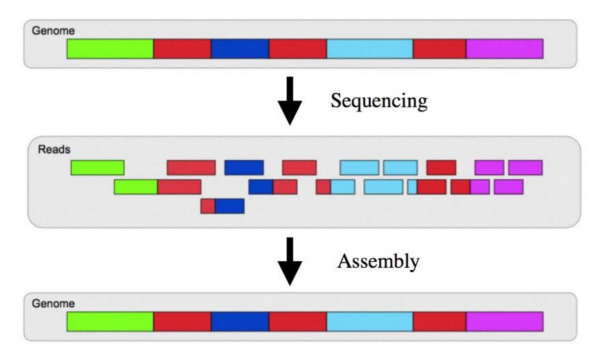
Genome assembly by short/long-read sequencing

2019 Dragon Star Bioinformatics Course (Day 2)

What is genome assembly?

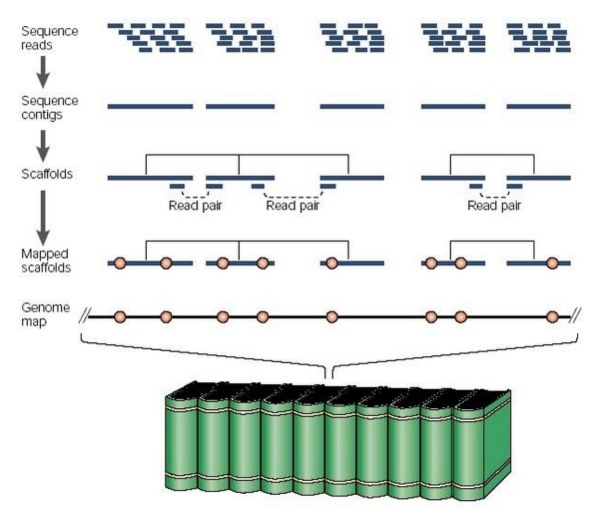
 The genome assembly is simply the genome sequence produced after chromosomes have been fragmented, those fragments have been sequenced, and the resulting sequences have been put back together.



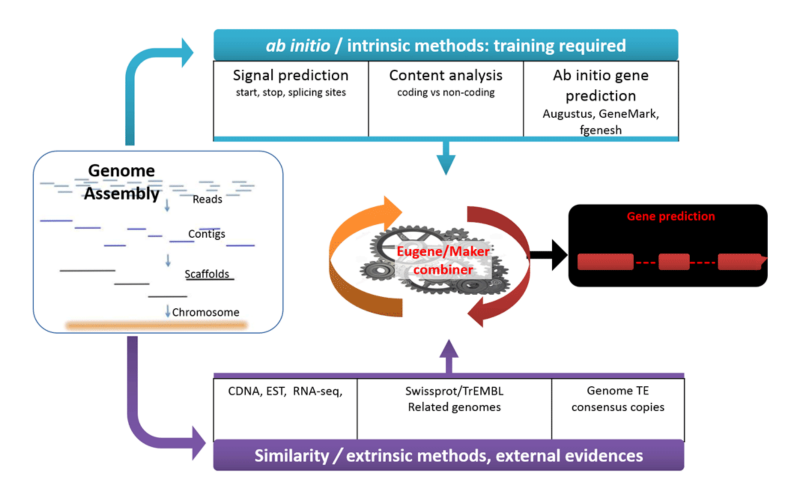
Why genome assembly?

- Accurate assembly of genomes is key to understanding genetic variation. The more accurate the reference genome, the easier it is to map reads and interpret the functional impacts of genetic mutations
- Some reference-free variant caller are available, but most of the human genetic data analysis relies on reference genome. Therefore, accurate genome assembly and more complete (that incorporate population variation and alternative haplotypes) is important for human genetic studies.

Long-range sequence assembly in wholegenome shotgun sequencing



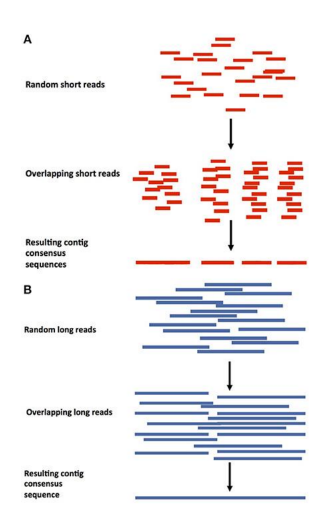
Why genome assembly?

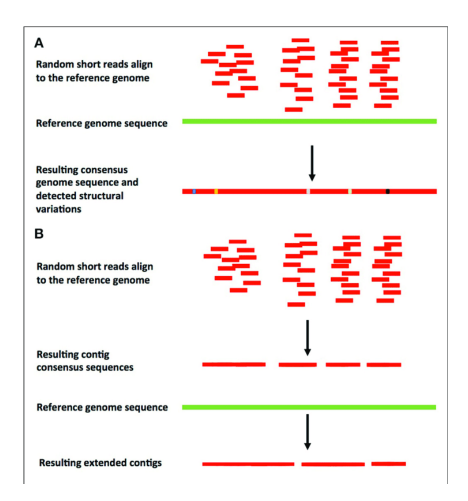


What types of genome assembly?

- de novo assembly: the entire assembled sequence is resolved from raw sequence data without comparison to a reference genome sequence.
- Comparative assembly: you have a "reference" genome to guide the assembly process. For example, sequence and assemble a new human genome, using GRCh38 as a guide. Or sequence a chimp genome using the existing human assembly as a guide.

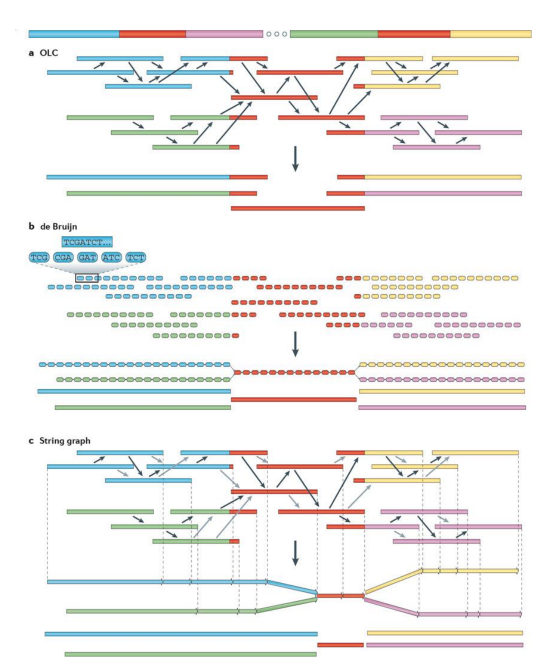
De novo vs reference-guided assembly





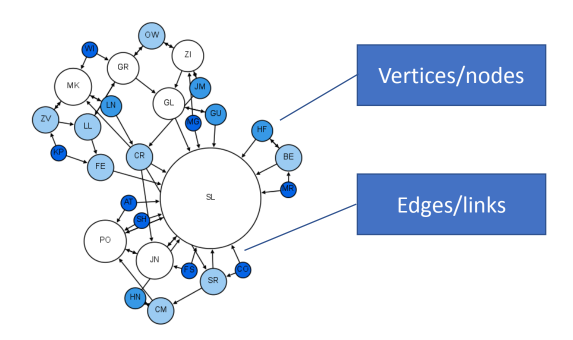
De Novo Assembly paradigms

- Overlap-layout-consensus (OLC) methods
 - greedy (TIGR Assembler, phrap, CAP3...)
 - graph-based (Celera Assembler, Arachne)
 - Examples are Arachne, Celera Assembler, CAP3, PCAP, Phrap, Phusion and Newbler
- De bruijn (k-mer) graph
 - Especially useful for assembly from short reads
 - Stacking overlapping sequences of genomic fragments of a defined size (the k-mer), generated by breaking each read into k-mer size.
 - Examples are Euler-USR, Velvet, ABySS, AllPath-LG and SOAPdenovo.
- Extensions and other approaches
 - String graph
 - Hybrid approach



