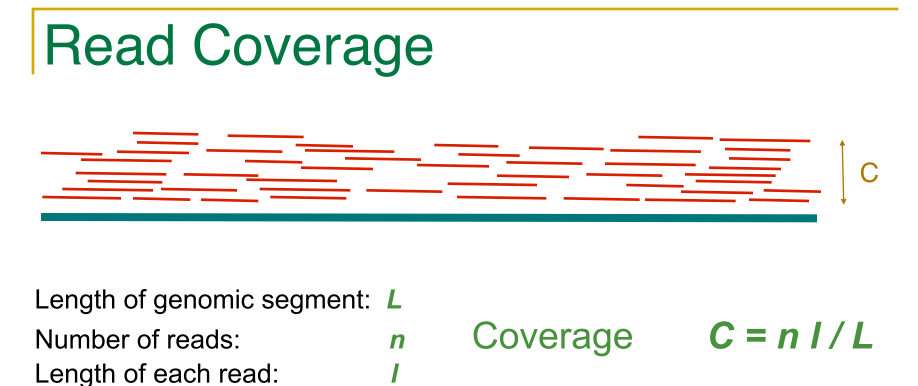
# Shotgun Sequencing

#### genomic segment



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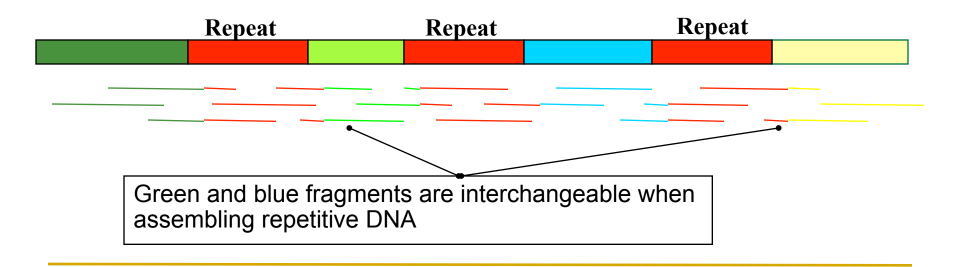
How much coverage is enough?

#### Lander-Waterman model:

Assuming uniform distribution of reads, *C*=10 results in 1 gapped region per 1,000,000 nucleotides

# **Challenges in Fragment Assembly**

- Repeats: A major problem for fragment assembly
- > 50% of human genome are repeats:
  - over 1 million Alu repeats (about 300 bp)
  - about 200,000 LINE repeats (1000 bp and longer)



## Triazzle: A Fun Example

The puzzle looks simple

BUT there are repeats!!!

The repeats make it very difficult.

Try it – only \$7.99 at www.triazzle.com



**Repeat Types** 

- Low-Complexity DNA (e.g. ATATATATACATA...) ٠
- Microsatellite repeats ٠

 $(a_1...a_k)^{N}$  where k ~ 3-6 (e.g. CAGCAGTAGCAGCACCAG)

Transposons/retrotransposons SINE Short Interspersed Nuclear Elements (e.g., Alu:  $\sim$ 300 bp long, 10<sup>6</sup> copies)

- Long Interspersed Nuclear Elements **I INF** • ~500 - 5,000 bp long, 200,000 copies
- LTR retroposons Long Terminal Repeats (~700 bp) at • each end **Gene Families** genes duplicate & then diverge
- Segmental duplications

~very long, very similar copies

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## Overlap-Layout-Consensus

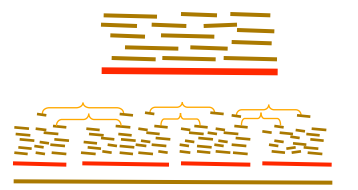
#### Assemblers: ARACHNE, PHRAP, CAP, TIGR, CELERA

**Overlap:** find potentially overlapping reads

*Layout:* merge reads into contigs and contigs into supercontigs

**Consensus:** derive the DNA sequence and correct read errors

..ACGATTACAATAGGTT..



