

#### Lecture 16: Combinatorial Pattern Matching

Study Chapter 9.1 – 9.5

Fall 2013

# **Repeat Finding**

- Example of repeats:
  - AT<u>GGTC</u>TA<u>GGTC</u>CTAGT<u>GGTC</u>
- Motivation to find them:
  - Phenotypes arise from copy-number variations
  - Genomic rearrangements are often associated with repeats
  - Trace evolutionary secrets
  - Many tumors are characterized by an explosion of repeats



# **Repeat Finding**

- Near matches are more difficult:
  - AT<u>GGTC</u>TA<u>GGAC</u>CTAGT<u>GTTC</u>
- Motivation to find them:
  - Phenotypes arise from copy-number variations
  - Genomic rearrangements are often associated with repeats
  - Trace evolutionary secrets
  - Many tumors are characterized by an explosion of repeats



### *l*-mer Repeats

- Long repeats are difficult to find
- Short repeats are easy to find
- Strategy for finding long repeats:
  - Find exact repeats of short *l*-mers (*l* is usually 10 to 13)
  - Extend *l*-mer repeated seeds into longer, *maximal* repeats



#### $\ell$ -mer Repeats (cont'd)

 There are typically many locations where an *l*-mer is repeated:

#### GCTTACAGATTCAGTCTTACAGATGGT

• The 4-mer TTAC starts at locations 3 and 17



# Extending *l*-mer Repeats

#### GCTTACAGATTCAGTCTTACAGATGGT

Extend these 4-mer matches:

#### G<u>CTTACAGAT</u>TCAGT<u>CTTAC</u>AGATGGT

- Maximal repeat: CTTACAGAT
- Maximal repeats cannot be extended in either direction
- To find maximal repeats in this way, we need ALL start locations of all *l*-mers in the genome
- **Hashing** lets us find repeats quickly in this manner



# Hashing: What is it?

- How hashing works...
  - Generate an integer "key" from an arbitrary record
  - Store record in an data structure indexed by this integer key
- Hashing is a very efficient way to store and retrieve data
  - e.g., Python directories are hashes



# Hashing: Definitions

- Hash table: array used in hashing
- <u>Records</u>: data stored in a *hash table*
- <u>Keys</u>: identify sets of *records*
- <u>Hash function</u>: uses a *key* to generate an index to insert at in *hash table*
- <u>Collision</u>: when more than one record is mapped to the same index in the hash table



# Hashing: Example

- Where do the animals eat?
- Records: each animal
- Keys: where each animal eats



Records	Keys
x	h(x)
Penguin	1
Octopus	4
Turtle	3
Mouse	2
Snake	3
Heron	1
Tiger	2
Iguana	3
Ăpe	2
Cricket	4
Sparrow	1

## Hashing DNA sequences

- Each *l*-mer can be translated into a binary string, key = quaternary(seq)
  (A, T, C, G can be represented as 0, 1, 2, 3)
- After assigning a unique integer per *l*-mer it is easy to store the starting locations of occurance of each *l*-mer in a genome of length *n* in O(*l n*) time



# Hashing: Maximal Repeats

- To find repeats in a genome:
  - For all *l*-mers in the genome, note its starting position and the sequence
  - Generate a hash table index for each unique *l*-mer sequence
  - In each index of the hash table, store all genome start locations of the *l*-mer which generated that index
  - Extend *l*-mer repeats to maximal repeats
- Problem as *l* gets big the number of possible patterns becomes larger than the genome's length (4<sup>*l*</sup> >> n)

# Hashing: Collisions

Generate hash keys *l*-mer #1 20 400 450 100from a reduced space *l*-mer #2 Ex. Key = quaternary(seq) % (N/*l*) *l*-mer #3 • Leads to possible collisions 1003 2003 503 43 Dealing with collisions: – "Chain" tuples of (*l*-mer, start location) Chained Locations of *l*-mers pairs in a linked list *l*-mer #n

# Hashing: Summary

- When finding genomic repeats from *l*-mers:
  - Generate a hash table index for each *l*-mer sequence
  - In each index, store all genome start locations of the *l*-mer which generated that index
  - Extend *l*-mer repeats to maximal repeats



