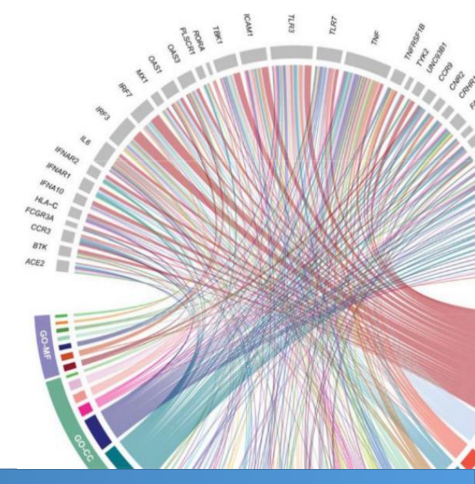
# Tècniques i Eines Bioinformàtiques

**Suffix Trees** 

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#### Acknowledgements

Many pictures and materials are taken from **Ben Langmead's course**.

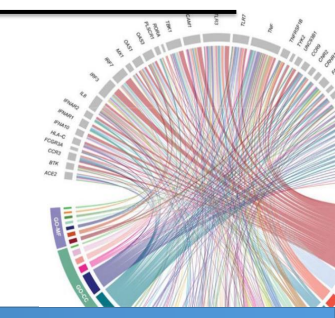
#### Course heavily inspired in:

- **Genome-Scale Algorithm Design**. Veli Mäkinen, Djamal Belazzougui, Fabio Cunial, Alexandru I. Tomescu. Cambridge University Press.
- Algorithms on Strings, Trees, and Sequences. Dan Gusfield.
   Cambridge University Press.
- An Introduction to Bioinformatics Algorithms. Neil C. Jones, Pavel A. Pevzner. MIT Press.



1

# **Suffix Trees**

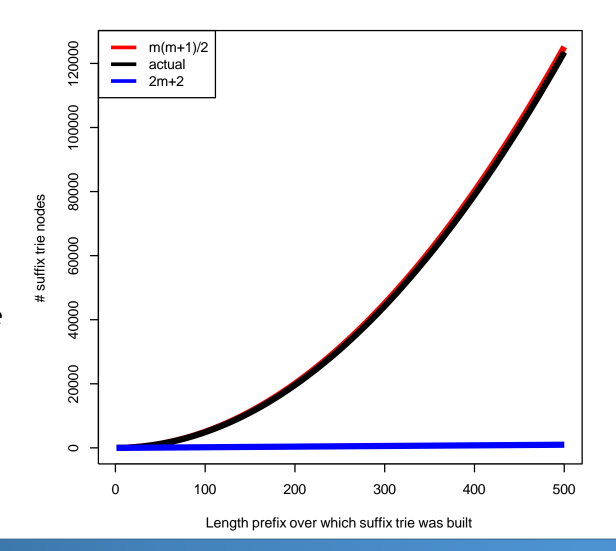


#### **Suffix Trie**

We saw the suffix trie, but we also saw its size grows quadratically with the length of the string

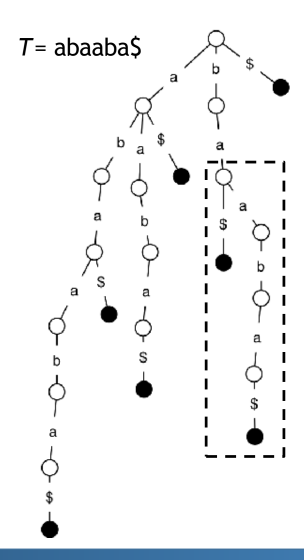
Human genome is 3 · 10<sup>9</sup> bases long.

