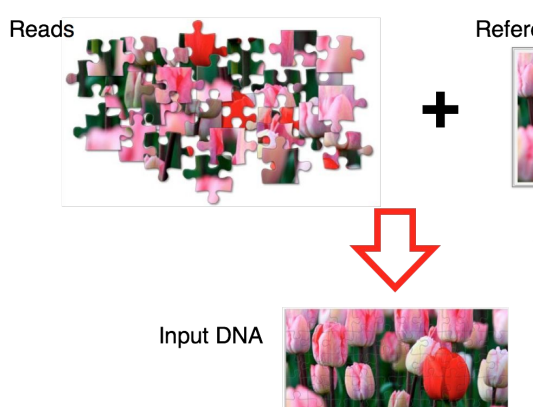
Shortest common superstring

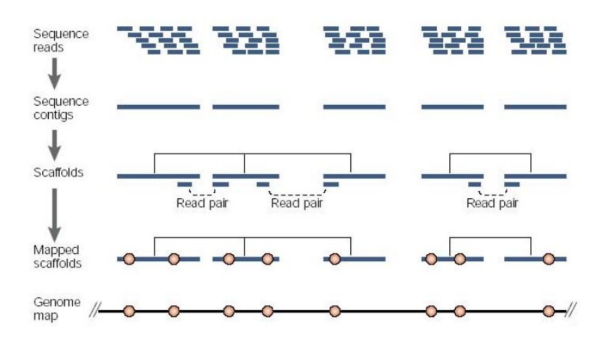


Reference genome



How to assemble puzzle without the benefit of knowing what the finished product looks like?

de novo whole-genome shotgun assembly



Whole-genome "shotgun" sequencing starts by copying and fragmenting the DNA

("Shotgun" refers to the random fragmentation of the whole genome; like it was fired from a shotgun)

Input: GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTTT

Copy: GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTTT
GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTTT
GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTTT
GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTTT

Fragment: GGCGTCTA TATCTCGG CTCTAGGCCCTC ATTTTTT
GGC GTCTATAT CTCGGCTCTAGGCCCTCA TTTTTT
GGCGTC TATATCT CGGCTCTAGGCCCT CATTTTTT
GGCGTCTAT ATCTCGGCTCTAG GCCCTCA TTTTTT

Assume sequencing produces such a large number fragments that almost all genome positions are *covered* by many fragments...

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

From these

GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTTT

...but we don't know what came from where

CTAGGCCCTCAATTTTT GGCGTCTATATCT CTCTAGGCCCTCAATTTTT TCTATATCTCGGCTCTAGG Reconstruct GGCTCTAGGCCCTCATTTTTT this From these CTCGGCTCTAGCCCCTCATTTT TATCTCGACTCTAGGCCCTCA GGCGTCGATATCT TATCTCGACTCTAGGCC GGCGTCTATATCTCG GGCGTCTATATCTCGGCTCTAGGCCCTCATTT

Key term: *coverage*. Usually it's short for *average* coverage: the average number of reads covering a position in the genome

CTAGGCCCTCAATTTTT
CTCTAGGCCCTCAATTTTT
GGCTCTAGGCCCTCATTTTTT
CTCGGCTCTAGCCCCTCATTTTT
TATCTCGACTCTAGGCCCTCA
TATCTCGACTCTAGGCC
TCTATATCTCGGCTCTAGG
GGCGTCTATATCTCG
GGCGTCGATATCT

177 nucleotides

35 nucleotides

GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTTT

GGCGTCTATATCT

Coverage could also refer to the number of reads covering a particular position in the genome:

```
CTAGGCCCTCAATTTTT
                CITCTAGGCCCTCAATTTTT
              GGCTCTAGGCCCTCATTTTT
           CTCGGCTCTAGCCCCTCATTTT
        TATCTCGACTCTAGGCCCTCA
      TATCTCGACTCTAGGCC
    TCTATATCTCGGGTCTAGG
GGCGTCTATATCTCG
GGCGTCGATATCT
GGCGTCTATATCT
GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTTT
```

Coverage at this position = 6

- Best case scenario: check every pair of (long) reads for overlaps
- Computationally expensive: n reads, ~n**2 operations
- Our task: how to get the best assemblies at the smallest expense in terms of sequencing and computational expenses

Read1 - TTTGGTGCTCTTCGAAAAGGGATCTTCGAGAGAGATCTCGCGATAAGGTTG

overlap

Basic principle: the more similarity there is between the end of one read and the beginning of another...