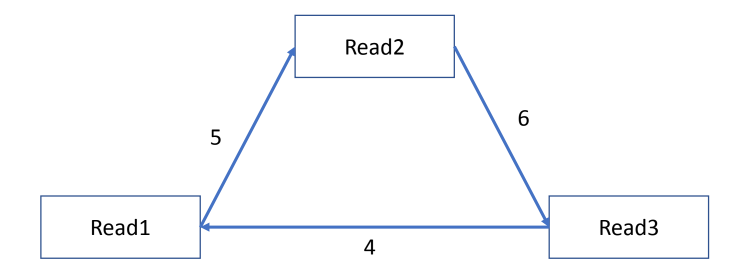
Graph theory

• Graph: mathematical structures used to model pairwise relations between objects. A graph is made up of vertices (also called nodes or points) which are connected by edges (also called links or lines).



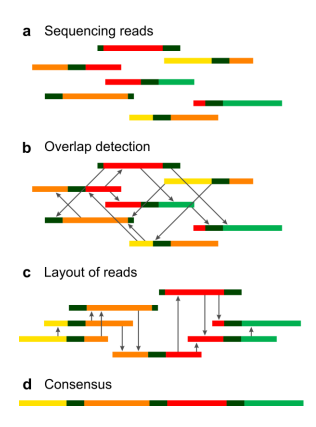
Directed graph

- G(V, E): a set of vertices (V) and a set of directed edges (E)
- Vertex and edge may have weights associated with them
- Here, V= {Read1, Read2, Read3}, E = {(Read1, Read2), (Read2, Read3), (Read3, Read3)}



Overlap-based approach

 Overlap graph: the nodes are initially formed by the sequences of the individual reads while the edges are represented by the sequence overlaps between these reads.



Three steps: overlap-layout-consensus

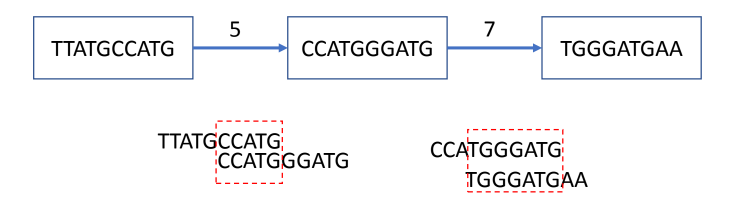
- Overlap: The identification of overlaps between all sequencing reads is the first and most time consuming step of the assembly process.
 Pairwise overlaps of variable length are identified between all reads in the dataset and represented as edges between these reads in the resulting overlap graph
- Layout: This information is used in the next step to layout the reads into the most probable contiguous sequence stretches
- Consensus: In the final step the consensus sequence is determined for each contig by choosing the nucleotide, which is represented by the majority of the overlapping reads for every sequence position

Constructing the overlap graph

- Create the overlap graph for the following string, where overlap means suffix/prefix match of >=5bp
 - TTATGCCATGGGATGAA
- Note that TT, CC, GG and AA are separated by 3 identical ATG
- Assume that we generated a few short reads from this string: TTATGCCATG, CCATGGGATG, TGGGATGAA
- Vertices/nodes are formed by the sequences of the individual reads; edges are represented by the sequence overlaps between these reads

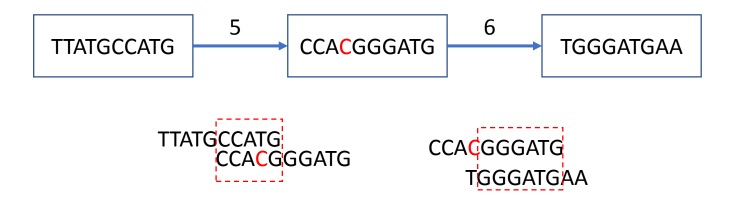
Constructing the overlap graph

- Genome: TTATGCCATGGGATGAA
- Vertices: {TTATGCCATG, CCATGGGATG, TGGGATGAA}
- Edges: {(TTATGCCATG, CCATGGGATG), (CCATGGGATG, TGGGATGAA)}



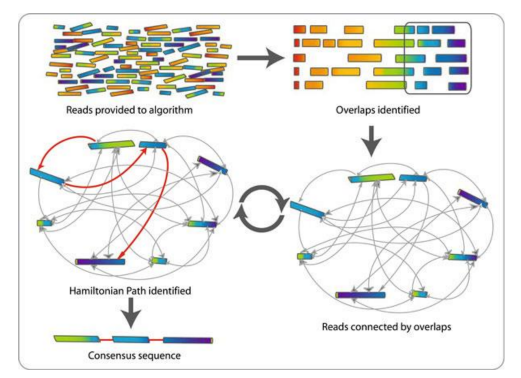
Sometimes we need to tolerate sequencing errors

- Genome: TTATGCCATGGGATGAA
- Vertices: {TTATGCCATG, CCATGGGATG, TGGGATGAA}
- Edges: {(TTATGCCATG, CCATGGGATG), (CCATGGGATG, TGGGATGAA)}



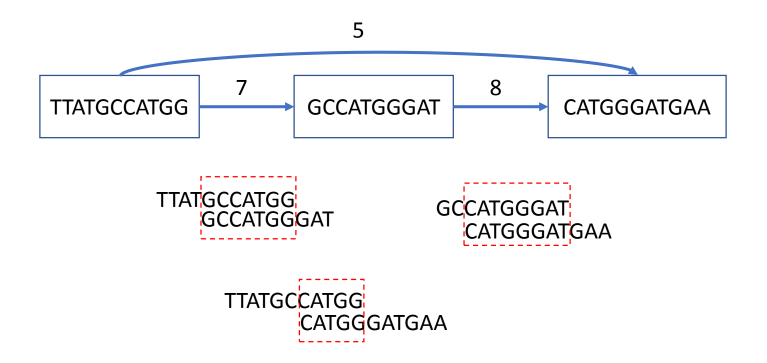
Layout step

• The algorithm determines the best path through the graph (Hamiltonian path). Redundant information (i.e., unused nodes and edges) is discarded.

