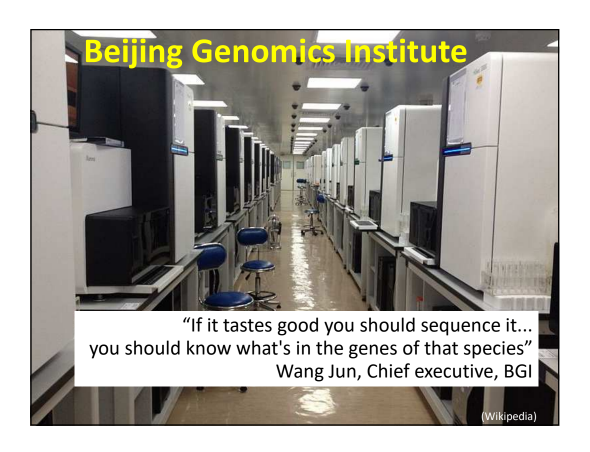
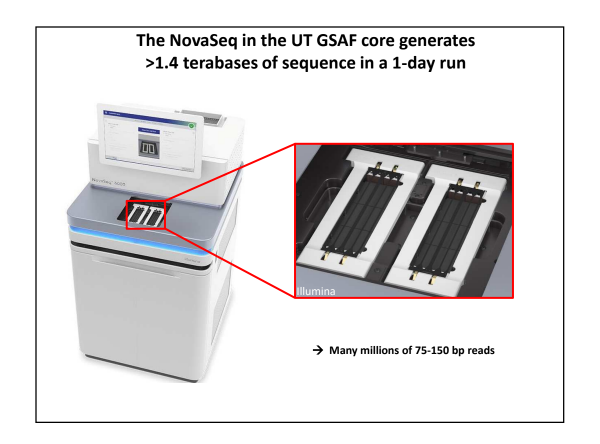
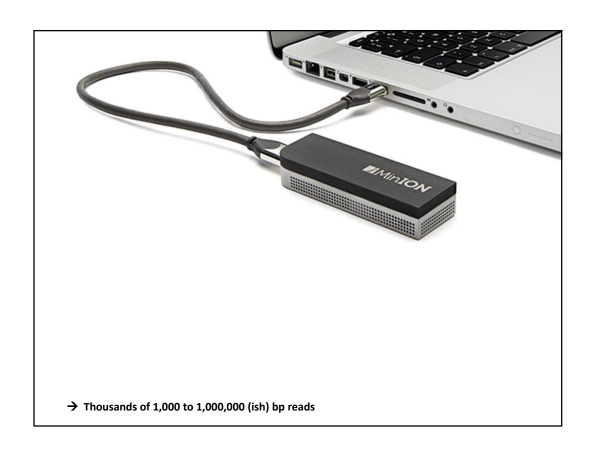
# **Assembling Genomes**

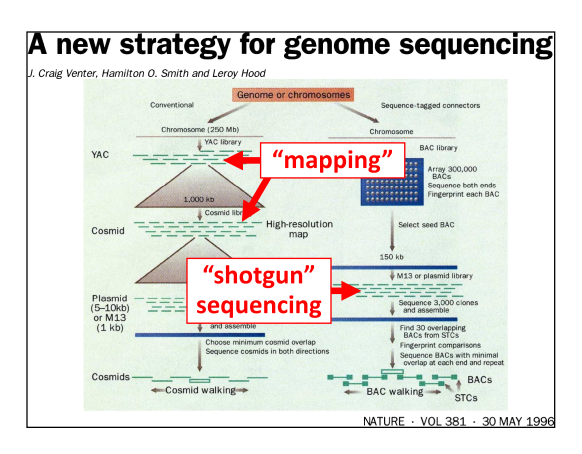
BCH394P/364C Systems Biology / Bioinformatics Edward Marcotte, Univ of Texas at Austin