If  $m = 3.10^9$ ,  $m^2$  is way huge, far beyond what we can store in memory

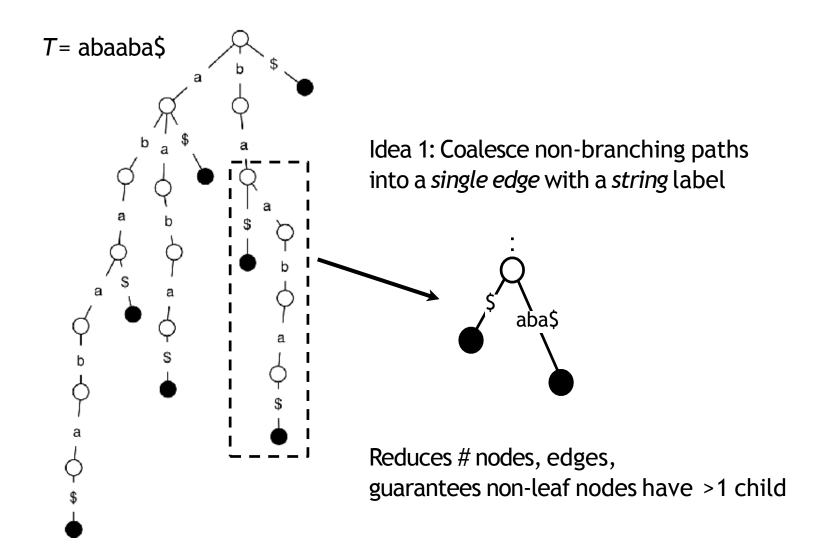




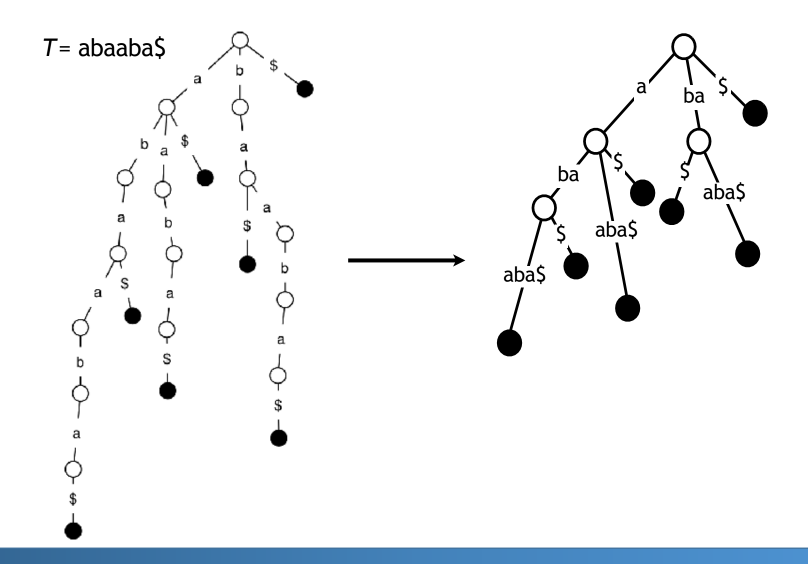
# Suffix trie: making it smaller



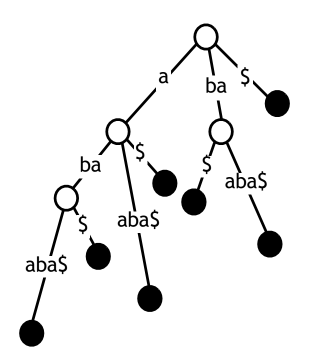
## Suffix trie: making it smaller



# Suffix trie: making it smaller



$$T$$
= abaaba\$  $|T|$ =  $m$ 



# leaves? m

# non-leaf nodes (bound)? ≤m-1

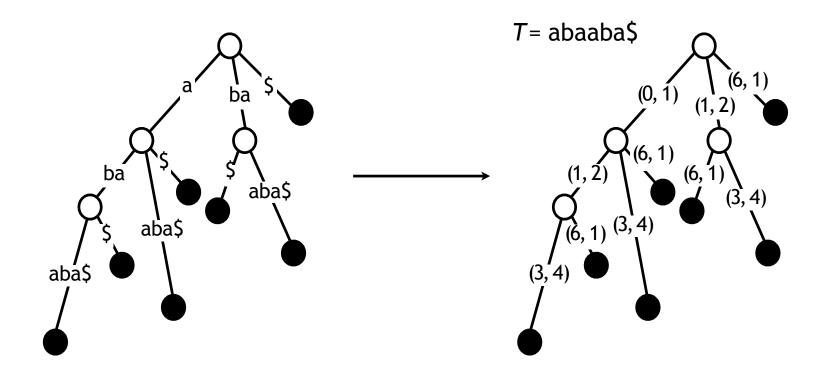
 $\leq 2m$  -1 nodes total -O(m)

Is total size O(m) now?