...the more likely they are to have originated from overlapping stretches of the genome:

TATCTCGACTCTAGGCC TCTATATCTCGGCTCTAGG GGCGTCTATATCTCGGCTCTAGGCCCTCATTTTTT

Say two reads truly originate from overlapping stretches of the genome. Why might there be

differences?

TATCTCGACTCTAGGCC
||||||| ||||||
TCTATATCTCGGCTCTAGG

1. Sequencing error

2. Difference between inherited *copies* of a chromosome. E.g. humans are diploid; we have two copies of each chromosome, one from mother, one from father. The copies can differ:

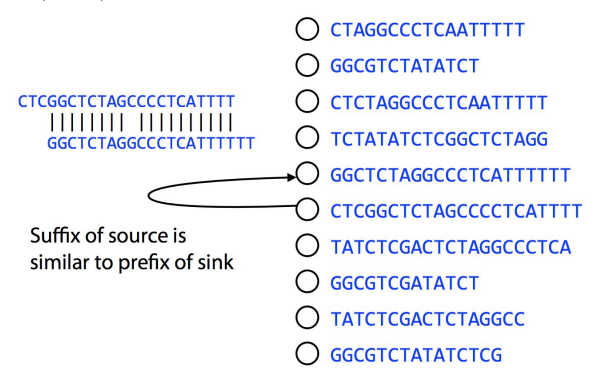
Read from Mother: TATCTCGACTCTAGGCC

Read from Father: TCTATATCTCGGCTCTAGG

Sequence from Mother: TCTATATCTCGACTCTAGGCC Sequence from Father: TCTATATCTCGGCTCTAGGCC

We'll mostly ignore ploidy, but real tools must consider it

Finding all overlaps is like building a *directed graph* where directed edges connect overlapping nodes (reads)



Graphs

Directed graph G=(V, E) is a pair consisting of *node set* (or vertex set) V and *edge set* (or *arc set*) $E \subseteq VxV$.

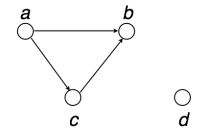
An **edge** $e = (u, v) \in E$ represents a connection from u to v. **Directed edge** is an ordered pair of vertices.

We call *u* and *v* the *source* and the *target* (*sink*), respectively, of *e*.

Directed graph also called digraph.

An *outdegree* of the node is the number of edges leaving it, and *indegree* of the node is the number of edges ending at it.

Graph theory developed by Euler (more info about this later...)



$$V = \{ a, b, c, d \}$$

 $E = \{ (a, b), (a, c), (c, b) \}$
Source Sink

Overlap graph

Below: overlap graph, where an overlap is a suffix/prefix match of at least 3 characters.

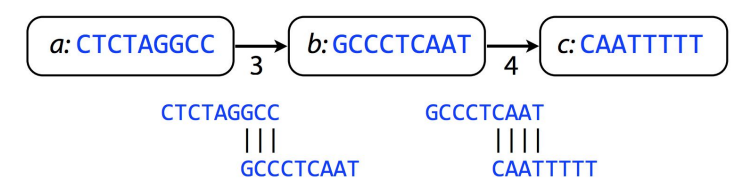
In overlap graph:

A vertex is a read.

A directed edge is an overlap between suffix of source and prefix of sink.

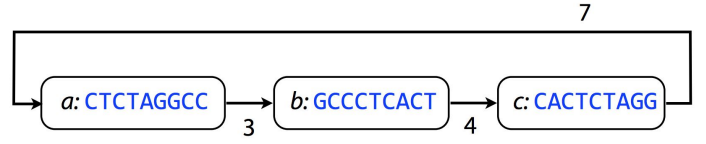
Vertices (reads): { a: CTCTAGGCC, b: GCCCTCAAT, c: CAATTTTT }

Edges (overlaps): $\{(a, b), (b, c)\}$



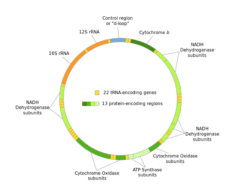
Overlap graph - cycles in graph

Overlap graph could contain *cycles*. A cycle is a path beginning and ending at the same vertex.