## Overlap

- Find the best match between the suffix of one read and the prefix of another
- Due to sequencing errors, need to use dynamic programming to find the optimal overlap alignment
- Apply a filtration method to filter out pairs of fragments that do not share a significantly long common substring

## **Overlapping Reads**

- Sort all *k*-mers in reads  $(k \sim 24)$
- Find pairs of reads sharing a k-mer
- Extend to full alignment throw away if not >95% similar

TACA TAGATTACACAGATTAC T GA

**Overlapping Reads and Repeats** 

- A k-mer that appears N times, initiates N<sup>2</sup> comparisons
- For an Alu that appears 10<sup>6</sup> times à 10<sup>12</sup> comparisons too much
- Solution:

Discard all *k*-mers that appear more than t´Coverage, (t ~ 10)

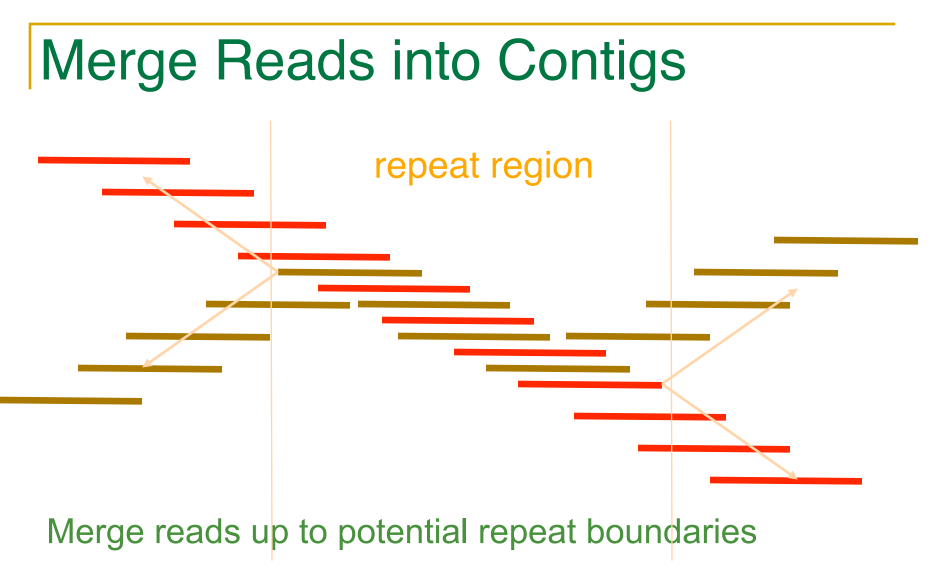
# Finding Overlapping Reads

# Create local multiple alignments from the overlapping reads



#### Layout

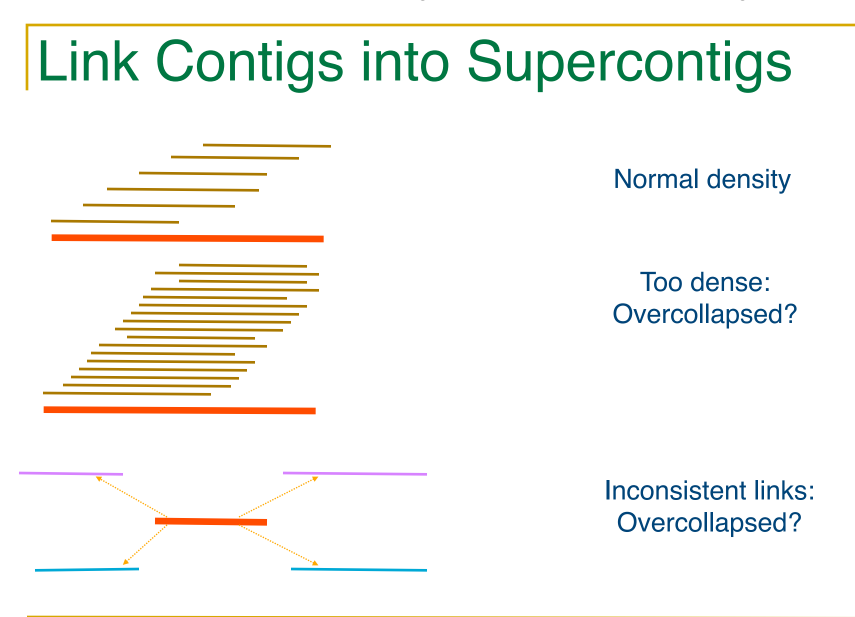
- Repeats are a major challenge
- Do two aligned fragments really overlap, or are they from two copies of a repeat?
- Solution: repeat masking hide the repeats!!!
- Masking results in high rate of misassembly (up to 20%)
- Misassembly means alot more work at the finishing step



#### Repeats, Errors, and Contig Lengths

- Repeats shorter than read length are OK
- Repeats with more base pair differencess than sequencing error rate are OK
- To make a smaller portion of the genome appear repetitive, try to:
  - Increase read length
  - Decrease sequencing error rate