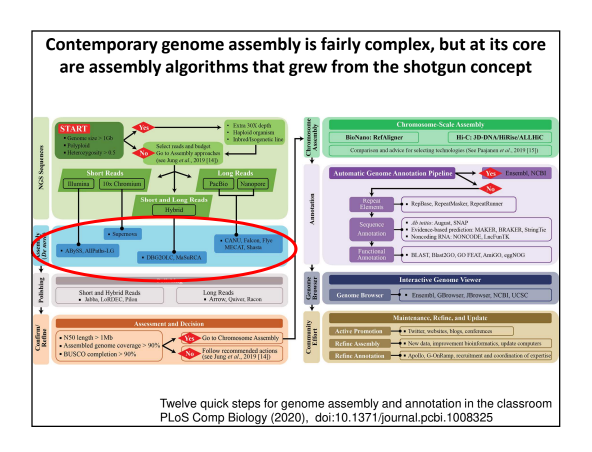


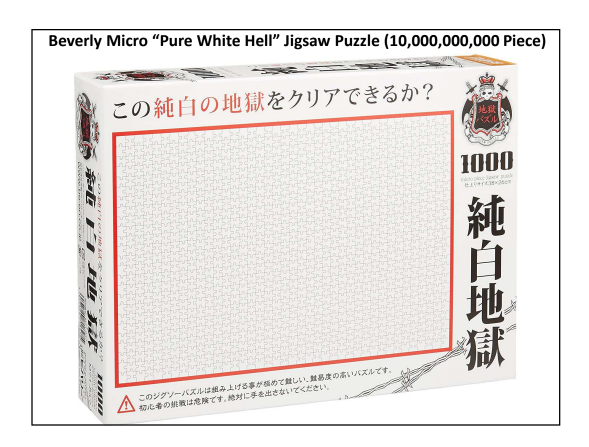




CLONE LIB	RARIES USED FOR GENON AND SEQUENCING	ME MAPPING
Vector	Human-DNA insert size range	Number of clones required to cover the human genome
Yeast artificial chromosome (YAC)	100–2,000 kb	3,000 (1,000 kb)
Bacterial artificial chromosome (BAC)	80–350 kb	20,000 (150 kb)
Cosmid	30-45 kb	75,000 (40 kb)
Plasmid	3–10 kb	600,000 (5 kb)
M13 phage	1 kb	3,000,000 (1 kb)

NATURE · VOL 381 · 30 MAY 1996





# Thinking about the basic shotgun concept

- Start with a very large set of random sequencing reads
- How might we match up the overlapping sequences?
- How can we assemble the overlapping reads together in order to derive the genome?

## Thinking about the basic shotgun concept

- At a high level, the first genomes were sequenced by comparing pairs of reads to find overlapping reads
- Then, building a graph (i.e., a network) to represent those relationships
- The genome sequence is a "walk" across that graph

# The "Overlap-Layout-Consensus" method

**Overlap**: Compare all pairs of reads

(allow some low level of mismatches)

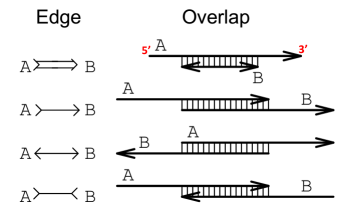
**<u>Layout</u>**: Construct a graph describing the overlaps

sequence overlap read

Simplify the graph read
Find the simplest path through the graph

<u>Consensus</u>: Reconcile errors among reads along that path to find the consensus sequence

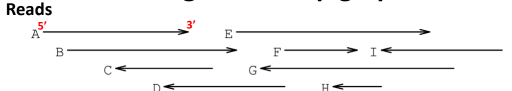
# **Building an overlap graph**



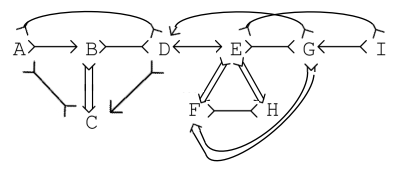


EUGENE W. MYERS. *Journal of Computational Biology*. Summer 1995, 2(2): 275-290

# Building an overlap graph

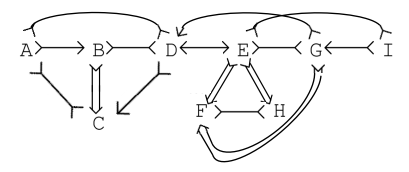


## Overlap graph



EUGENE W. MYERS. *Journal of Computational Biology*. Summer 1995, 2(2): 275-290 (more or less)

# Simplifying an overlap graph



1. Remove all contained nodes & edges going to them

EUGENE W. MYERS. *Journal of Computational Biology*. Summer 1995, 2(2): 275-290 (more or less)

# Simplifying an overlap graph



2. Transitive edge removal: Given A - B - D and A - D, remove A - D

EUGENE W. MYERS. *Journal of Computational Biology*. Summer 1995, 2(2): 275-290 (more or less)

# Simplifying an overlap graph

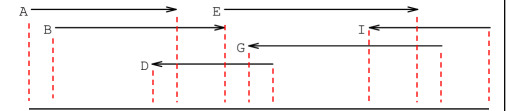
$$A \longrightarrow B \longrightarrow C \longrightarrow E \longrightarrow G \longleftarrow I$$

3. If un-branched, calculate consensus sequence
If branched, assemble un-branched bits and then decide
how they fit together

EUGENE W. MYERS. *Journal of Computational Biology*. Summer 1995, 2(2): 275-290 (more or less)

# Simplifying an overlap graph

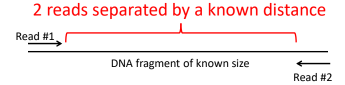
$$A \longrightarrow B \longrightarrow C \longrightarrow E \longrightarrow G \longleftarrow I$$



"contig" (assembled contiguous sequence)

EUGENE W. MYERS. *Journal of Computational Biology*. Summer 1995, 2(2): 275-290 (more or less)

# This basic strategy was used for most of the early genomes. Also useful: "mate pairs"



Contigs can be ordered using these paired reads

Contig #1

Contig #2

to produce "scaffolds"