These happen when the DNA string itself is circular. E.g. bacterial genomes are often circular; mitochondrial DNA is circular.

Cycles could also be due to *repetitive* DNA, as we'll see.



Finding overlaps



How do we build the overlap graph?

What constitutes an overlap?

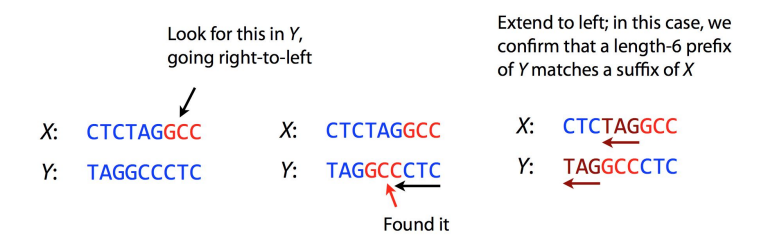
(**Def:**?) Assume for now an "overlap" is when a suffix of **X** of **length** ≥ **I** exactly matches a prefix of **Y**, where *I* is given.

Finding overlaps

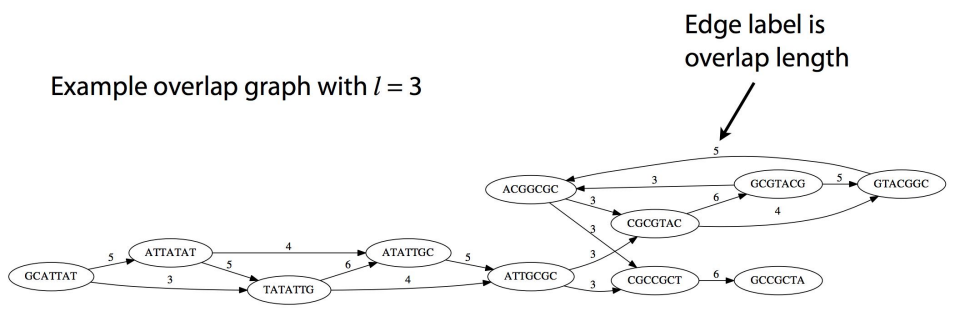
Overlap: length-/ suffix of X matches length-/ prefix of Y, where / is given.

Simple idea: look in Y for occurrences of length-/ suffix of X. Extend matches to the left to confirm whether entire prefix of Y matches.

For I=3



Finding overlaps



Finding overlaps is important, and we'll return to it, but our ultimate goal is to recreate (assemble) the genome.

Every sequence (read) is a part of the genome, and we're trying to recreate a genome from reads. We need a *superstring* of the reads - string (genome) that contains all reads we have. Reasonable approximation is that we want not any, but *shortest superstring*.

How do we formulate this problem?

First attempt: the shortest common superstring (SCS) problem:

Given a set of strings, find a shortest string that contains all of them.

Given a collection of strings S, find SCS(S): the shortest string that contains all strings in S as substrings.

Without requirement of "shortest," it's easy: just concatenate them

```
Example: S: BAA AAB BBA ABA ABB BBB AAA BAB
```

```
Concatenation: BAAAABBBAABBBBBAAABAB

SCS(S): AAABBBBABAA

AAA

AAB

ABB

ABB

BBB

BBA

BAB

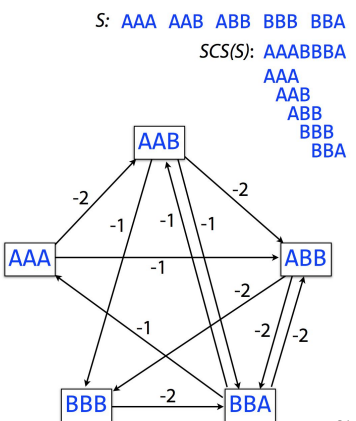
ABA

ABA
```

How can we solve it?