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#### Consensus

- A consensus sequence is derived from a profile of the assembled fragments
- A sufficient number of reads is required to ensure a statistically significant consensus
- Reading errors are corrected

#### **Derive Consensus Sequence**

TAGATTACACAGATTACTGA TTGATGGCGTAA CTA TAGATTACACAGATTACTGACTTGATGGCGTAAACTA TAG TTACACAGATTATTGACTTCATGGCGTAA CTA TAGATTACACAGATTACTGACTTGATGGCGTAA CTA TAGATTACACAGATTACTGACTTGATGGGGGTAA CTA

TAGATTACACAGATTACTGACTTGATGGCGTAA CTA

Derive multiple alignment from pairwise read alignments

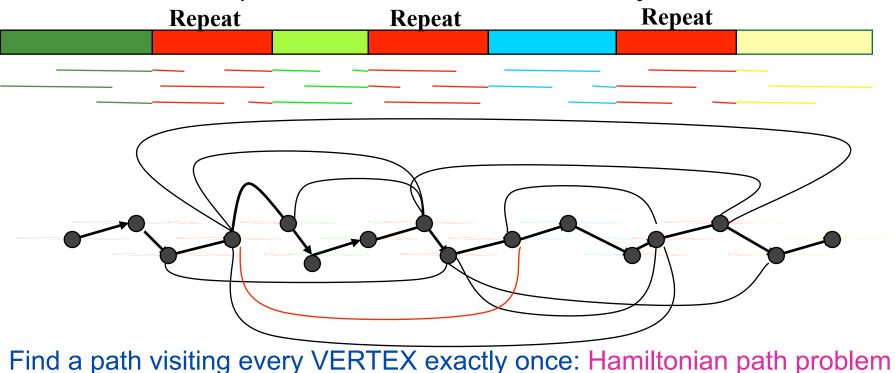
Derive each consensus base by weighted voting

#### EULER - A New Approach to Fragment Assembly

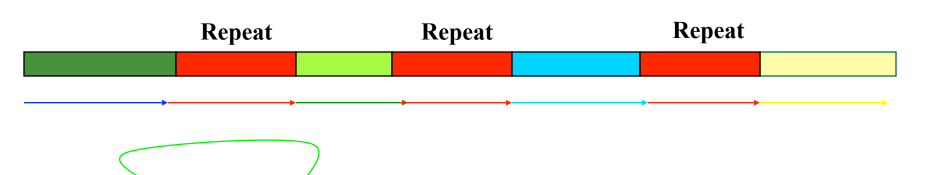
- Traditional "overlap-layout-consensus" technique has a high rate of mis-assembly
- EULER uses the Eulerian Path approach borrowed from the SBH problem
- Fragment assembly without repeat masking can be done in linear time with greater accuracy

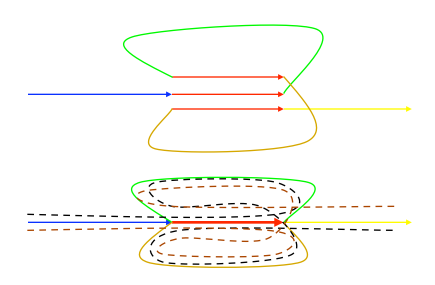
#### Overlap Graph: Hamiltonian Approach

Each vertex represents a read from the original sequence. Vertices from repeats are connected to many others.



#### **Overlap Graph: Eulerian Approach**



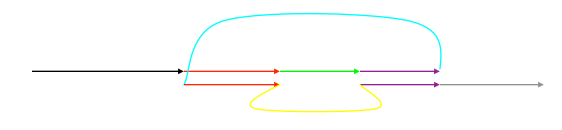


Placing each repeat edge together gives a clear progression of the path through the entire sequence.

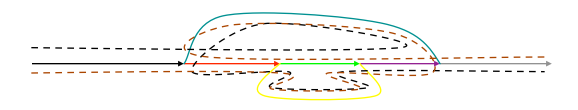
Find a path visiting every EDGE exactly once: Eulerian path problem

#### **Multiple Repeats**





Can be easily constructed with any number of repeats



## **Construction of Repeat Graph**

 <u>Construction of repeat graph from k – mers</u>: emulates an SBH experiment with a huge (virtual) DNA chip.

<u>Breaking reads into k – mers</u>: Transform sequencing data into virtual DNA chip data.