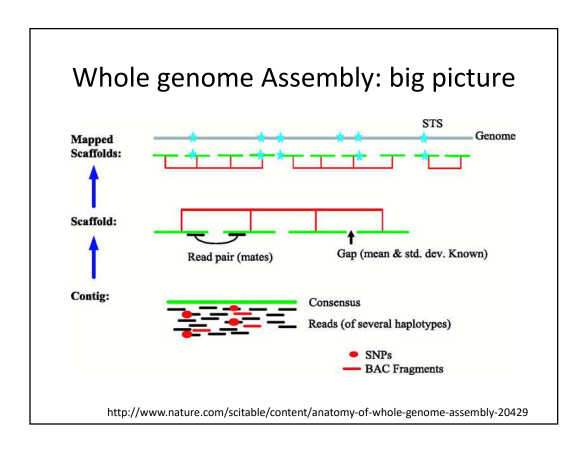
# GigAssembler (used to assemble the public human genome project sequence)



Jim Kent

**David Haussler** 

Let's take a little walk through history to see what they did...

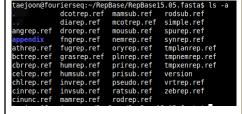


# GigAssembler - Preprocessing

- 1. Decontaminating & Repeat Masking.
- 2. Aligning of mRNAs, ESTs, BAC ends & paired reads against initial sequence contigs.
  - psLayout → BLAT
- 3. Creating an input directory (folder) structure.

```
chr1/
chr1/contig1.e
chr1/contig1.a
chr1/contig1.b
chr1/contig1.d
chr3/
chr2/
chr2/contig2.d
chr2/contig2.b
chr2/contig2.a
chr2/contig2.a
```

# RepBase + RepeatMasker

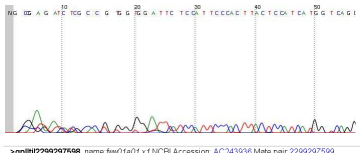


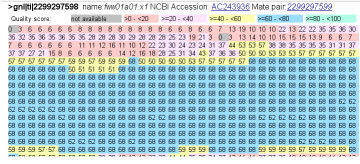
# GigAssembler: Build merged sequence contigs ("rafts")



**Figure 1** Two sequences overlapping end to end. The sequences are represented as dashes. The aligning regions are joined by vertical bars. End-to-end overlap is an extremely strong indication that two sequences should be joined into a contig.







# Sequencing quality (Phred Score)

$$Q = -10 \; \log_{10} P \hspace{-0.4cm} \longleftarrow \hspace{-0.4cm} \begin{array}{c} \textit{Base-calling} \\ \textit{Error} \\ \textit{Probability} \end{array}$$

or

$$P = 10^{\frac{-Q}{10}}$$

### Phred quality scores are logarithmically linked to error probabilities

Phred Quality Score	Probability of incorrect base call	Base call accuracy
10	1 in 10	90 %
20	1 in 100	99 %
30	1 in 1000	99.9 %
40	1 in 10000	99.99 %
50	1 in 100000	99.999 %

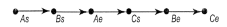
http://en.wikipedia.org/wiki/Phred quality score

We're going to skip the remaining details of GigAssembler (mainly of historical interest now) to get to the key strategy for assembling all of the various contigs and paired end reads into a genome

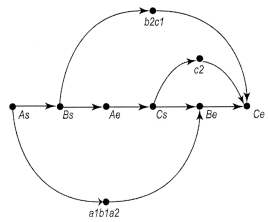
# GigAssembler: Build a "raft-ordering" graph



**Figure 4** Three overlapping draft clones: A, B, and C. Each clone has two initial sequence contigs. Note that initial sequence contigs a1, b1, and a2 overlap as do b2 and c1.



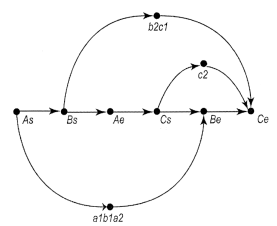
**Figure 5** Ordering graph of clone starts and ends. This represents the same clones as in Fig. 4. (As) The start of clone A; (Ae) the end of clone A. Similarly Bs, Be, Cs, and Ce represent the starts and ends of clones B and C.



**Figure 6** Ordering graph after adding in rafts. The initial sequence contigs shown in Fig. 4 are merged into rafts where they overlap. This forms three rafts: a1b1a2, b2c1, and c2. These rafts are constrained to lie between the relevant clone ends by the addition of additional ordering edges to the graph shown in Fig. 5.