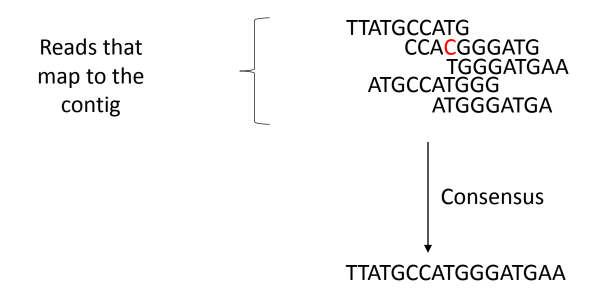


Sometimes graphs can be reduced by eliminating transitive edges

Genome: TTATGCCATGGGATGAA



Consensus step: inferring the most likely nucleotide at each position

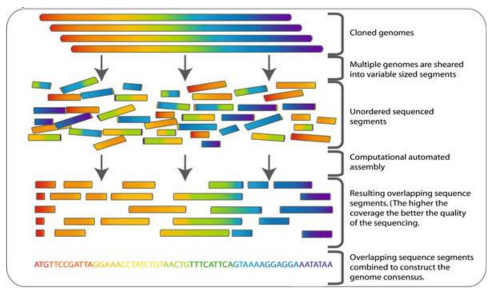


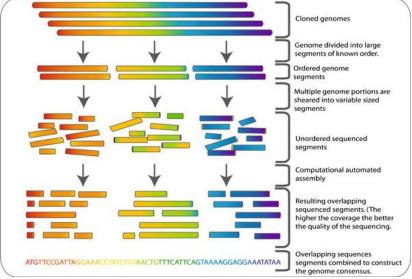
Note: sequencing error or mapping error (map to repetitive regions) or ploidy can result in mismatches

Note: For long-read sequencing, it is common to do error correction first to reduce error rates of reads, and then use a fraction of the longest reads for overlap-based assembly

Comparison of two approaches for large genomes such as human genomes

Whole genome shotgun sequencing vs hierarchical approach





Now let's go back 30 years

- How should we sequence and assemble a human genome?
- Begun formally in 1990, the U.S. Human Genome Project was a 13year effort coordinated by the U.S. Department of Energy (DOE) and the National Institutes of Health (NIH; http://www.genome.gov/).
- Primary goals were to discover the complete set of human genes and make them accessible for further biological study, and determine the complete sequence of DNA bases in the human genome.
- Two thoughts: whole-genome shotgun versus physical mapping followed by BAC-by-BAC sequencing

Primary Human Genome Project Sequencing Sites

- U.S.DOE Joint Genome Institute, Walnut Creek, California, USA*
- Baylor College of Medicine Human Genome Sequencing Center, Department of Molecular and Human Genetics, Houston, Texas, USA*
- The Wellcome Trust Sanger Institute, The Wellcome Trust Genome Campus, Hinxton, Cambridgeshire, United Kingdom*
- Washington University School of Medicine Genome Sequencing Center, St. Louis, Missouri, USA*
- Whitehead Institute/MIT Center for Genome Research, Cambridge, Massachusetts, USA*

Additional Contributors to HGP Research

- · Australian Genome Research Facility
- Beijing Genomics Institute/Human Genome Center, Institute of Genetics, Chinese Academy of Sciences, Beijing, China*
- Chromosome 21 Consortium
- Cold Spring Harbor Laboratory, Lita Annenberg Hazen Genome Center, Cold Spring Harbor, New York, USA*
- Chromosome 13 Sanger Centre, UK
- · Computational Biology at Oak Ridge National Laboratory, Oak Ridge, TN, USA
- · European Bioinformatics Institute
- · Foundation Jean Dausset-CEPH
- Généthon
- Genome Database
- Genoscope and CNRS UMG-8030, Evry, France*
- · German Human Genome Project
- · Gesellschaft fur Biotechnologische Forschung mbH, Braunschweig, Germany*
- GTC Sequencing Center, Genome Therapeutics Corporation, Waltham, Massachusetts, USA*
- Department of Genome Analysis, Institute of Molecular Biotechnology, Jena, Germany*
- The Institute for Systems Biology, Seattle, Washington, USA*
- · Japan Science and Technology Corporation Sequencing Projects
- Department of Molecular Biology, Keio University School of Medicine, Tokyo, Japan*
- Max Planck Institute for Molecular Genetics, Berlin, Germany*

How should we assemble a human genome?

 Two thoughts: whole-genome shotgun versus physical mapping followed by BAC-by-BAC sequencing

PERSPECTIVE

Human Whole-Genome Shotgun Sequencing

James L. Weber^{1,3} and Eugene W. Myers²

¹Center for Medical Genetics, Marshfield Medical Research Foundation, Marshfield, Wisconsin 54449; ²Department of Computer Science, University of Arizona, Tucson, Arizona 85721

Large-scale sequencing of the human genome is now under way (Boguski et al. 1996; Marshall and Pennisi 1996). Although at the beginning of the Genome Project, many doubted the scientific value of sequencing the entire human genome, these doubts have evaporated almost entirely (Gibbs 1995; Olson 1995). Primary reasons for generating the human genomic sequence are listed in Table 1.

Table 5. Costs of Human Genomic Sequencing

Clone by clone
\$0.30 per finished base
\$130 million per year for 7 years
Total \$900 million spent by end of 2003
Shotgun
\$0.01 per raw base
\$130 million for 3 years would provide
10× coverage plus an additional \$90 million
for informatics

 Two thoughts: whole-genome shotgun versus physical mapping followed by BAC-by-BAC sequencing

PERSPECTIVE

Against a Whole-Genome Shotgun

Philip Green¹

Department of Molecular Biotechnology, University of Washington, Seattle, Washington 98195

The human genome project is entering its decisive final phase, in which the genome sequence will be determined in large-scale efforts in multiple laboratories worldwide. A number of sequencing groups are in the process of scaling up their throughput; over the next few years they will need to attain a collective capacity approaching half a gigabase per year to complete the 3-Gb genome sequence by the target date of 2005. At present, all contributing groups are using a clone-by-clone approach, in

MIT Center for Genome Research, http://www-genome.wi.mit.edu], with several intensively mapped chromosomes already exceeding it (Nagaraja et al. 1997, Bouffard et al. 1997), and BACs average 130 kb or more in size in current libraries (Kim et al. 1996), this STS density should be adequate to obtain contiguous clone coverage of much of the genome; most gaps that remain should be closable by developing new STSs directly from the sequence adjacent to the gap and rescreening the library.

At present, all contributing groups are using a clone-by-clone approach, in which mapped bacterial clones (typically 40–400 kb in size) from known chromosomal locations are sequenced to completion. Among other advantages, this permits a variety of alternative sequencing strategies and methods to be explored independently without redundancy of effort.

Draft human genome

- Public effort: BAC based sequencing (Lander et al., Nature, Feb. 15, 2001). Performed by a consortium of government labs and universities.
 - Break genome into ~100-300kb pieces and create Bacterial Artificial Chromosomes (BACs) from each piece, assemble each piece by Sanger sequencing, then stitch overlapping BACs together to draft genome.

- Private effort: whole genome random shotgun sequencing (Venter et al., Science, Feb. 16, 2001). Performed by Celera, which is a company.
 - In 1998, Craig Venter announced that he was forming Celera that within three years would sequence human genome —seven years before the projected finish of the U.S. government's Human Genome Project.