#### Construction of Repeat Graph (cont'd)

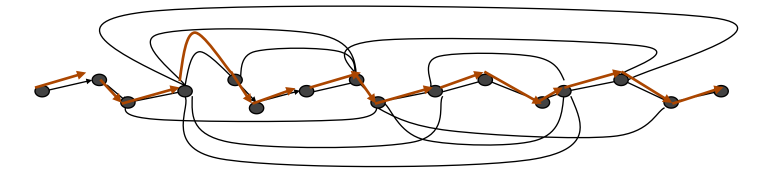
 Error correction in reads: "consensus first" approach to fragment assembly. Makes reads (almost) error-free BEFORE the assembly even starts.

Using reads and mate-pairs to simplify the repeat graph (Eulerian Superpath Problem).

**Approaches to Fragment Assembly** 

Find a path visiting every VERTEX exactly once in the OVERLAP graph:

Hamiltonian path problem

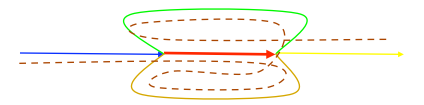


NP-complete: algorithms unknown

Approaches to Fragment Assembly (cont'd)

Find a path visiting every EDGE exactly once in the REPEAT graph:

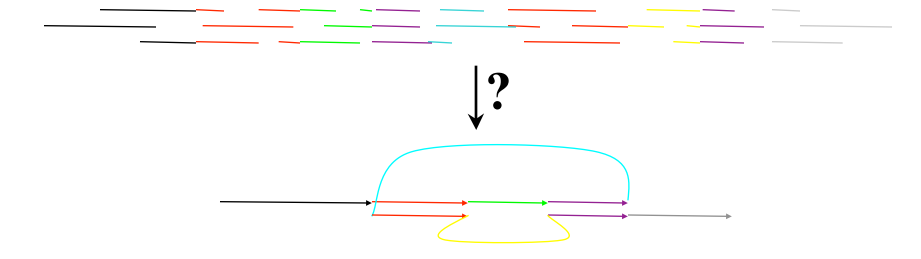
Eulerian path problem



Linear time algorithms are known

#### Making Repeat Graph Without DNA

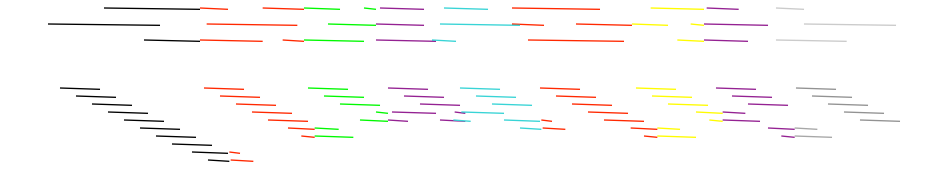
Problem: Construct the repeat graph from a collection of reads.



Solution: Break the reads into smaller pieces.

# Repeat Sequences: Emulating a DNA Chip

 Virtual DNA chip allows the biological problem to be solved within the technological constraints.

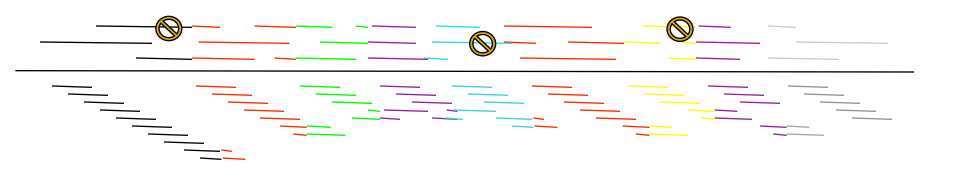


# Repeat Sequences: Emulating a DNA Chip (cont'd)

- Reads are constructed from an original sequence in lengths that allow biologists a high level of certainty.
- They are then broken again to allow the technology to sequence each within a reasonable array.

# **Minimizing Errors**

 If an error exists in one of the 20-mer reads, the error will be perpetuated among all of the smaller pieces broken from that read.



## Minimizing Errors (cont'd)

- However, that error will not be present in the other instances of the 20-mer read.
- So it is possible to eliminate most point mutation errors before reconstructing the original sequence.

#### Conclusions

- Graph theory is a vital tool for solving biological problems
- Wide range of applications, including sequencing, motif finding, protein networks, and many more

#### References

- Simons, Robert W. Advanced Molecular Genetics Course, UCLA (2002). http://www.mimg.ucla.edu/bobs/C159/ Presentations/Benzer.pdf
- Batzoglou, S. Computational Genomics Course, Stanford University (2004). http://www.stanford.edu/class/cs262/ handouts.html