# GigAssembler: Build a "raft-ordering" graph

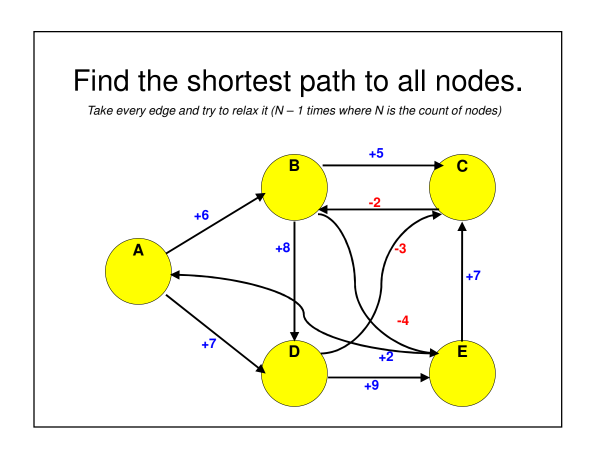
- Add information from mRNAs, ESTs, paired plasmid reads, BAC end pairs: building a "bridge"
  - Different weight to different data type: (mRNA ~ highest)
  - Conflicts with the graph as constructed so far are rejected.
- Build a sequence path through each raft.
- · Fill the gap with N's.
  - 100: between rafts
  - 50,000: between bridged barges

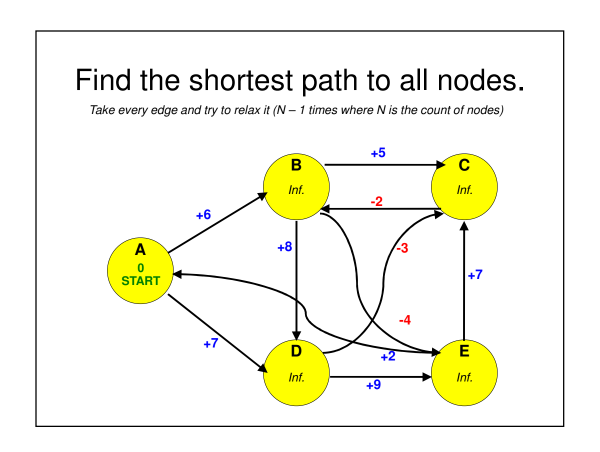


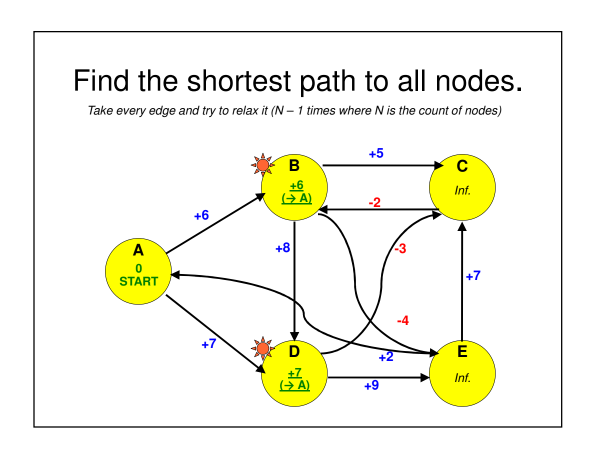
**Figure 6** Ordering graph after adding in rafts. The initial sequence contigs shown in Fig. 4 are merged into rafts where they overlap. This forms three rafts: a1b1a2, b2c1, and c2. These rafts are constrained to lie between the relevant clone ends by the addition of additional ordering edges to the graph shown in Fig. 5.

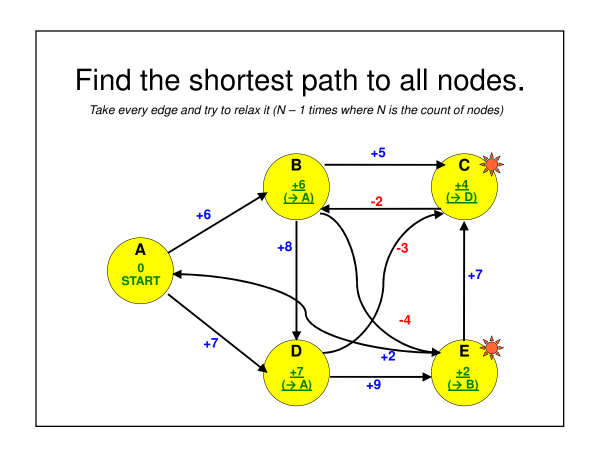
# Finding the shortest path across the ordering graph using the Bellman-Ford algorithm

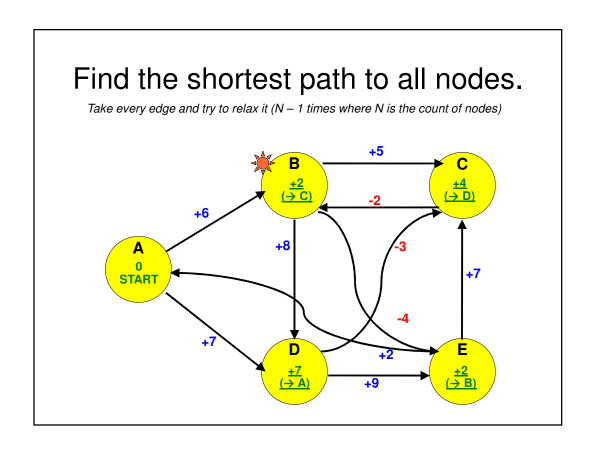
http://compprog.wordpress.com/2007/11/29/one-source-shortest-path-the-bellman-ford-algorithm/