On the sequencing of the human genome

Robert H. Waterston*†, Eric S. Lander‡, and John E. Sulston§

*Genome Sequencing Center, Washington University, Saint Louis, MO 63108; [‡]Whitehead Institute/Massachusetts Institut Genome Research, Cambridge, MA 02142; and [§]Wellcome Trust Sanger Institute, Wellcome Trust Genome Campus, Hinxto United Kingdom

Communicated by Aaron Klug, Medical Research Council, Cambridge, United Kingdom, December 21, 2001 (received for r

Two recent papers using different approaches reported draft sequences of the human genome. The international Human Genome Project (HGP) used the hierarchical shotgun approach, whereas Celera Genomics adopted the whole-genome shotgun (WGS) approach. Here, we analyze whether the latter paper provides a meaningful test of the WGS approach on a mammalian genome. In the Celera paper, the authors did not analyze their own WGS data. Instead, they decomposed the HGP's assembled sequence into a "perfect tiling path", combined it with their WGS data, and assembled the merged data set. To study the implications of this approach, we perform computational analysis and find that a perfect tiling path with 2-fold coverage is sufficient to recover virtually the entirety of a genome assembly. We also examine the manner in which the assembly was anchored to the human genome and conclude that the process primarily depended on the HGP's sequence-tagged site maps, BAC maps, and clone-based

sequences. Our analysis indicates that the Celera paper provides neither a meaningful test of the WGS approach nor an independent sequence of the human genome. Our analysis does not imply that a WGS approach could not be successfully applied to assemble a draft sequence of a large mammalian genome, but merely that the

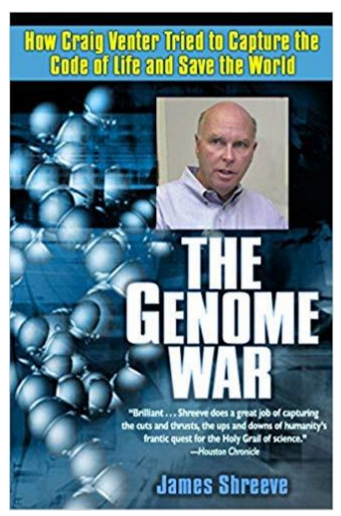
problem is straightforward, becausequence can typically be mergemisassembly. The relatively few gap to produce complete sequences. The successfully to produce complete such as plasmids, viruses, organ genome shotgun data alone also had 15-fold redundancy (5) to produce uchromatic portion of the *Droceontent*), although a clone-based convert this to a finished sequence

A greater challenge arises in tag a large proportion of repeat seq misassembly. Two alternative appr

Hierarchical shotgun (HS) assemble nome is first broken up into a intermediate clones such as bact (BACs). The sequence of each Basequencing, and the sequence of merging the sequences of the BAC a guaranteed route for producing

What happened from Craig's perspective

- Calling his company Celera (from the Latin for "speed"), he assembled a small group of scientists in an empty building in Rockville, Maryland, and set to work.
- At the same time, the leaders of the government program began to mobilize an unexpectedly unified effort to beat Venter to the prize—knowledge that had the potential to revolutionize medicine and society.
- It is also the story of how one man's ambition created a scientific Camelot where, for a moment, it seemed that the competing interests of pure science and commercial profit might be gloriously reconciled—and the national repercussions that resulted when that dream went awry.



How Perl Saved the Human Genome Project

How Perl Saved the Human Genome Project

by Lincoln Stein

Reprinted courtesy of the Perl Journal, http://www.tpj.com

Lincoln Stein's website is http://stein.cshl.org

DATE: Early February, 1996

LOCATION: Cambridge, England, in the conference room of the largest DNA sequencing center in Europe.

OCCASION: A high level meeting between the computer scientists of this center and the largest DNA sequencing center in the United States.

THE PROBLEM: Although the two centers use almost identical laboratory techniques, almost identical databases, and almost identical data analysis tools, they still can't interchange data or meaningfully compare results.

THE SOLUTION: Perl.

The human genome project was inaugurated at the beginning of the decade as an ambitious international effort to determine the complete DNA sequence of human beings and several experimental animals. The justification for this undertaking is both scientific and medical. By understanding the genetic makeup of an organism in excruciating detail, it is hoped that we will be better able to understand how organisms develop from single eggs into complex multicellular beings, how food is metabolized and transformed into the constituents of the body, how the nervous system assembles itself into a smoothly functioning ensemble. From the medical point of view, the wealth of knowledge that will come from knowing the complete DNA sequence will greatly accelerate the process of finding the causes of (and potential cures for) human diseases.

Six years after its birth, the genome project is ahead of schedule. Detailed maps of the human and all the experimental animals have been completed (mapping out the DNA using a series of landmarks is an obligatory first step before determining the complete DNA sequence). The sequence of the smallest model organism, yeast, is nearly completed, and the sequence of the next smallest, a tiny soil-dwelling worm, isn't far behind. Large scale sequencing efforts for human DNA started at several centers a number of months ago and will be in full swing within the year.

The scale of the human DNA sequencing project is enough to send your average Unix system administrator running for cover. From the information-handling point of view, DNA is a very long string consisting of the four letters G, A, T and C (the letters are abbreviations for the four chemical units that form the "rungs" of the DNA double helix ladder). The goal of the project is to determine the order of letters in the string. The size of the string is impressive but not particularly mind-boggling: 3 x 10⁴ letters long, or some 3 gigabytes of storage space if you use 1 byte to store each letter and don't use any compression techniques.

Random web quotes: "Perl and the human genome are almost perfectly matched; both are almost incomprehensible, with no central design, accreted haphazardly over a long time."

Lander-Waterman statistics in genome assembly

- Originally developed to guide the planning of a genome assembly project (how to select random clones and declare overlap of clones)
- Provide guidelines on expected number of contigs

GENOMICS 2, 231-239 (1988)

Genomic Mapping by Fingerprinting Random Clones: A Mathematical Analysis

ERIC S. LANDER*, † AND MICHAEL S. WATERMAN‡

*Whitehead Institute for Biomedical Research, 9 Cambridge Center, Cambridge, Massachusetts 02142; †Harvard University, Cambridge, Massachusetts 02138; and ‡Departments of Mathematics and Molecular Biology, University of Southern California, Los Angeles, California 90089

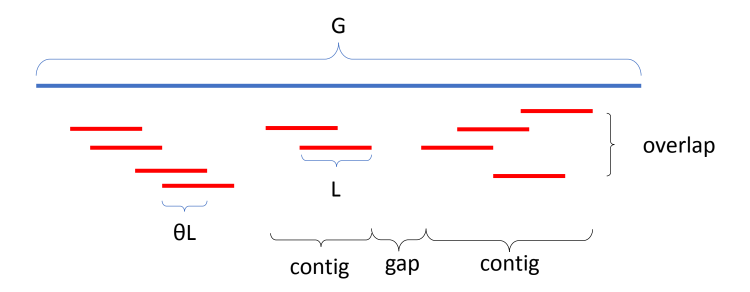
Received January 13, 1988; revised March 31, 1988

Back to today

- Gigabase genomes can be easily sequenced and assembled using PacBio/Nanopore long-read assemblers, often with a N50 value over 10Mb, which are orders of magnitude higher than conventional approaches that generate sub-kb reads
- Whole-genome shotgun is the predominant approach to be used today for assembling these large genomes, but optical mapping or physical map of specific genetic markers can still provide clue in placing large contigs/scaffolds in genomic regions.

Lander-Waterman statistics

- Let N = # of reads, L = length of a read (average length), G = genome length, c = coverage = N*L/G
- Main questions to address:
 - What's the fraction of genome that are covered by reads?
 - How many contigs are generated?



Coverage at a position can be modelled by Poisson distribution

- c = coverage = N*L/G
- Use Poisson distribution

$$P(k,\lambda) = e^{-\lambda} \times \frac{\lambda^k}{k!}$$

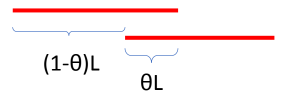
- Probability that a base is NOT covered = $P(0,c) = e^{-c}$
- For genome size G, # of uncovered bases = G* P(0,c) = G * e^{-c}

What's the fraction of genome that are covered by reads?