Imagine a modified overlap graph where each edge has **cost** = - (length of overlap)

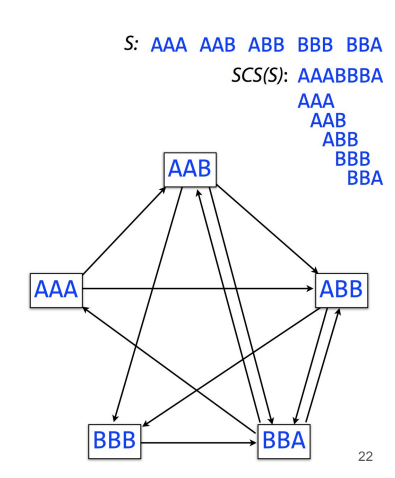
SCS corresponds to a path that visits every node once, minimizing total cost along path That's the Traveling Salesman Problem (TSP), which is NP-hard!



Say we disregard edge weights and just look for a path that visits all the nodes exactly once

That's the Hamiltonian Path problem: NP-complete

Indeed, it's well established that SCS is NP-hard



NP problems

NP: Decision problems (yes or no answers), if "yes" can be checked in polynomial time

NP complete: Solve one, solved them all!

- Traveling salesman problem
- Knapsack problem
- Integer linear programming
- SAT problem ...

(Karp's 21 NP-complete problems)

NP hard: at least hard as NP

Hamiltonian Path

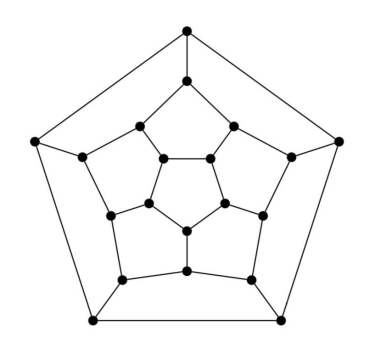
Hamiltonian Cycle Problem:

Find a cycle in a graph that visits every vertex exactly once.

Input: A graph G.

Output: A cycle in G that visits every vertex exactly once (if exists).

- Sir Richard Hamilton created a game:
 - Graph whose vertices are labeled with the name of 25 cities
 - Visit every city only once
 - Sold via London game dealer for 25 pounds
 - Failed miserably, but defined "Hamiltonian Path" problem which occupied mathematicians and engineers in the times to come
 - NP-complete



Sir Richard Hamilton's Icosian Game

Hamiltonian Path



Sir Richard Hamilton's Icosian Game

NP, NP-hard, NP-complete

Shortest common superstring & friends: For refreshers on Traveling Salesman, Hamiltonian Path, NP-hardness and NP-completeness, see Chapters 34 and 35 of "Introduction to Algorithms" by Cormen, Leiserson, Rivest and Stein;

or Chapters 8 and 9 of "Algorithms" by Dasgupta, Papadimitriou and Vazirani (free online: http://www.cs.berkeley.edu/~vazirani/algorithms)

TI;dr - stack overflow - What are the differences between NP, NP-Complete and NP-Hard?

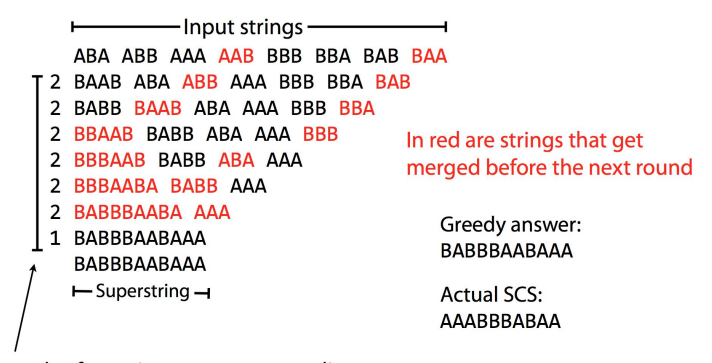
Greedy approach

Let's take the hint, give up on finding the shortest possible superstring.

Non-optimal superstrings can be found with a *greedy* algorithm.

At each step, the greedy algorithm "greedily" chooses longest remaining overlap, merges its source and sink.

Greedy-SCS algorithm in action (l = 1):



Rounds of merging, one merge per line. Number in first column = length of overlap merged before that round.

Greedy algorithm is not guaranteed to choose overlaps yielding SCS.