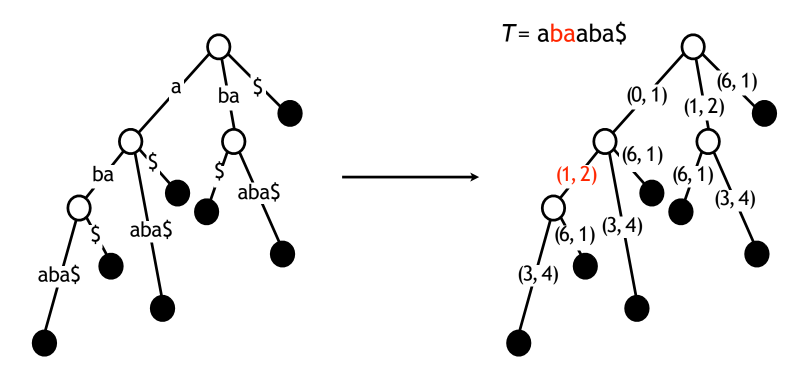
**No:** total length of edge labels grows with  $m^2$ 



Idea 2: Store *T* itself in addition to the tree. Convert tree's edge labels to (offset, length) pairs with respect to *T* 



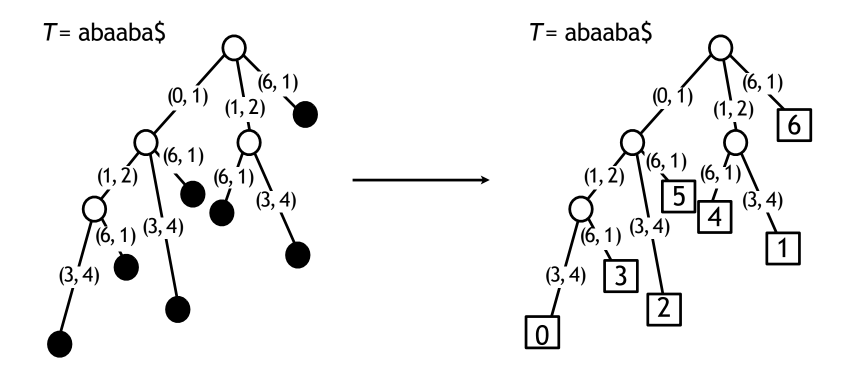
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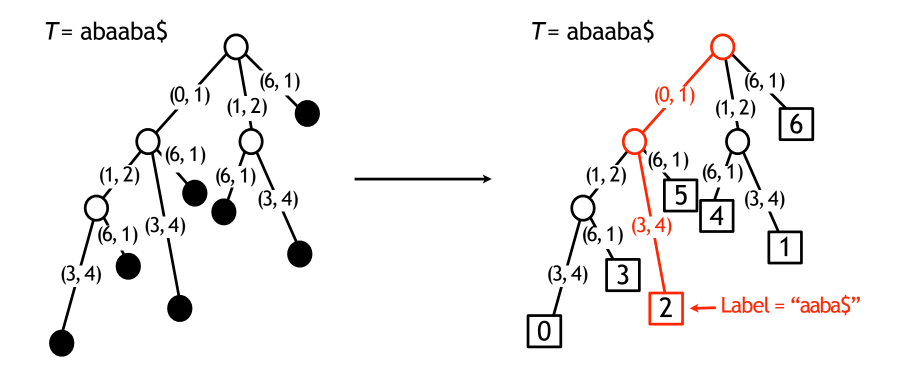
Space is now O(m) Suffix trie was  $O(m^2)$ !



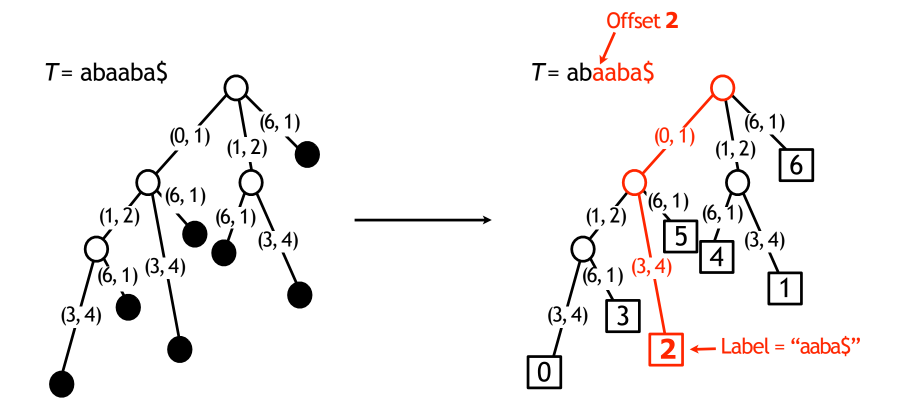
#### Suffix tree: leaves hold offsets



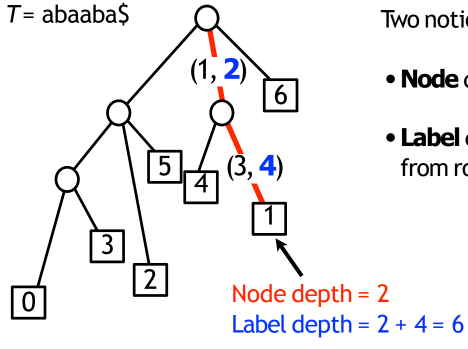
#### Suffix tree: leaves hold offsets



#### Suffix tree: leaves hold offsets



#### Suffix tree: labels



Two notions of depth:

- **Node** depth: # edges from root to node
- **Label** depth: total length of edge labels from root to node

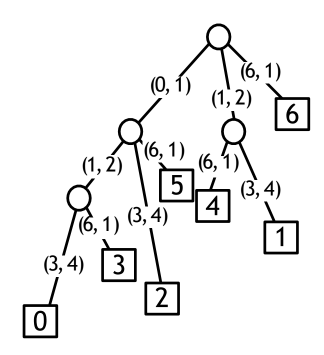
### Suffix tree: building

Method 1: build suffix trie, coalesce non- branching paths, relabel edges

 $O(m^2)$  time,  $O(m^2)$  space

Method 2: build single-edge tree representing longest suffix, augment to include the 2<sup>nd</sup>-longest, augment to include 3<sup>rd</sup>-longest, etc (Gusfield 5.4)

 $O(m^2)$  time, O(m) space



### Suffix tree: building

Canonical method: Ukkonen's algorithm

Ukkonen, Esko. "On-line construction of suffix trees."

Algorithmica 14.3 (1995): 249-260.

O(m) time and space!

Won't cover it in class; see Gusfield Ch. 6 for details



#### On-Line Construction of Suffix Trees1

E. Ukkonen<sup>2</sup>

Abstract. An on-line algorithm is presented for constructing the suffix tree for a given string in time linear in the length of the string. The new algorithm has the desirable property of processing the string symbol by symbol from left to right. It always has the suffix tree for the scanned part of the string ready. The method is developed as a linear-time version of a very simple algorithm for (quadratic size) suffix tries. Regardless of its quadratic worst case this latter algorithm can be a good practical method when the string is not too long. Another variation of this method is shown to give, in a natural way, the well-known algorithms for constructing suffix automata (DAWGs).

Key Words. Linear-time algorithm, Suffix tree, Suffix trie, Suffix automaton, DAWG.

