# Lecture 4: Genome assembly & annotation

- Genome assembly
  - o de Bruijn graphs
- Genome annotation
  - Hidden Markov Models

#### Why sequence the genome?

- Determine the "complete" sequence of a haploid genome.
  - Previously "snippets" of the genome were available.
- Identify the sequence and location of every protein coding gene.
- Use as a "map" with which to track the location and frequency of genetic variation.
- Unravel the genetic architecture of inherited and somatic traits/diseases.
- To understand genome and species evolution.

"All the News That's Fit to Print"

# The New Hork Times

Late Edition

New York: Today, afternoon thunderstorms, high 88. Tonight, showers end, low 67. Tomorrow, partly cloudy with showers late, high 81. Yesterday, high 88, low /4. Weather map, Page D8.

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\$1 beyont the greater New York metropolitan area.

The New York Times

75 CENTS

#### Genetic Code of Human Life Is Cracked by Scientists

#### JUSTICES REAFFIRM MIRANDA RULE, 7-2; A PART OF 'CULTURE'

#### By LINDA GREENHOUSE

WASHINGTON, June 26 - The Sapreme Court reaffirmed the Miranda decision teday by a 7-to-2 vote that erased a shadow over one of the most famous rulings of modern times and acknowledged that the Miranda warnings "have become part of our national culture."

The court said in an opinion by Chief Justice William H. Rehnquist that because the 1966 Miranda decision "armounced a constitutional rule," a statute by which Congress had sought to overrule the decision was itself unconstitutional.

Miranda had appeared to be in jeopardy, both because of that longignored but recently rediscovered law, by which Cengress had tried to overrule Miranda 32 years ago, and because of the court's perceived hostility to the original decision.

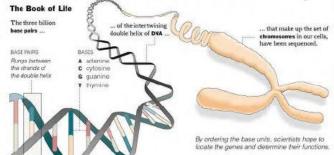
The chief justice said, though, that the 1968 law, which replaced the Miranda warnings with a case-by-case test of whether a confession was voluntary, could be upheld only if the Supreme Court decided to overturn Miranda, But with Miranda having

Justices Antonin Scalia and Clarence Thomas cast the dissenting

The decision overtumed a ruling last year by the federal appeals court ir Richmond, Va., which held that Congress was entitled to the last word because Miranda's presumption that a confession was not volunlary unless preceded by the warnings was not required by the Consti-

The decision today -only 14 pages long, ir Chief Justice Rehnquist's typically spare style - brought an abrupt and to one of the odder episodes in the court's recent history, an intense and strangely delayed relighting of a previous generation's battle over the rights of criminal suspects. Miranda v. Arizona was a hallmank of the Warren Court, and Chief Justice Rehnquist, despite his record as an early and tenacious critic of the decision, evidently did not want its repudiation to be an imprint of his own tenure.

There was considerable drama in the courtroom today as the chief justice announced that he would de-



#### **Science Times** A specialissue

- · Putting the genome to work.
- Some information has already paid research dividends
- Two research methods two results.
- From Mendel to helix to genome.
- More articles. charts and photos of the genome effort.

Section F

Francis S. Collins. head of the Human Genome Project. left, with J. Craig Venter, head of Celera Genomics. after the announcement vesterday that they had finished the first survey of

the human genome.



mics, and Dr. Francis S. Collins, director of the National Human Genome Research Institute, praised each other's contributions and signaled a spirit of cooperation from now on, even though the two efforts

The human genome, the ancient script that has now been deciphered.

#### A SHARED SUCCESS

#### 2 Rivals' Announcement Marks New Medical Era, Risks and All

#### By NICHOLAS WADE

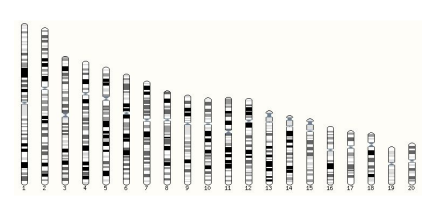
WASHINGTON, June 26 - In an achievement that represents a pinnacle of human self-knowledge, two rival groups of scientists said today that they had deciphered the hereditary script the set of instructions that defines the human organism.