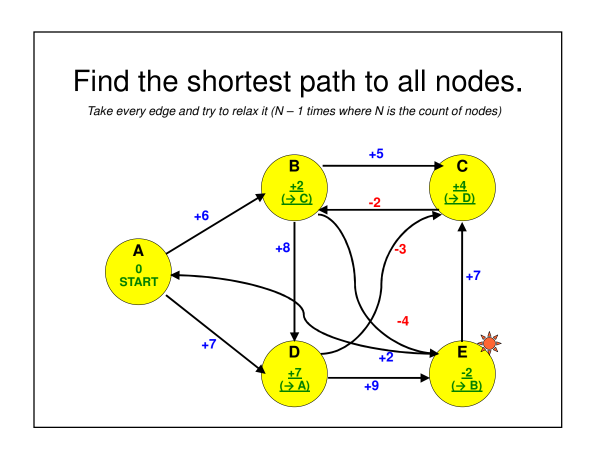


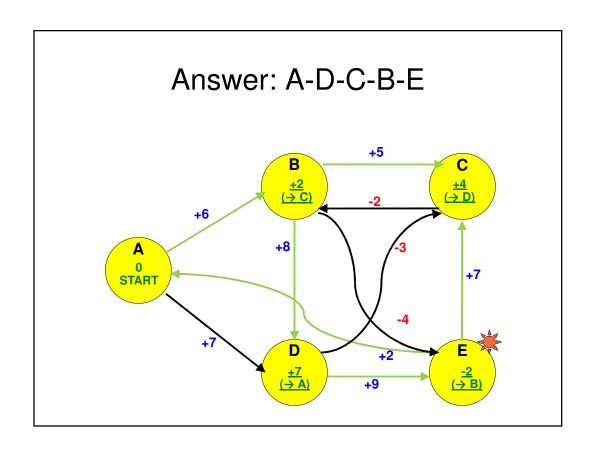


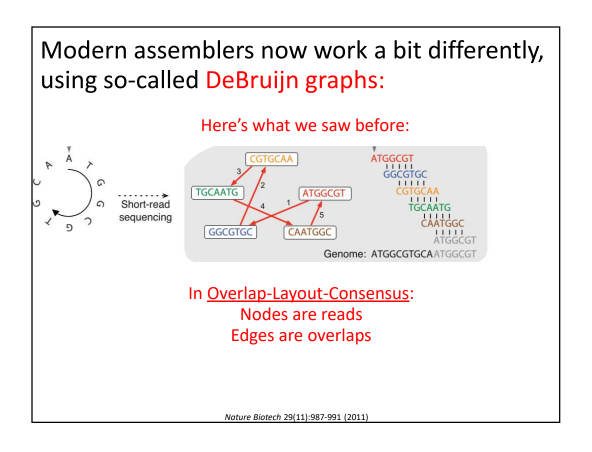


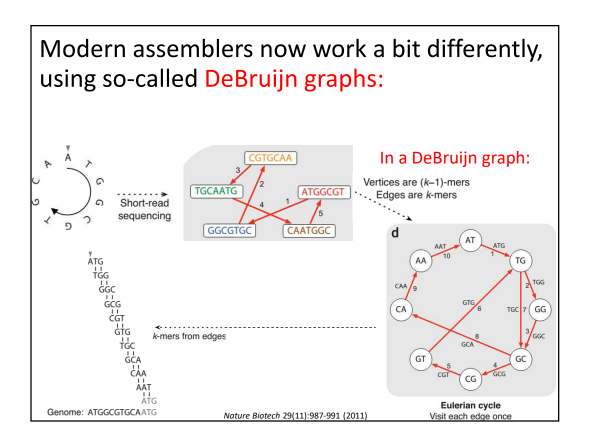












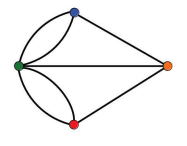
## Why Eulerian?

From Leonhard Euler's solution in 1735 to the 'Bridges of Königsberg' problem:

Königsberg (now Kaliningrad, Russia) had 7 bridges connecting 4 parts of the city. Could you visit each part of the city, walking across each bridge only once, & finish back where you started?