- This is the same question as "how many positions have coverage>0"?
- For genome size G, number of positions with read coverage: G * (1-P(0,c)) = G * (1- e^{-c}).

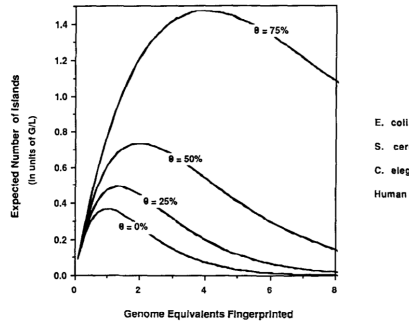
How many contigs are generated?

- How many contigs are generated?
- The same question: how many gaps are there? Or, how many reads do not have overlap > θ with any other reads (Remember that we declare two reads as overlap only if the fraction of overlap > θ)
- For each read, we calculate the probability that zero reads start at (1- θ)L: $e^{-(1-\theta)c} = e^{-(1-\theta)(NL/G)}$
- For all N reads, number of contigs are: N * $e^{-(1-\theta)c}$



Theory is different from reality!

• If there is no repeat, no polymorphism, no region bias, and no sequence error, sequence assembly could be very easy when >8X coverage can be generated even with $\theta = 25\%$



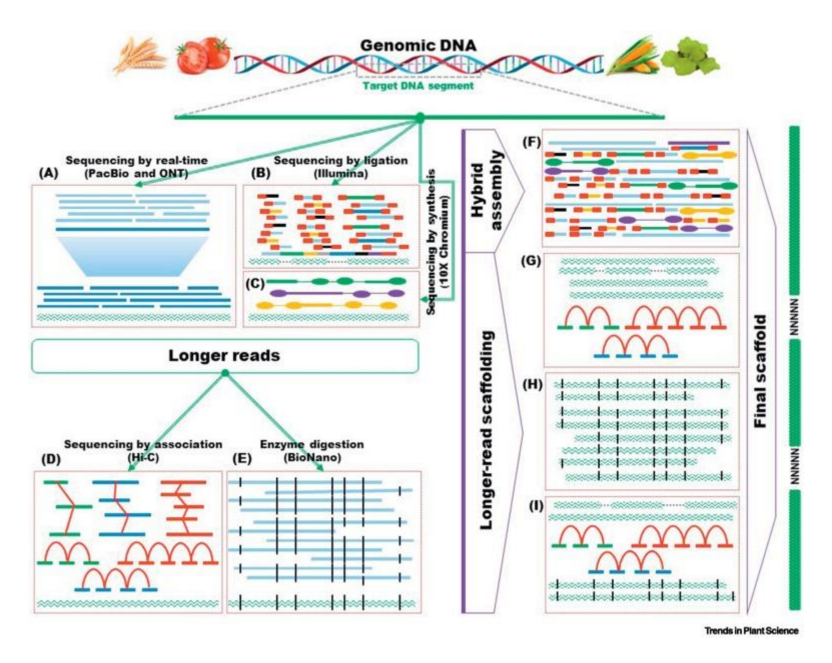
Approximate value of G/L				
Phage (15kb)	Cosmid (40kb)	Yeast (1Mb)		
267	100	4		
1333	500	20		
5,667	2,125	8.5		
200,000	75,000	3,000		
	Phage (15kb) 267 1333 5,667	Phage (15kb) Cosmid (40kb) 267 100 1333 500 5,667 2,125		

Assembling large genomes today

• Since 2013, de novo assembly of large genomes has shifted from short-read sequencing to synthetic or true long-read sequencing.

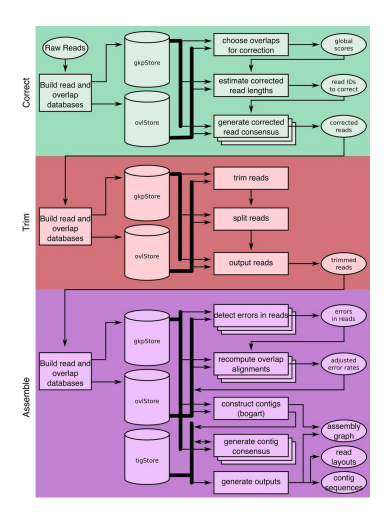
Table 1 Summary of Selected Long-Read Sequencers for De Novo Assemblies of Large Fukanyotic Genomes^{a,t}

Pros and Cons	10X Genomics	Pacific Biosciences	Oxford Nanopore	BioNano	Dovetail ^c
Pros and Cons	Chromium ^c (HiSeq 4000)	(SEQUEL/Cell)	(MinilON)	(Saphyr/Chip)	(HiSeq 4000)
Compatible platforms	Illumina	RS II	GridION ^d and PromethION ^d	Irys	Illumina
Minimum input	~3 ng	~20 µg	~1 µg	~200 ng	~5 µg
Long-read	Synthetic	True	True	True	Synthetic
Average/maximum read length	~300 bp (PE)/ ~150 kb (LLR)	~12 kb/~150 kb	~12 kb/~2 Mb	~350 kb/~1 Mb	~150 kb/~1 Mb (SLR)
Throughput	~1500 Gb	0.7 Gb-20 Gb (SEQUEL)	50 Gb-15 Tb (PromethION)	~640 Gb	~1500 Gb
Reads	~5 Billion (B)	0.07 million (M)–2 M	1.5–5 M	~2 M (image file)	~5 B
Runtime	~3 Days	6–10 h	2 h to 6 days	~1 day	~3 days
Quality scores	>30	>10	>10	NA (only nonsequence based method)	>30
Error profile	<1% (GC/AT biased and substitutions)	5-10% (indels)	5–15% (indels and substitutions)	Sizing error, false sites, and missing sites	<1% (GC/AT-biased and substitutions)
Output format	Fasta Fastq	Bam Fasta Fastq Hdf5 (RS II)	Fast5	BNX C/S/XMAP SVMerge TIFF	Fasta Fastq
General assembly software	Supernova	CANU Falcon/Falcon-Unzip Flye HGAP Minimap/Miniasm	CANU Minimap/Miniasm TULIP	RefAligner	3D-DNA HiRise LACHESIS Meraculous SALSA



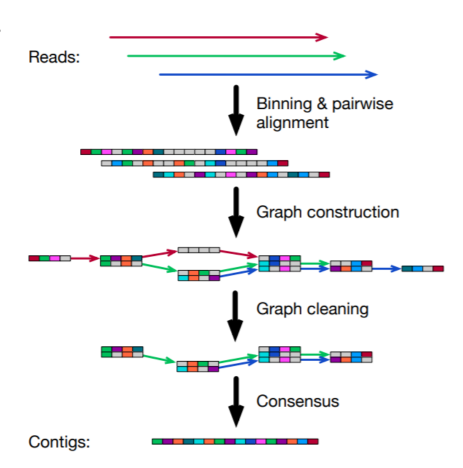
Canu assembler for long-read assembly

- Canu was branched from Celera Assembler in 2015, and specialized for long-read sequencing
- A full Canu run includes three stages: correction (green), trimming (red), and assembly (purple).