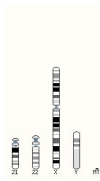
"Today we are learning the language in which God created life." President Clinton said at a White House ceremony attended by members of the two teams, Dr. James D. Watson, co-discoverer of the structure of DNA, and, via satellite, Prime Minister Tony Blair of Britain. [Excernts. Page D8.1

The teams' leaders, Dr. J. Craig Verter, president of Celera Genowill remain firmly independent.

consists of two sets of 23 giant DNA

The Human Genome – Summary



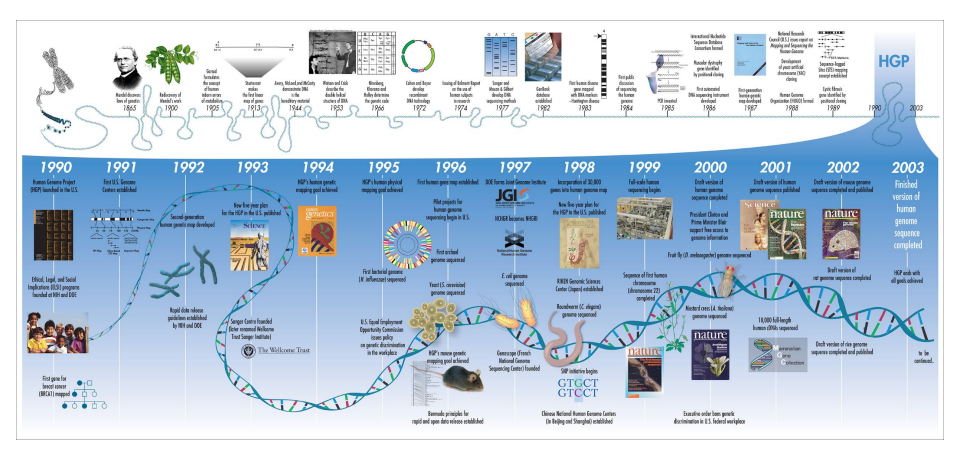


Assembly GRCh38.p12 (Genome Referei **Base Pairs** 3,609,003,417 Golden Path Length 3,096,649,726 Annotation provider Ensembl Annotation method Full genebuild Genebuild started Jan 2014 Genebuild released Jul 2014 Genebuild last updated/patched Jul 2018 **Database version** 95.38 **GENCODE 29** Gencode version

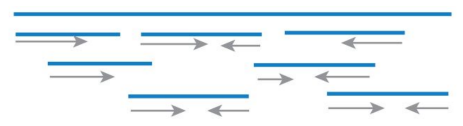
#### Gene counts (Primary assembly)

incl 650 readthrough)
ncl 284 readthrough)
incl 8 readthrough)

http://useast.ensembl.org/Homo sapiens/Location/Genome



#### Genome assembly & annotation – Overview



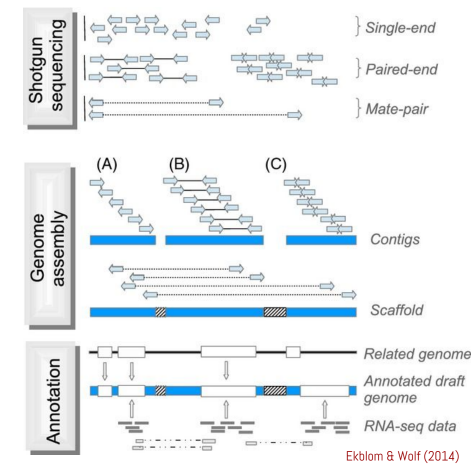
**read**: a short/long word that comes out of sequencer

**mate pair**: a pair of reads from two ends of the same insert fragment

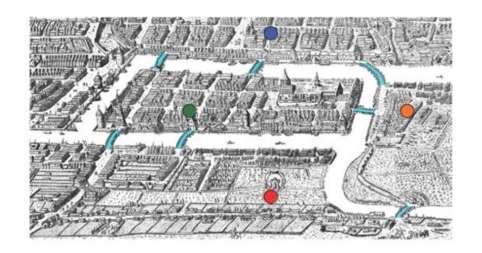
**contig**: a contiguous sequence formed by several overlapping reads with no gaps

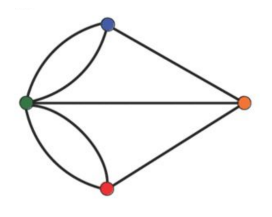
**scaffold**: an ordered and oriented set of contigs, usually by mate pairs

**consensus sequence**: derived from the sequence multiple alignment of reads in a contig



# Introduction to graph theory





#### Bridges of Königsberg

Can every part of the city be visited by walking across each of the seven bridges exactly once and returning to one's starting location?