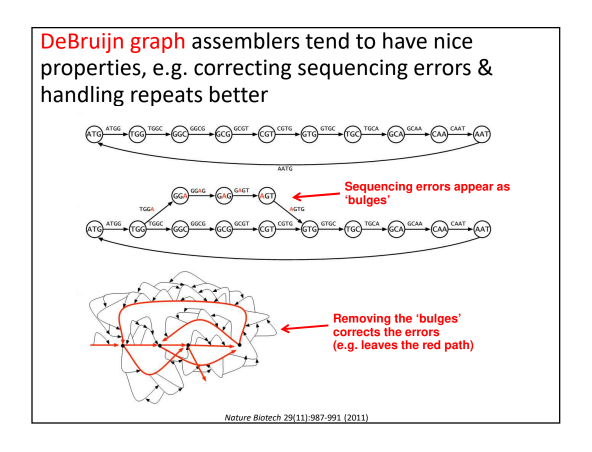


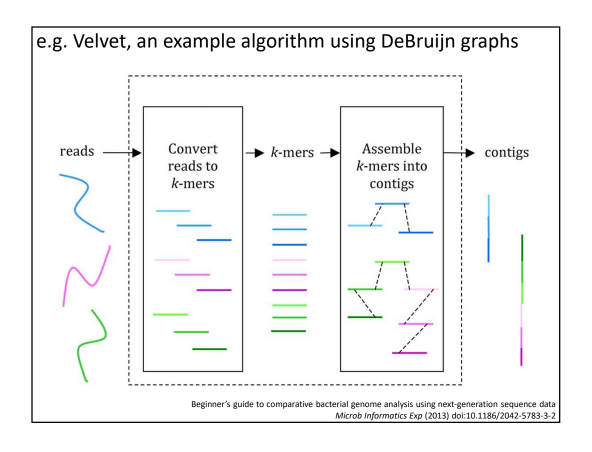
(Visiting every edge once = an Eulerian path)



Euler conceptualized it as a graph: Nodes = parts of city Edges = bridges

Nature Biotech 29(11):987-991 (2011)





# Once a reference genome is assembled, new sequencing data can 'simply' be mapped to the reference.

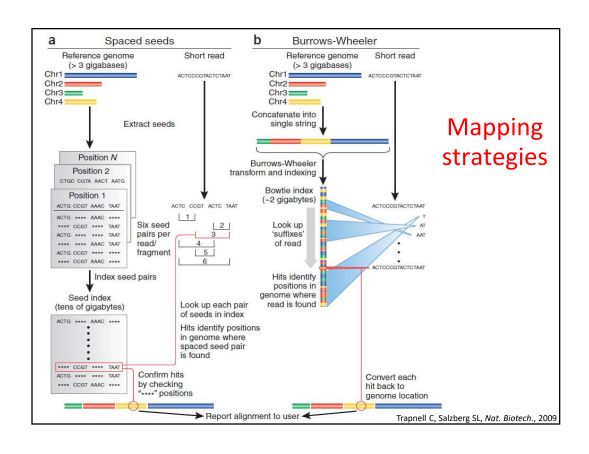
	reads		
	<u> </u>	<u>=</u>	
Reference genome 💳			

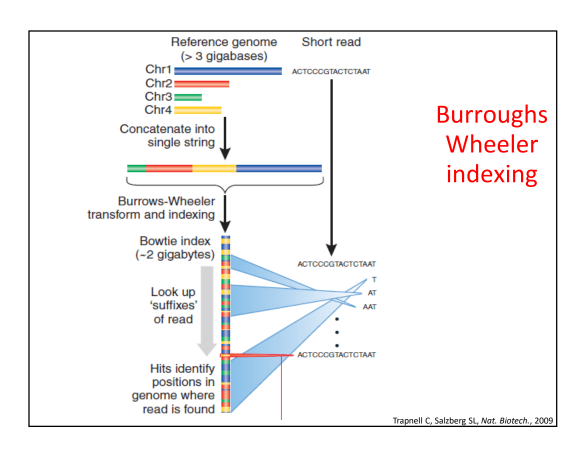
# Mapping reads to assembled genomes

Program	Website	Open source?	Handles ABI color space?	Maximum read length
Bowtie	http://bowtie.cbcb.umd.edu	Yes	No	None
BWA	http://maq.sourceforge.net/bwa-man.shtml	Yes	Yes	None
Maq	http://maq.sourceforge.net	Yes	Yes	127
Mosaik	http://bioinformatics.bc.edu/marthlab/Mosaik	No	Yes	None
Novoalign	http://www.novocraft.com	No	No	None
SOAP2	http://soap.genomics.org.cn	No	No	60
ZOOM	http://www.bioinfor.com	No	Yes	240

The list is a little longer now! e.g. see https://en.wikipedia.org/wiki/List\_of\_sequence\_alignment\_software#Short-Read\_Sequence\_Alignment