Canonical algorithm for O(m) time & space suffix tree construction

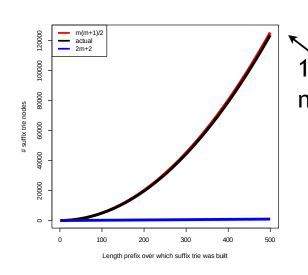


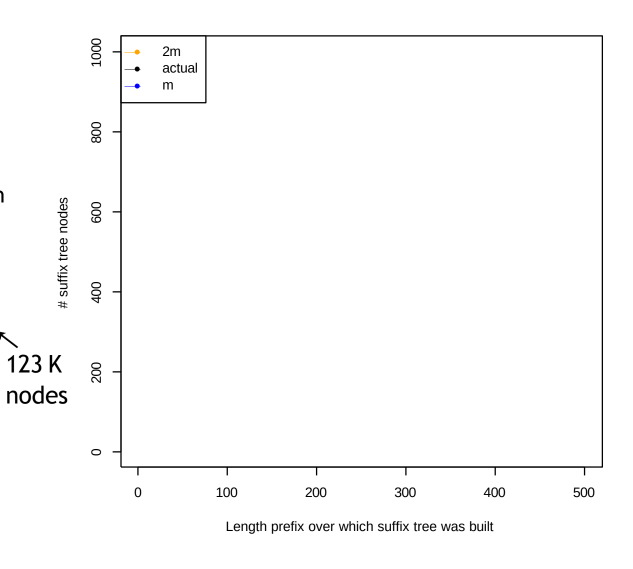
### Suffix tree: actual growth

Built suffix trees for the first 500 prefixes of the lambda phage virus genome

Black curve shows # nodes increasing with prefix length

#### Remember suffix trie plot:



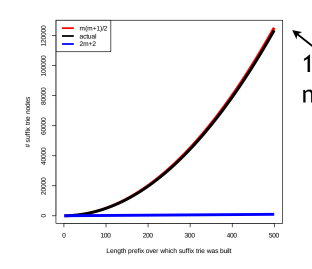


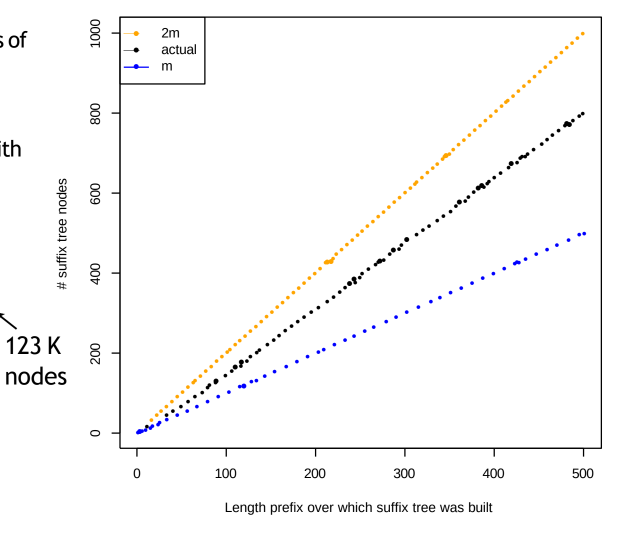
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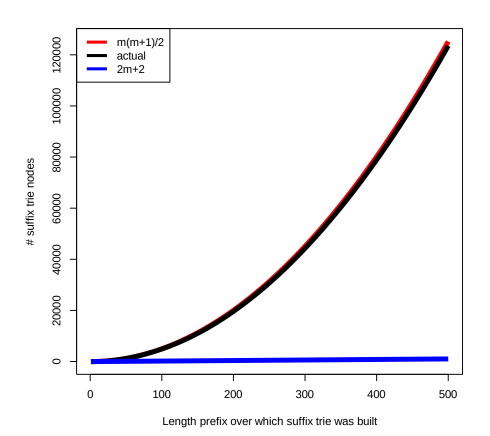
Remember suffix trie plot:





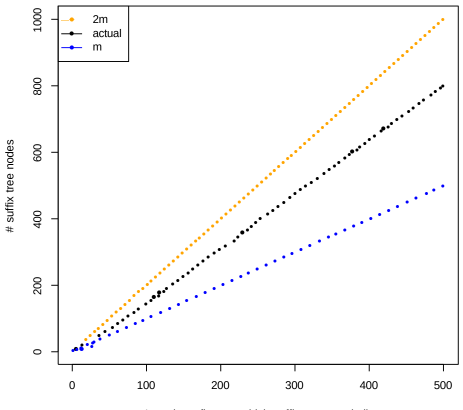


#### >100K nodes

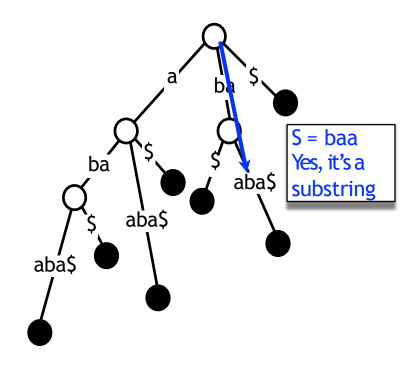


## Suffix tree

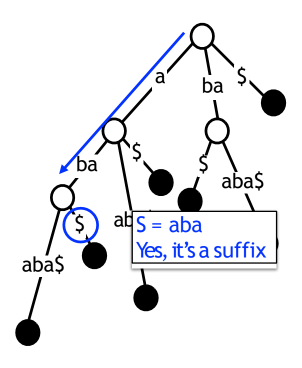
#### <1K nodes



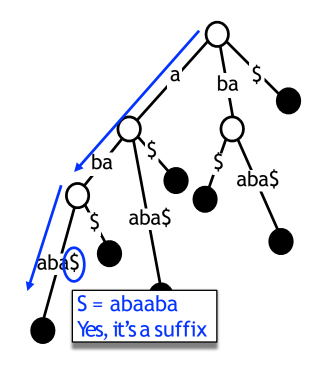
How do we check whether a string S is a substring of T?