Graph: (Nodes, Edges, Weights)

(Eulerian) Path exists if the graph contains zero or two vertices that have an odd degree.

## de Bruijn graphs: the 'superstring problem'

Find a shortest circular 'superstring' that contains all possible 'substrings' of length k (k-mers) over a given alphabet.

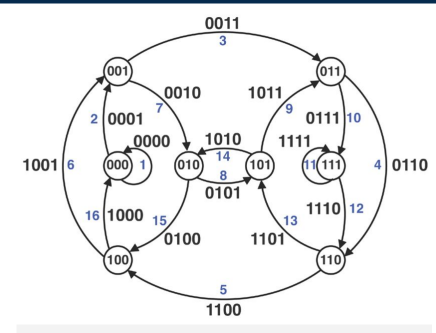
There exist n<sup>k</sup> k-mers in an alphabet containing n symbols:

- alphabet: 0 & 1
- all 3-mers: 000, 001, 010, 011, 100, 101, 110, 111.
- The circular superstring **0001110100** contains all 3-mers & each 3-mer exactly once.



How can we construct a superstring for all k-mers in the case of an arbitrary value of k and an arbitrary alphabet?

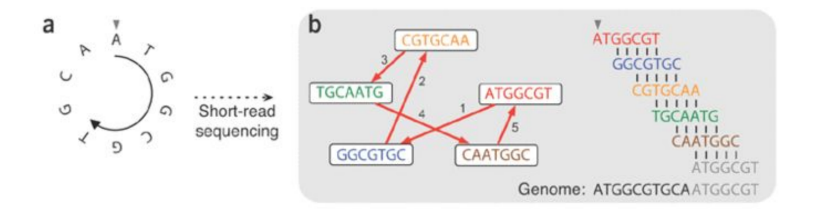
Construct a **directed graph**: all prefixes & suffixes as nodes; all k-mers as edges.



k = 4 | Two-character alphabet: digits 0 & 1

Does this graph have an Eulerian cycle? [Balanced?]

Following the blue numbered edges in order from 1 to 16 traces the cyclic superstring 0000110010111101.



Nodes: Reads

**Edges**: Alignments between reads (≥ 5 bases)

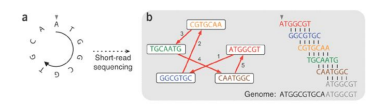
**Genome**: Walk along a Hamiltonian cycle (visit each vertex exactly once) to combine alignments between successive reads and reconstruct the full circular genome.

Four hidden assumptions that **do not hold** for real sequencing:

- 1. We can generate all k-mers present in the genome.
- All k-mers are error free.
- 3. Each k-mer appears at most once in the genome.
- 4. The genome consists of a single circular chromosome.

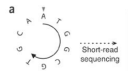
E.g., a technology that generates 100-nucleotide long reads:

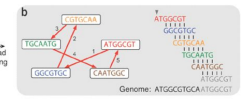
- may miss some 100-mers present in the genome (even if the read coverage is high)
- the 100-mers that it does generate typically have errors.



Break reads into shorter k-mers!

Resulting k-mers often represent nearly all k-mers from the genome for sufficiently small k.





Nodes: k-mers

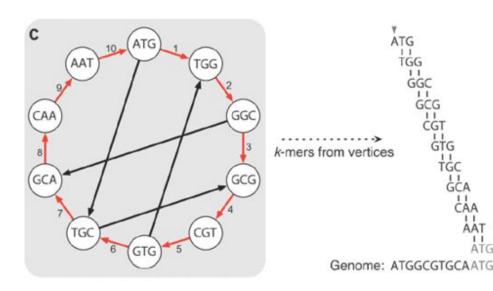
Edges: Overlap between k-mers

Genome: Walk the Hamiltonian cycle

Large genomes result in too many reads:

- $10^6$  reads  $\rightarrow 10^{12}$  pairwise alignments.
- $10^9$  reads  $\rightarrow 10^{18}$  alignments.

There is no known efficient algorithm for finding a Hamiltonian cycle in a large graph with millions (let alone billions) of nodes.



Finding a path that visits all *edges* of a graph exactly once is much easier!

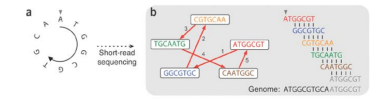
Nodes: (k-1)-mers [in- & out-degrees?]