Trapnell C, Salzberg SL, Nat. Biotech., 2009

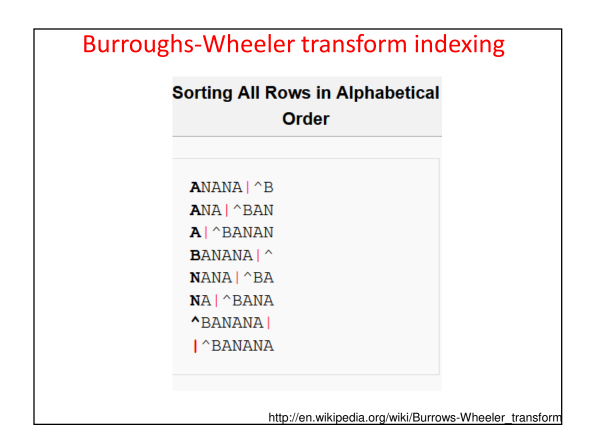




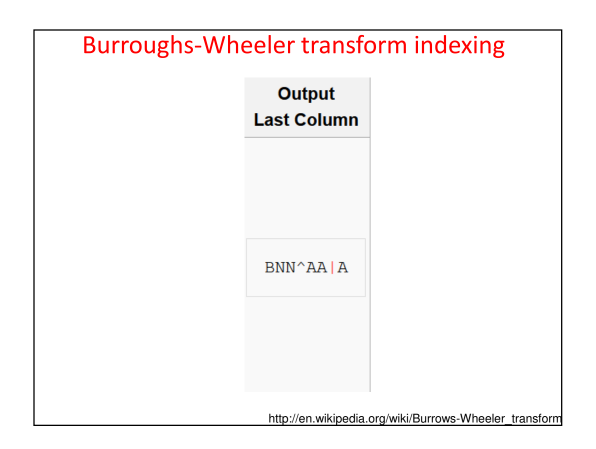
# BWT is often used for file compression (like bzip2), here used to make a fast 'lookup' index in a genome BWT = 'reversible block-sorting' Input SIX.MIXED.PIXIES.SIFT.SIXTY.PIXIE.DUST.BOXES This sequence is more compressible Output TEXYDST.E.IXIXIXXSSMPPS.B..E.S.EUSFXDIIOIIIT Reverse BWT Recovered SIX.MIXED.PIXIES.SIFT.SIXTY.PIXIE.DUST.BOXES input http://en.wikipedia.org/wiki/Burrows-Wheeler\_transform

# Input ABANANA | http://en.wikipedia.org/wiki/Burrows-Wheeler\_transform

# Burroughs-Wheeler transform indexing All Rotations ^BANANA | | ^BANANA A | ^BANAN NA | ^BANA ANA | ^BAN NANA | ^BA ANANA | ^B BANANA | ^B



# Taking Last Column ANANA | ^B ANA | ^BANA A | ^BANAN BANANA | ^BA NA | ^BANA NANA | ^BA NANA | ^BANA NANA | ^BANA NANA | ^BANA NANA | ^BANA NANA | ^BANANA ABANANA ABANANA ABANANA ABANANA http://en.wikipedia.org/wiki/Burrows-Wheeler\_transform



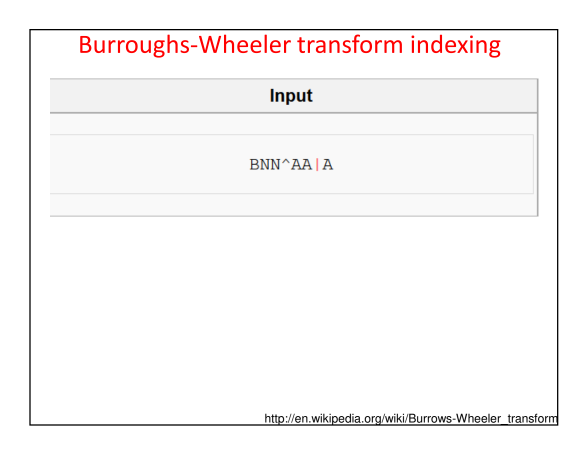
# Burroughs-Wheeler transform indexing

Transformation				
Input	All Rotations	Sorting All Rows in Alphabetical Order	Taking Last Column	Output Last Column
^BANANA	^BANANA     ^BANANA A   ^BANAN NA   ^BANA ANA   ^BAN NANA   ^BA ANANA   ^B BANANA   ^	ANANA   ^B ANA   ^BAN A   ^BANAN BANANA   ^ NANA   ^BA NA   ^BANA ^BANANA     ^BANANA	ANANA   ^B ANA   ^BAN A   ^BANAN BANANA   ^ NANA   ^BA NA   ^BANA ^BANANA     ^BANANA	BNN^AA   A

http://en.wikipedia.org/wiki/Burrows-Wheeler transform

# BWT is remarkable because it is reversible.

Any ideas as how you might reverse it?