Wtdbg2 assembler for long-read assembly

- Wtdbg2 groups 256 base pairs into a bin (bins with the same color suggest they share kmers)
- Wtdbg2 performs all-vs-all alignment between binned reads, ignoring detailed base sequences.
- In the fuzzy-Bruijn assembly graph, a vertex is a 4-bin segment.
- Much faster than other assemblers for long reads

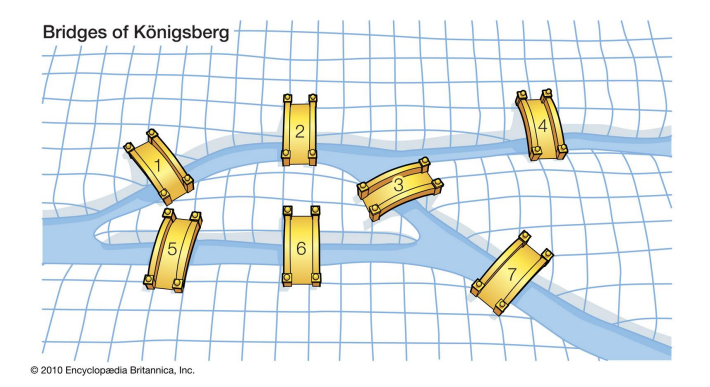


Sparse hierarchical minimizers (SHIMMER)

- The most common approach to long-read assembly, using an overlaplayout-consensus (OLC) paradigm, requires all-to-all read comparisons, which quadratically scales in computational complexity with the number of reads.
- We assert that recently achievements in sequencing technology (i.e. with accuracy ~99% and read length ~10-15k) enables a fundamentally better strategy for OLC that is effectively linear rather than quadratic
- The Peregrine tool index reads, avoiding the need for an all-to-all read comparison step. Peregrine can assemble 30x human PacBio CCS read datasets in less than 30 CPU hours and around 100 wallclock minutes to a high contiguity assembly (N50 > 20Mb).

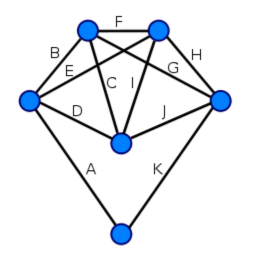
Seven Bridges of Königsberg Problem

 Königsberg bridge problem: whether a citizen could take a walk through the town in such a way that each bridge would be crossed exactly once.



Seven Bridges of Königsberg Problem

- In 1736, the Swiss mathematician Leonhard Euler demonstrated that the answer is no
- In graph theory, an **Eulerian trail** (or **Eulerian path**) is a trail in a finite graph which visits every edge exactly once (allowing for revisiting vertices). Similarly, an Eulerian circuit or Eulerian cycle is an Eulerian trail which starts and ends on the same vertex.



Following the edges in alphabetical order gives an Eulerian circuit/cycle.

Euler's Theorem on directed graphs

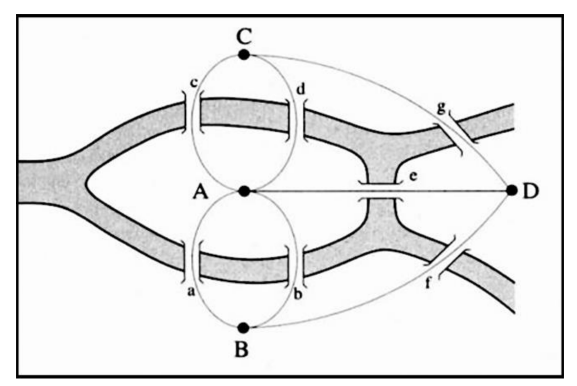
- For directed graphs, the cycle will need to follow the direction of the edges.
 - Indegree(v) = # edges coming into v
 - Outdegree(v) = # edges leaving v
- A directed graph has an Eulerian path if and only if indegree(v)=outdegree(v) for all but 2 nodes (x and y), where indegree(x)=outdegree(x)+1, and indegree(y)=outdegree(y)-1.
- A directed graph has an Eulerian cycle if and only if indegree(v)=outdegree(v) for all nodes

Euler's Theorem on undirected graphs

- For undirected graphs, the degree of a vertex is the number of edges that are incident to the vertex
- An undirected graph has an Eulerian cycle if and only if every vertex has even degree
- An undirected graph has an Eulerian trail if and only if exactly zero or two vertices have odd degree

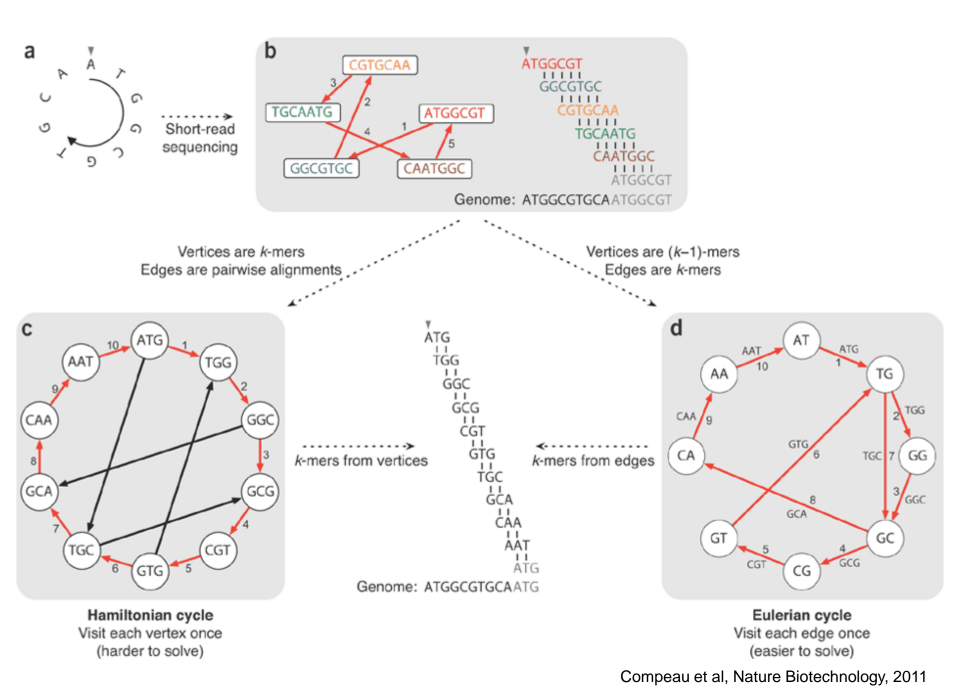
The Königsberg graph is not Eulerian

 For the existence of Eulerian trails it is necessary that zero or two vertices have an odd degree. If there are no vertices of odd degree, all Eulerian trails are circuits. If there are exactly two vertices of odd degree, all Eulerian trails start at one of them and end at the other.



Hamiltonian vs Eulerian path

- Hamiltonian path: a path that travels to every node exactly once and ends at the starting node.
- Eulerian path: a path that visits all edges of a graph exactly once.
- When analyzing sequencing data, instead of assigning each k-mer contained in some read to a node, we will assign each such k-mer to an edge. This allows the construction of a 'de Bruijn graph'.