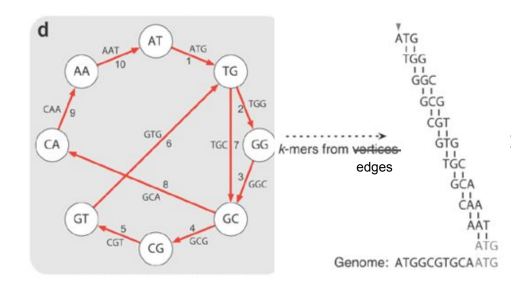
Edges: k-mers

**Genome**: Walking the Eulerian cycle

Send out an ant, find a cycle; Note edges traversed, send out another ant, ... until all of the graph's edges have been explored. Combine all cycles to form an Eulerian cycle!

Computationally tractable; time roughly proportional to the number of edges.

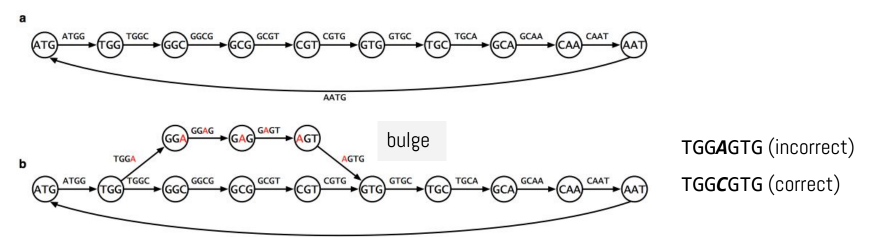


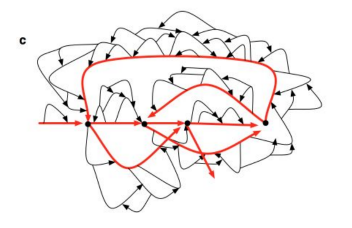


#### de Bruijn graphs for sequence assembly – not so easy w/ real data

- Not all k-mers in the genome
  - Read breaking
- Errors in reads
  - Error correcting reads
  - Removing bulges in de Bruijn graphs
- DNA repeats
  - Incorporating k-mer multiplicity
- Multiple and linear chromosomes
  - Cycles to paths
- Unsequenced regions
  - Scaffolds

## de Bruijn graphs from reads with sequencing errors



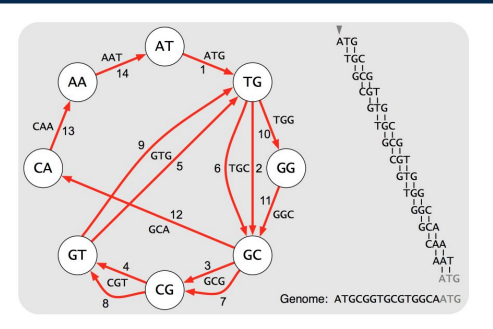


The process of bulge removal should leave only the red edges remaining, yielding an Eulerian path in the resulting graph.

#### de Bruijn graphs for sequence assembly – not so easy w/ real data

- Not all k-mers in the genome
  - Read breaking
- Errors in reads
  - Error correcting reads
  - Removing bulges in de Bruijn graphs
- DNA repeats [E.g., ATGCATGC → four 3-mers: ATG, TGC, GCA & CAT]
  - Incorporating k-mer multiplicity
- Multiple and linear chromosomes
  - Cycles to paths
- Unsequenced regions
  - Scaffolds

## de Bruijn graphs for dealing with repeats



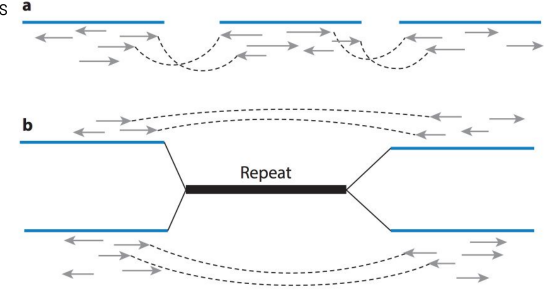
Genome: ATGCGTGCGTGGCA

Incorporate k-mer multiplicity:

- Four 3-mers TGC, GCG, CGT, and GTG: multiplicity 2
- Six 3-mers ATG, TGG, GGC, GCA, CAA, and AAT: multiplicity 1