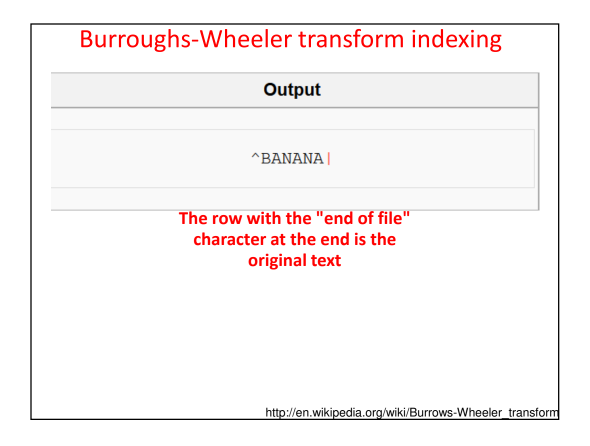


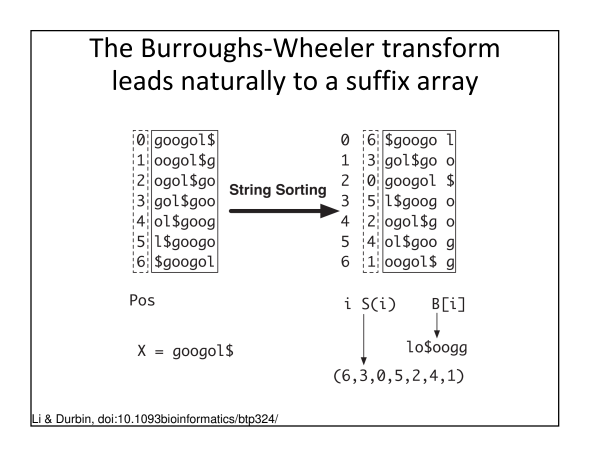
Add 1	Sort 1	Add 2	Sort 2
В	A	BA	AN
N	A	NA	AN
N	A	NA	A
^	В	^B	BA
A	N	AN	NA
A	N	AN	NZ
1	^	^	^E
A	1	A	1′
Write the	Sort it	Add the	Sort those

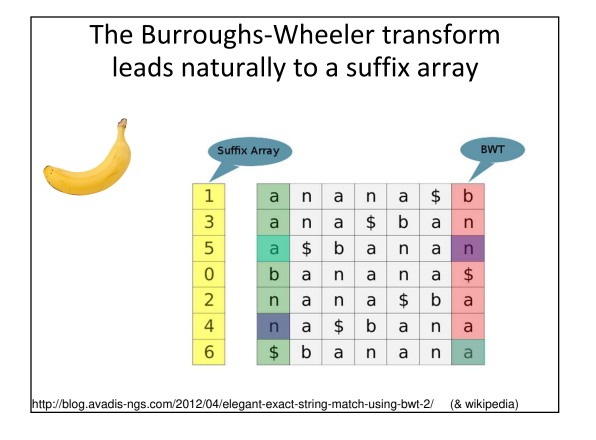
Add 3	Sort 3	Add 4	Sort 4
BAN	ANA	BANA	ANAN
NAN	ANA	NANA	ANA
NA	A ^	NA   ^	A ^E
^BA	BAN	^BAN	BANA
ANA	NAN	ANAN	NANA
ANA	NA	ANA	NA ^
^B	^BA	^BA	^BAN
A   ^	^B	A ^B	^BA
Add the	Sort those	Add the	Sort those

Add 5	Sort 5	Add 6	Sort 6
BANAN	ANANA	BANANA	ANANA
NANA	ANA   ^	NANA   ^	ANA ^I
NA ^B	A ^BA	NA ^BA	A ^BAN
^BANA	BANAN	^BANAN	BANANA
ANANA	NANA	ANANA	NANA   ′
ANA   ^	NA ^B	ANA ^B	NA ^BA
^BAN	^BANA	^BANA	^BANAI
A ^BA	^BAN	A ^BAN	^BANA
Add the	Sort those	Add the	Sort those

	Add 8	Sort 7	Add 7
	BANANA   ^	ANANA   ^	BANANA
The row with	NANA ^BA	ANA ^BA	NANA ^B
the "end of file	NA   ^BANA	A ^BANA	NA   ^BAN
<ul> <li>character at th</li> </ul>	^BANANA	BANANA	^BANANA
end is the	ANANA   ^B	NANA ^B	ANANA   ^
original text	ANA ^BAN	NA ^BAN	ANA ^BA
	^BANANA	^BANANA	^BANAN
	A   ^BANAN	^BANAN	A ^BANA
	Add the	Sort those	Add the columns



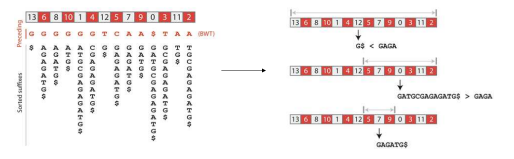




"If string W is a substring of X, the position of each occurrence of W in X will occur in an interval in the suffix array. This is because all the suffixes that have W as prefix are sorted together."

Li & Durbin, doi:10.1093bioinformatics/btp324/

e.g. applying BWT to construct the suffix array of GATGCGAGAGATG



The search can be even more efficient by using compression & various other extensions

http://blog.thegrandlocus.com/2016/07/a-tutorial-on-burrows-wheeler-indexing-methods

# Why is this efficient?

Searching a suffix array in this way cuts the search space in half at each step, so...

A suffix array of the human genome (3.2 billion bases) takes at most

$$log_2(3.2 billion) + 1 = 32 steps$$

to determine if a query sequence is present or not

There are few more steps to find all the occurrences, build an efficient real-world implementation, use compression to reduce memory and storage space, etc., but this still illustrates the massive savings in time and memory from constructing an index