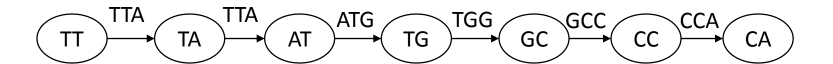


de Bruijn graph

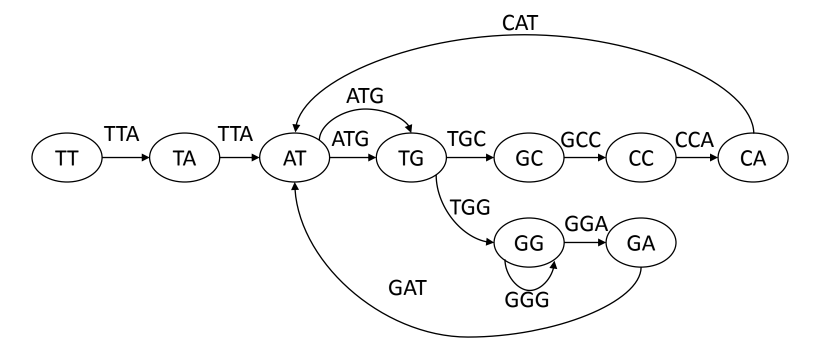
- The reads are k-mers and each corresponds to one edge in the graph
- The vertices are (k-1)-mers that appear in some read, and edges defined by overlap of k-2 nucleotides.
- Small values of k produce small graphs.
- Note this creates a directed graph ("digraph")!

- Create the de Bruijn graph for the following string, using k=3
 - TTATGCCATGGGATGAA
- Note that TT, CC, GG and AA are separated by 3 identical ATG
- vertices are 2-mers
- edges are 3-mers

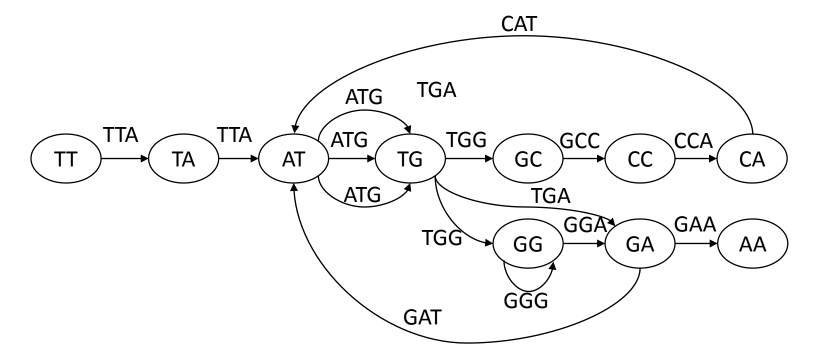
- Create the de Bruijn graph for the following string, using k=3
 - TTATGCCATGGGATGAA



- Create the de Bruijn graph for the following string, using k=3
 - TTATGCCATGGGATGAA

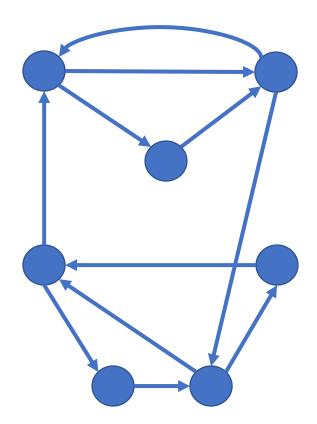


- Create the de Bruijn graph for the following string, using k=3
 - TTATGCCATGGGATGAA

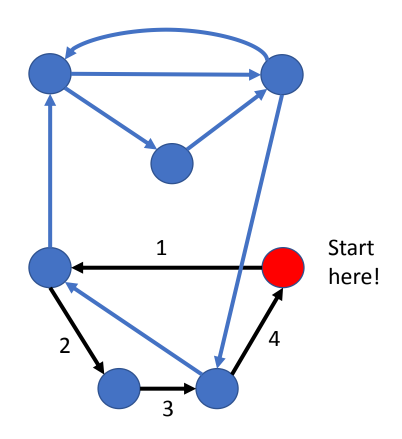


How to find a Eulerian cycle?

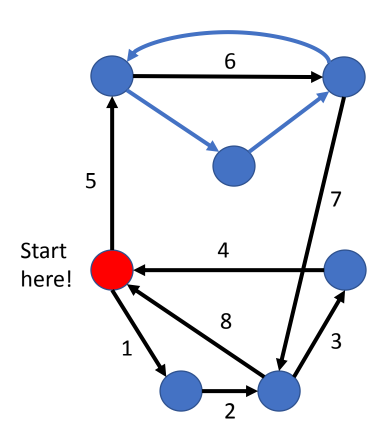
Random walking of an ant in the graph



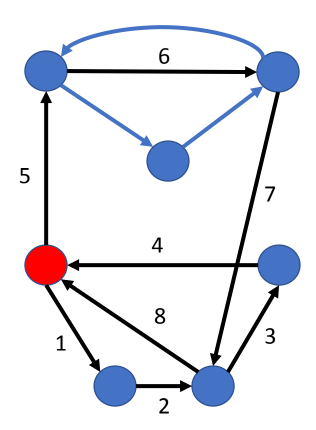
• Random walking of an ant: get stuck in the starting vertex



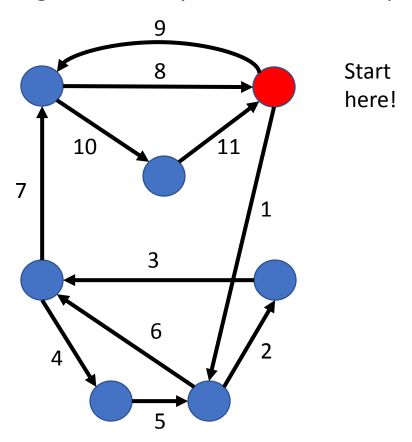
 Random walking of an ant: now start from a different vertex with unused edge, finishing the same cycle first before exploring new path



 Random walking of an ant: now get stuck again without finishing all edges

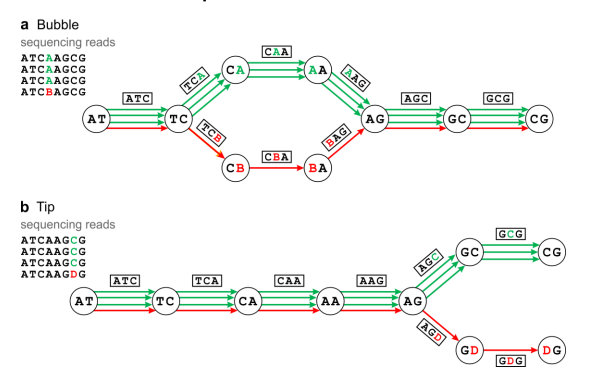


 Random walking of an ant: now start from a different vertex with unused edge, finishing the same cycle first before exploring new path



Error correction in de bruijn graph

• Erroneous reads (marked in red) introduce k false k-mers to the graph, resulting in additional spurious branches, so-called "bubbles" and "tips"

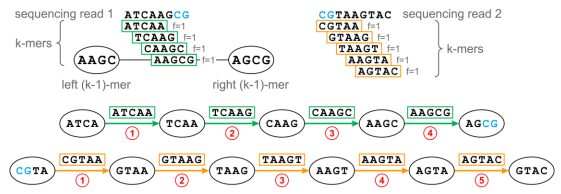


What is the appropriate k-mer?

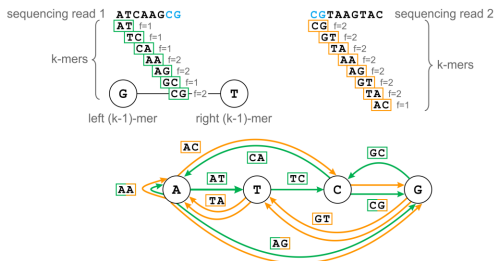
- Tools such as kmerGenie (http://kmergenie.bx.psu.edu/) and Velvet Advisor (dna.med.monash.edu.au/~torsten/velvet_advisor) have been developed to predict the optimal value for k for a given read dataset, based on read length and k-mer frequencies.
- Other strategies may include the merging of assemblies produced with different k-mer sizes or the use of a multi k-mer assembly method.