## de Bruijn graphs for sequence assembly – not so easy w/ real data

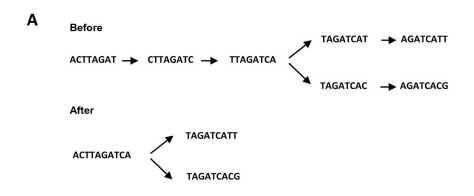
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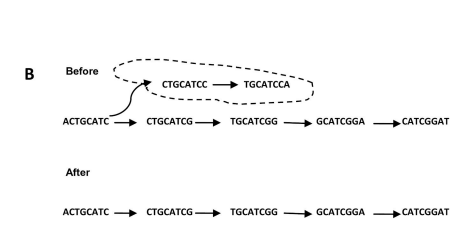


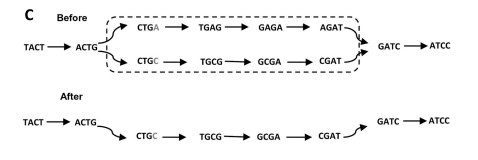
## Algorithms for genome assembly

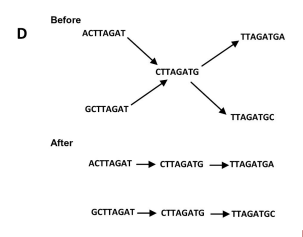
- Error detection and correction based on sequence composition of the reads.
- Graph construction to represent reads/k-mers and their shared sequence.
- Graph adjustments:
  - Reduction of simple non-intersecting paths to single nodes.
  - Removal of error-induced paths (recognized as spurs or bubbles).
  - Collapse of polymorphism-induced complexity (bubbles).
  - Simplification of tangles (using information outside the graph: individual, paired-end, or mate-pair reads to constraints on path distance & outcome.
- Conversion of reduced paths to contigs and scaffolds.
- Reduction of alignments to a consensus sequence.

#### Simplifying de Bruijn graphs

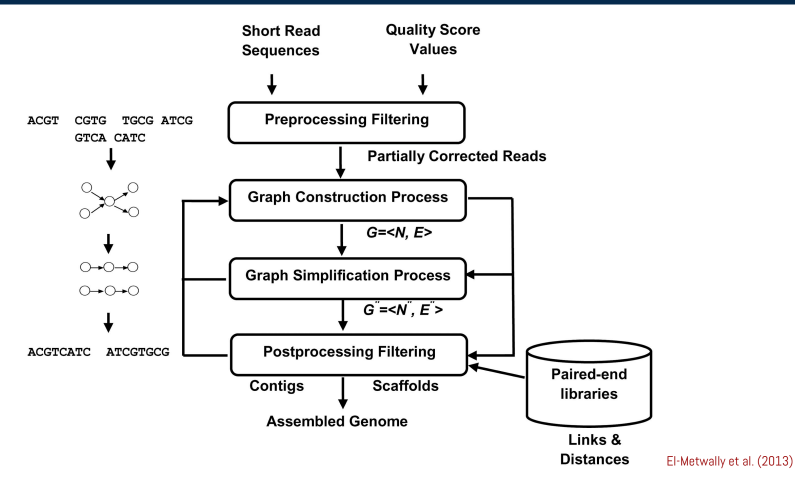








## Algorithms for genome assembly



## Algorithms for genome assembly

- ullet New sequencing technologies o a different best computational strategy.
- Factors that influence the choice of algorithms:
  - Quantity of data (read length and coverage)
  - Quality of data (including error rates)
  - Genome structure (e.g., GC content and the number and size of repeated regions).
- de Bruijn graphs are best suited for current short-read sequencing technologies
  - Produce very large numbers of reads
  - Can represent genomes with repeats [Overlap methods need to mask repeats > read length]
- Long-read technology growing at a rapid pace.

# Some modern genome assemblers

Assemblers	Technology	Availability	Notes
Genome assen	nblers		
ALLPATHS-LG	Illumina, Pacific Biosciences	ftp://ftp.broadinstitute.org/pub/ crd/ALLPATHS/Release-LG	Requires a specific sequencing recipe (BOX 3)
SOAPdenovo	Illumina	http://soap.genomics.org.cn/ soapdenovo.html	Also used for transcriptome and metagenome assembly
Velvet	Illumina, SOLiD, 454, Sanger	http://www.ebi.ac.uk/~zerbino/ velvet	May have substantial memory requirements for large genomes
ABySS	Illumina, SOLiD, 454, Sanger	http://www.bcgsc.ca/platform/ bioinfo/software/abyss	Also used for transcriptome assembly

#### Paper discussion

- Ask questions & offer answers/thoughts.
  - Let's collectively engage.
  - Good for you & the presenters.
- Also be ready with all the questions you had from your reading.