But greedy algorithm is a good approximation; i.e. the superstring yielded by the greedy algorithm won't be more than ~2.5 times longer than true SCS (see *Gusfield 16.17.1*)

Greedy-SCS algorithm in action again (I = 3):

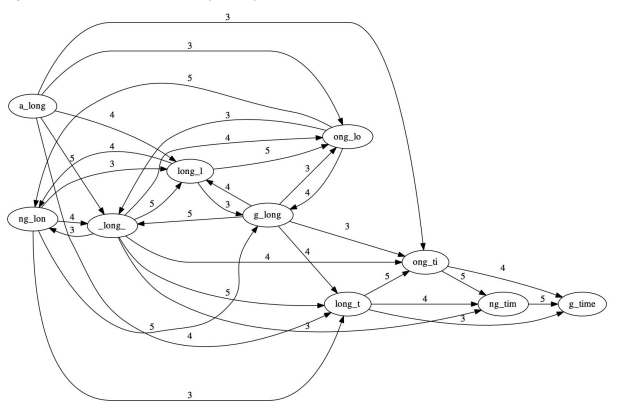
————Input strings — ATTATAT CGCGTAC ATTGCGC GCATTAT ACGGCGC TATATTG GTACGGC GCGTACG ATATTGC 6 TATATTGC ATTATAT CGCGTAC ATTGCGC GCATTAT ACGGCGC GTACGGC GCGTACG CGCGTACG TATATTGC ATTATAT ATTGCGC GCATTAT ACGGCGC GTACGGC CGCGTACG TATATTGCGC ATTATAT GCATTAT ACGGCGC GTACGGC 5 CGCGTACGGC TATATTGCGC ATTATAT GCATTAT ACGGCGC 5 CGCGTACGGCGC TATATTGCGC ATTATAT GCATTAT 5 CGCGTACGGCGC GCATTATAT TATATTGCGC 5 CGCGTACGGCGC GCATTATATTGCGC 3 GCATTATATTGCGCGTACGGCGC GCATTATATTGCGCGTACGGCGC ——Superstring ——

What happened?

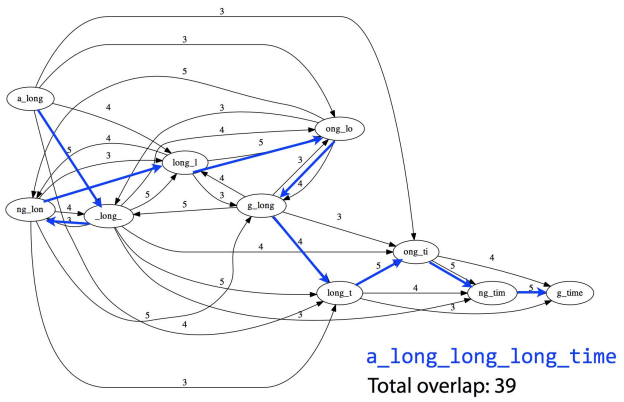
Another setup for Greedy-SCS: assemble all substrings of length 6 from string a_long_long_time. I = 3.

```
ng lon long a long long long ti ong lo long t g long g time ng tim
 5 ng time ng lon long a long long long ti ong lo long t g long
 5 ng time g long ng lon a long long long ti ong lo long t
 5 ng_time long_ti g_long_ ng_lon a_long long_l ong_lo
 5 ng time ong lon long ti g long a long long l
 5 ong lon long time g long a long long l
 5 long lon long time g long a long
 5 long long long time a long
 5 long long time a long
 4 a long long time
   a long long time
I only got back: a long long time (missing one long)
```

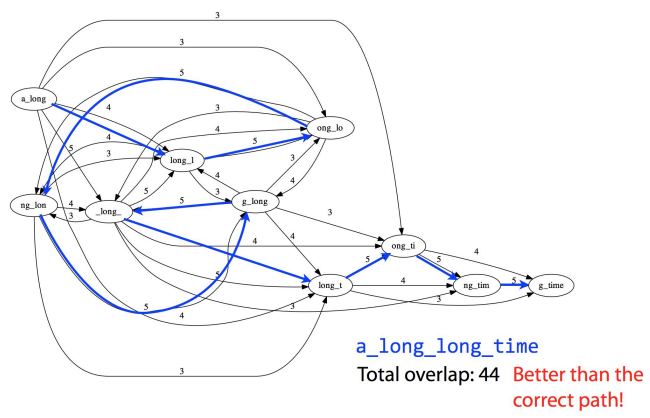
The overlap graph for that scenario (I = 3):



The overlap graph for that scenario (I = 3):



The overlap graph for greedy solution scenario (I = 3):



Same example, but increased the substring length from 6 to 8

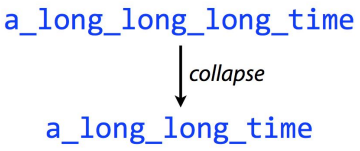
Got the whole thing: a_long_long_time

Why are substrings of length 8 long enough for Greedy-SCS to figure out there are 3 copies of long?