Influence of k-mer length on assembly

a de Bruijn graph for high k-mer lengths



b de Bruijn graph for low k-mer lengths



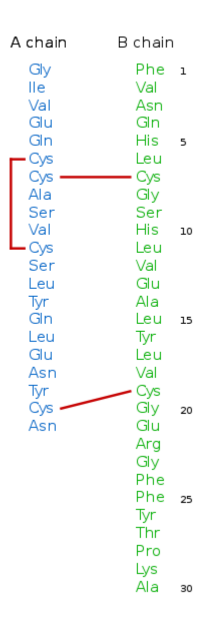
A fun hypothetical case study: Frederick Sanger's insulin sequencing study

- Frederick Sanger (1918 2013) was a British biochemist who twice won the Nobel Prize in Chemistry,
- In 1958, he was awarded a Nobel Prize in Chemistry "for his work on the structure of proteins, especially that of insulin"
- In 1980, Walter Gilbert and Sanger shared half of the chemistry prize "for their contributions concerning the determination of base sequences in nucleic acids"



Sequencing insulins

- Sanger determined the complete amino acid sequence of the two polypeptide chains of bovine insulin, A and B, in 1952 and 1951, respectively.
- In determining these sequences, Sanger proved that proteins have a defined chemical composition



Details on sequencing insulin

FREDERICK SANGER

The chemistry of insulin

Nobel Lecture, December 11, 1958

It is great pleasure and privilege for me to give an account of my work on protein structure and I am deeply sensitive of the great honour that has been done to me in recognizing my work in this way. Since the work on insulin has extended over about 12 years it will be necessary to give a somewhat simplified account and to omit most of the work that did not contribute directly to the main problem, the determination of the structure of a protein.

The Amino-acid Sequence in the Phenylalanyl Chain of Insulin

1. THE IDENTIFICATION OF LOWER PEPTIDES FROM PARTIAL HYDROLYSATES

By F. SANGER (Beit Memorial Fellow) AND H. TUPPY*
Biochemical Laboratory, University of Cambridge

(Received 17 January 1951)

When insulin is oxidized with performic acid, the —S—S— bridges of the cystine residues are broken by conversion to —SO₃H groups (Sanger, 1949a) and the molecule is split into its separate polypeptide chains. From the oxidized insulin two fractions could be isolated: an acidic fraction A, which contained only glycyl N-terminal residues (see below) and no basic amino-acids, and a basic fraction B, having phenylalanyl N-terminal residues. These appeared to be the only significant fractions present.

From a study of the partial hydrolysis products of the dinitrophenyl (DNP) derivatives of the two fractions it was possible to determine the sequence of the amino-acids adjoining the N-terminal residues partial hydrolysis might yield considerable information about the overall amino-acid sequence in these fractions. Consden, Gordon & Martin (1947) have described a method for the fractionation of lower peptides using paper chromatography which was successfully used to determine the pentapeptide sequence of 'gramicidin S' (Consden, Gordon, Martin & Synge, 1947). The present paper describes the application of this technique to partial acid hydrolysates of fraction B and the determination of a number of amino-acid sequences.

Throughout this paper the abbreviations for the amino-acid residues suggested by Brand & Edsall (1947) are used. These are listed in Table 1. In

Deduction of short peptides

- Sanger partially hydrolysed the insulin into short peptides, either with hydrochloric acid or using an enzyme such as trypsin.
- The mixture of peptides was fractionated in two dimensions on a sheet of filter paper, first by electrophoresis in one dimension and then, perpendicular to that, by chromatography in the other.
- The different peptide fragments of insulin, detected with ninhydrin, moved to different positions on the paper, creating a distinct pattern that Sanger called "fingerprints".
- By repeating this type of procedure Sanger was able to determine the sequences of the many peptides generated using different methods for the initial partial hydrolysis. These could then be assembled into the longer sequences to deduce the complete structure of insulin.

The peptides that Sanger found in fraction B

- We want to do a fun exercise:
 - Only Dipeptides and Tripeptides are shown below

		Table 15. Pepe	tides obtained from	phenylalanyl cha	in of insulin (fraction B)		
Dipeptides	$(B4\beta2)$ $(B1\alpha2)$	His Leu.CySO ₃ H	Thr. Pro Lys. Ala (B188) (B2γ4)	Gly.Glu Arg.Gly $(B 1\delta 1)$ $(B 2\gamma 5)$ Glu.Arg $(B 1\gamma 3)$	Tyr.Leu Val.CySO ₂ H $(B3\beta10)$ $(B1\alpha4)$ Leu.Val CySO ₂ H.Gly $(B3\beta7)$ $(B1\alpha1)$	Ser. His Leu. Val Glu. Ala (B2y2) (B3\$7) (B1\$4) His. Leu Val. Glu (B2y8) (B1\$7)	Ala. Leu $(B 1 \delta 11)$ Gly. Phe $(B 1 \gamma 12)$
Tripeptides	Phe. Val. Asp (B 1 \textit{\beta}13) Val. Asp. Glu (B 1 \textit{\beta}10)	His.Leu.CySO ₃ H $(B 1\gamma 4)$ Leu.CySO ₃ H.Gly $(B 1\infty 5)$ Glu.His.Leu $(B 1\gamma 7)$	Pro. Lys. Ala (B 5 y 6)	Gly.Glu.Orn (B 5 y 1) [Glu, Arg, Gly] (B 1 y 2)	Tyr. Leu. Val $(B3\beta12)$ Leu. Val. CySO ₃ H $(B1\alpha8)$ Val. CySO ₂ H. Gly $(B1\alpha3)$	Ser. His. Leu $(B2\sqrt{7})$ Val. Glu. Ala $(B1\sqrt{10})$ Leu. Val. Glu $(B2\sqrt{10})$	Ala.[Tyr, Leu] (B 3 \(\rho\rho\rho\rho)

Fun exercise: assemble insulin from dipeptides and tripeptides for Sanger

• What if Sanger knew de Bruijn graph in 1951? Can he use the information below to assemble isulin to a few contigs?

Dipeptide	Ala.Leu		Tripeptide				
Dipeptide	Arg.Gly		Tipeptiae				
	Asp.Glu						
	Cys.Gly			Glu.Arg.Gly			
	Cys.Gly			Glu.His.Leu			
	Glu.Ala			Gly.Glu.Arg			
	Glu.Arg						
	Glu.His			His.Leu.Cys			
	Gly.Glu			Leu.Cys.Gly			
	Gly.Phe			Leu.Val.Cys			
	His.Leu			Leu.Val.Glu			
	His.Leu			Phe.Val.Asp			
	Leu.Cys						
	Leu.Val			Pro.Lys.Ala			
	Leu.Val			Ser.His.Leu			
	Lys.Ala			Tyr.Leu.Val			
	Phe.Val			Val.Asp.Glu			
	Ser.His			Val.Cys.Gly			
	Thr.Pro			Val.Glu.Ala			
	Tyr.Leu			van Ora. Aria			
	Val.Asp						
	Val.Cys						
	Val.Glu						