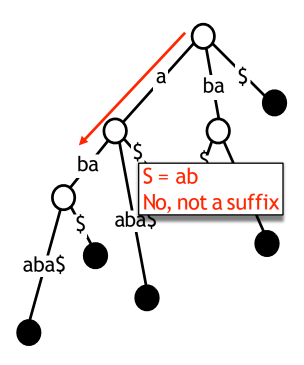
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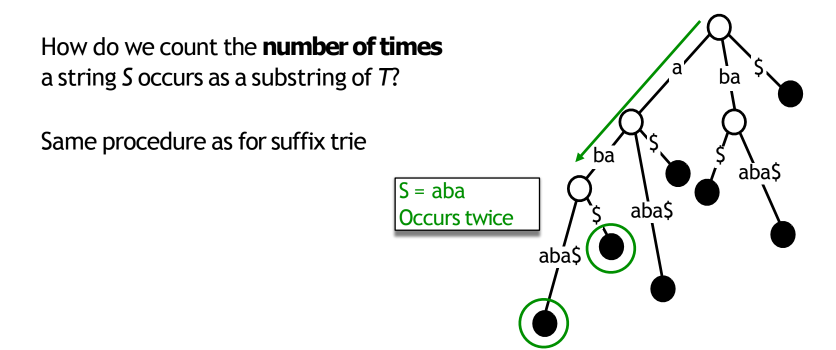


How do we check whether a string S is a suffix of T?



How do we check whether a string S is a suffix of T?





We can also **count or find** all the matches of P to T. Let k = # matches.

E.g., 
$$P = ab$$
,  $T = abaaba$ \$

O(n) Step 1: walk down ab path

If we "fall off" there are no matches

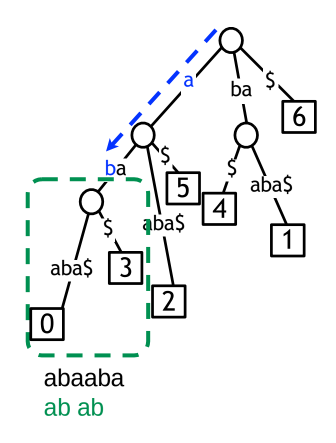
Step 2: visit all leaf nodes below

Report each leaf offset as match offset

# leaves in subtree is is k,

# non-leaves is  $\leq$  k-1

O(n + k) time overall



#### Suffix tree: some bounds

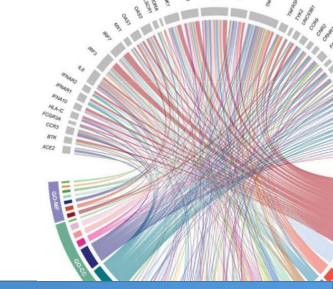
	<b>Suffix tree</b>
Time: Does Poccur?	<i>O</i> ( <i>n</i> )
Time: Count <i>k</i> occurrences of P	O(n + k)
Time: Report <i>k</i> locations of P	O(n + k)
Space	O(m)

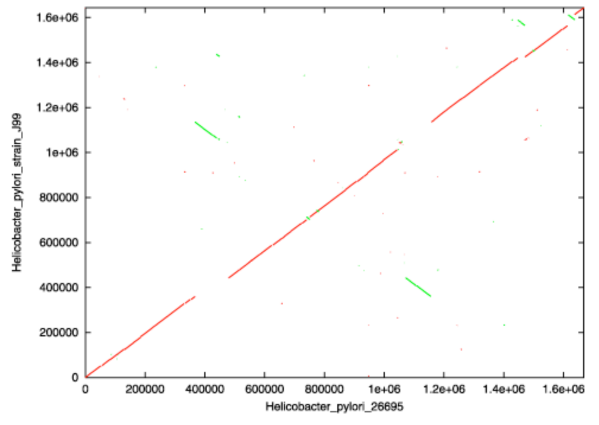
m = |T|, n = |P|, k = # occurrences of P in T



2

# **Longest Common Substring**





Dots are maximal unique matches (MUMs), a kind of long substring shared by two sequences

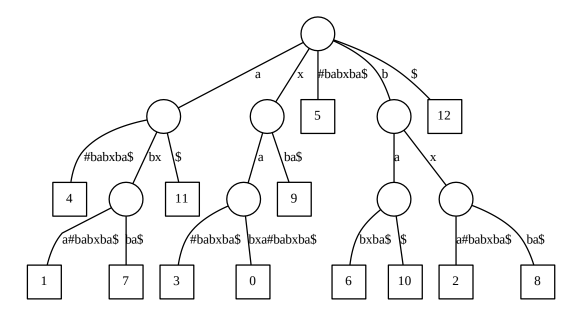
Red = match between like strands green = different strands

Axes are strains of Helicobacter pylori, bacterium found in stomach & associated with ulcers



Find longest common substring (LCS) of X and Y, make a new string X#Y\$ where #, \$ are both terminal symbols. Build a suffix tree for X#Y\$.