One length-8 substring spans all three longs

Repeats often foil assembly. They certainly foil SCS, with its "shortest" criterion! Reads might be too short to "resolve" repetitive sequences.

This is why sequencing vendors try to increase read length. Algorithms that don't pay attention to repeats (like our greedy SCS algorithm) might collapse them



The human genome is ~ 50% repetitive!

Basic principle: repeats foil assembly!

Another example using Greedy-SCS:

```
Input: it_was_the_best_of_times_it_was_the_worst_of_times
```

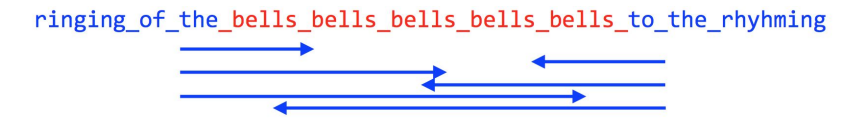
Extract every substring of length k, then run Greedy-SCS. Do this for various *I* (min overlap length) and k.

l, k	output
3, 5	the_worst_of_times_it_was_the_best_o
3, 7	s_the_worst_of_times_it_was_the_best_of_t
3, 10	_was_the_best_of_times_it_was_the_worst_of_tim
3, 13	<pre>it_was_the_best_of_times_it_was_the_worst_of_times</pre>

Basic principle: repeats foil assembly

Longer and longer substrings allow us to "anchor" more of the repeat to its non-repetitive context:

Often we can "walk in" from both sides. When we meet in the middle, the repeat is resolved:



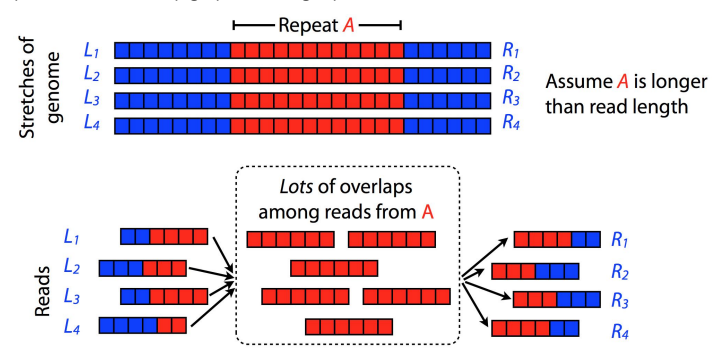
Basic principle: repeats foil assembly

Yet another example using Greedy-SCS:

```
Input: swinging_and_the_ringing_of_the_bells_bells_bells_bells
```

longer and longer substrings allow us to "reach" further into the repeat

Picture the portion of the overlap graph involving repeat A



Even if we avoid collapsing copies of A, we can't know which paths in correspond to which paths out.

SCS is too computationally complex as a way of formulating the assembly problem

- There is no practical way to find SCS
- Greedy approach: we might get too long answers (at least ~2.5 times)
- Repeat handling: SCS might collapse repeat sequences
 - In this case answers might get too short

Need solutions that are: a) tractable b) handle repeats as best as possible

Note: Repeats are problem in any assembly algorithm. This is a property of read length and repetitiveness of the genome.

Taxonomy of assembly approaches

Search for most parsimonious explanation of the reads (shortest superstring):

- Exact solutions are intractable (e.g. TSP), but a greedy approximation is possible
- Any solution will collapse repeats spuriously

Search for "maximum likelihood" explanation of the reads; i.e. force solution to be consistent with uniform coverage:

No solutions (that I know of) are tractable

Give up on unresolvable repeats and use a tractable algorithm to assemble the resolvable portions. This is what real tools do.