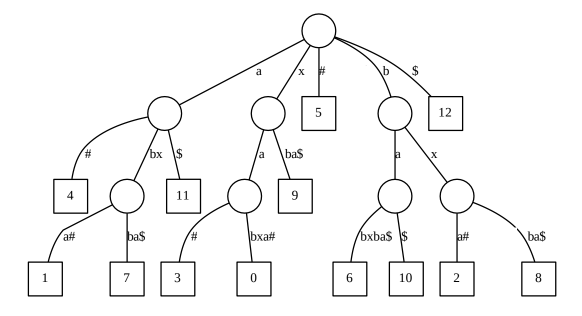
X= xabxa Y= babxba
X#Y\$= xabxa#babxba\$



For clarity, if a suffix includes part of both strings, let's hide the portion after the #

Find longest common substring (LCS) of X and Y, make a new string X#Y\$ where #, \$ are both terminal symbols. Build a suffix tree for X#Y\$.

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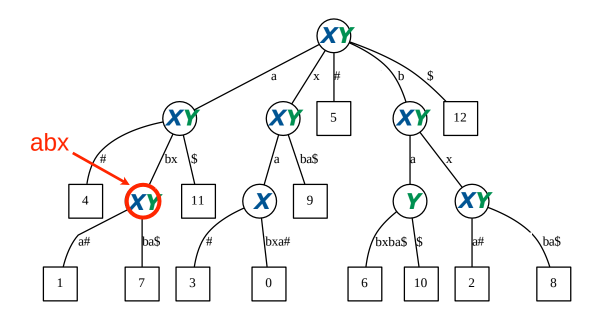


For clarity, if a suffix includes part of both strings, let's hide the portion after the #

Now suffixes of **X**end in # and suffixes of **Y**end in \$

Find longest common substring (LCS) of X and Y, make a new string X#Y\$ where #, \$ are both terminal symbols. Build a suffix tree for X#Y\$.

Leaves with labels in [0, 5] are suffixes of X#, labels of [6, 12] are suffixes of Y\$



Traverse tree, annotating each node according to whether leaves below include suffixes of X, Y or both

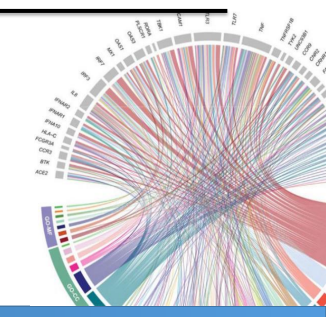
Node w/ greatest label depth annotated **XY** corresponds to LCS

O(|X| + |Y|) time and space!



3

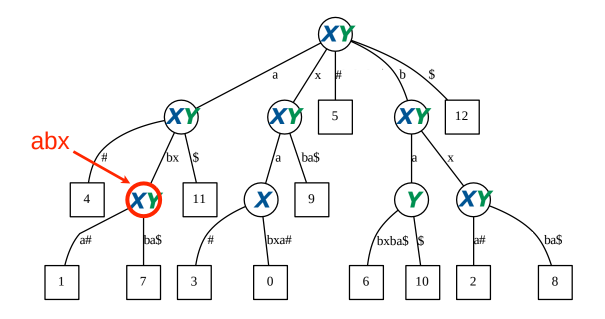
# **Other Applications**



#### **Generalized Suffix Tree**

It's often useful to build a suffix tree of many strings at once

This is a *generalized suffix tree*. See *Gusfield* 6.4.



#### Suffix trees in the real world

#### Alignment of whole genomes (MUMmer):

Delcher, Arthur L., et al. "Alignment of whole genomes." Nucleic Acids Research 27.11 (1999): 2369-2376.

Delcher, Arthur L., et al. "Fast algorithms for large-scale genome alignment and comparison." Nucleic Acids Research 30.11 (2002): 2478-2483.

Kurtz, Stefan, et al. "Versatile and open software for comparing large genomes." Genome Biol 5.2 (2004): R12.

~ 4,000 citations

http://mummer.sourceforge.net

#### Computing and visualizing repeats in whole genomes (REPuter):

Kurtz, Stefan, and Chris Schleiermacher. "REPuter: Fast computation of maximal repeats in complete genomes." *Bioinformatics* 15.5 (1999): 426-427.

Kurtz, Stefan, et al. "REPuter: the manifold applications of repeat analysis on a genomic scale." Nucleic acids research 29.22 (2001): 4633-4642.

> 1,000 citations

http://bibiserv.techfak.uni-bielefeld.de/reputer

#### Identifying sequence motifs

Marsan, Laurent, and Marie-France Sagot. "Algorithms for extracting structured motifs using a suffix tree with an application to promoter and regulatory site consensus identification." *Journal of Computational Biology* 7.3-4 (2000): 345-362.

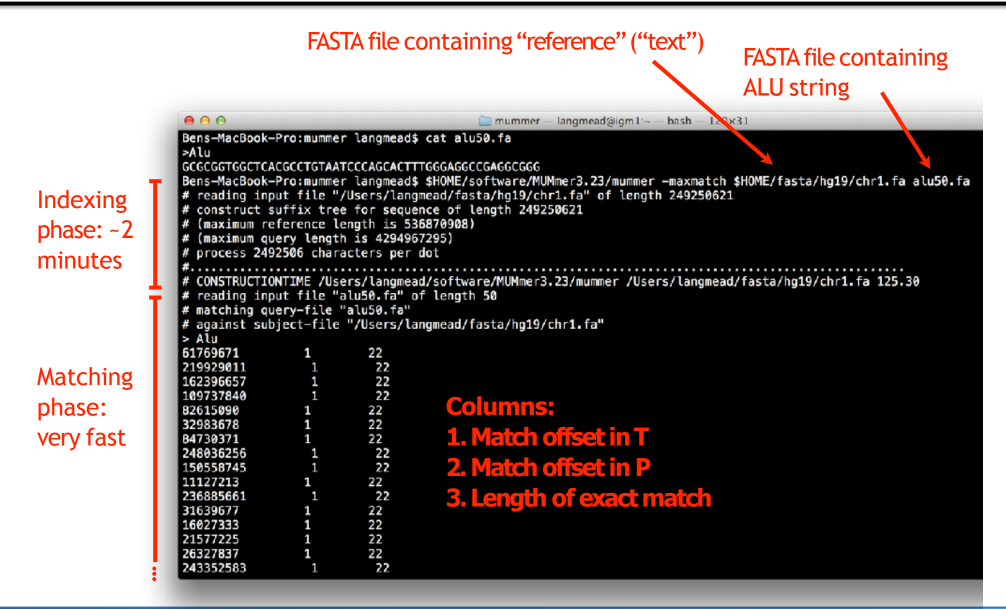
Sagot, Marie. "Spelling approximate repeated or common motifs using a suffix tree." LATIN'98: Theoretical Informatics (1998): 374-390.

~ 600 citations

#### Also used in: multiple alignment



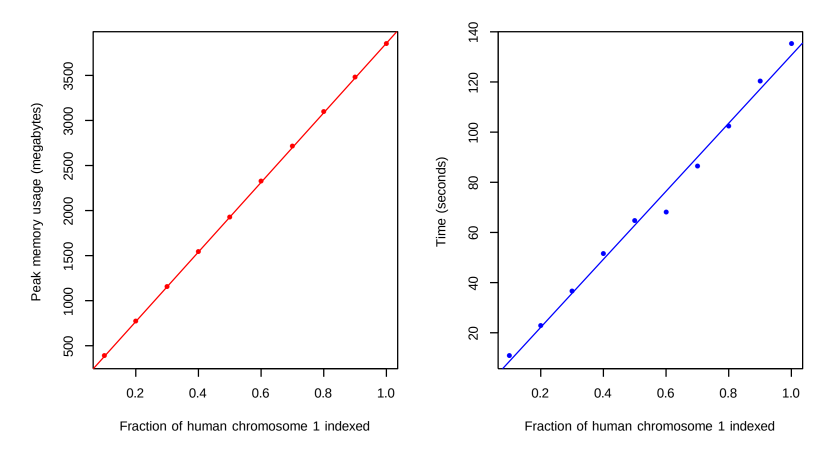
#### Suffix trees in the real world (MUMmer)





### Suffix trees in the real world (MUMmer)

MUMmer v3.32 time and memory scaling when indexing increasingly larger fractions of human chromosome 1



For whole chromosome 1, took 2m:14s and used 3.94 GB memory

#### Suffix trees in the real world: the constant factor

*O*(*m*) is desirable, but "constant factor" is significant, sometimes making the suffix tree inconvenient

Constant factor varies depending on implementation:

MUMmer constant factor = 3.94 GB / 250 million nt ≈ **15.76 bytes per nt** 

Kurtz, Stefan. "Reducing the space requirement of suffix trees." Software Practice and Experience 29.13 (1999): 1149-1171.

Suffix tree of human genome will be >45GB, perhaps much larger depending on exact data structures underlying suffix tree nodes/edges

# **Exercises**