### Pattern Matching

- What if, instead of finding repeats in a genome, we want to find all positions of a particular sequences in given sequence?
- This leads us to a different problem, the *Pattern Matching Problem*



## Pattern Matching Problem

- <u>Goal</u>: Find all occurrences of a pattern in a text
- <u>Input</u>: Pattern  $p = p_1 \dots p_n$  and text  $t = t_1 \dots t_m$
- <u>Output</u>: All positions 1≤ *i* ≤ (*m n* + 1) such that the *n*-letter substring of *t* starting at *i* matches *p*
- Motivation: Searching database for a known pattern



#### Exact Pattern Matching: A Brute-Force Algorithm

PatternMatching(p,t) $1 n \leftarrow$  length of pattern p $2 m \leftarrow$  length of text t $3 \text{ for } i \leftarrow 1 \text{ to } (m - n + 1)$  $4 \text{ if } t_{i} \dots t_{i+n-1} = p$ 5 output i



#### Exact Pattern Matching: An Example

- *PatternMatching* algorithm for:
  - Pattern GCAT
  - Text CGCATC

CGCATC CCAT CCATC CGCATC CGCATC





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#### Exact Pattern Matching: Running Time

- *PatternMatching* runtime: O(*nm*)
- Probability-wise, it's more like O(*m*)
  - Rarely will there be close to *n* comparisons in line 4
- Worse case: Find "AAAAT" in "AAAAAAAAAAAAAAA
- Better solution: suffix trees
  - Can solve problem in O(m) time

Comp 555

– Conceptually related to keyword trees

#### Keyword Trees: Example

root

p

- *Keyword tree*:
  - Apple

- *Keyword tree*:
  - Apple
  - Apropos



root





- Keyword tree:
  - Apple
  - Apropos
  - Banana





- Keyword tree:
  - Apple
  - Apropos
  - Banana
  - Bandana





- Keyword tree:
  - Apple
  - Apropos
  - Banana
  - Bandana
  - Orange





### Keyword Trees: Properties

- Stores a set of keywords in a rooted labeled tree
- Each edge labeled with a letter from an alphabet
- Any two edges coming out of the same vertex have distinct labels
- Every keyword stored can be spelled on a path from root to some leaf
- Searches are performed by "threading" the target pattern through the tree



- Thread "appeal"
  - <u>a</u>ppeal





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- Thread "appeal"
  - <u>ap</u>peal





- Thread "appeal"
  - <u>app</u>eal





- Thread "appeal"
  - <u>app</u>eal





- Thread "apple"
  - <u>a</u>pple





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- Thread "apple"
  - <u>ap</u>ple



- Thread "apple"
  - <u>app</u>le





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- Thread "apple"
  - <u>appl</u>e



• Thread "apple"

– <u>apple</u>

Now thread "band", "or", and the nonsense word "apro"

How do you tell "real" words from nonsense? (Homework problem)



#### Multiple Pattern Matching Problem

- <u>Goal</u>: Given a set of patterns and a text, find all occurrences of any of patterns in text
- <u>Input</u>: *k* patterns  $\mathbf{p}^1, \dots, \mathbf{p}^k$ , and text  $\mathbf{t} = t_1 \dots t_m$
- <u>Output</u>: Positions 1 ≤ *i* ≤ *m* where substring of **t** starting at *i* matches **p**<sub>*i*</sub> for 1 ≤ *j* ≤ *k*
- Motivation: Searching database for known multiple patterns



#### Multiple Pattern Matching: Straightforward Approach

- Can solve as *k* "Pattern Matching Problems"
  - Runtime:

#### O(kmn)

using the *PatternMatching* algorithm *k* times

- -m length of the text
- -n average length of the pattern



#### Multiple Pattern Matching: Keyword Tree Approach

- Or, we could use keyword trees:
  - Build keyword tree in O(N) time; N is total length of all patterns
  - With naive threading: O(N + nm)
  - Aho-Corasick algorithm: O(N + m)



# Keyword Trees: Threading

- To match patterns in a text using a keyword tree:
  - Build keyword tree of patterns
  - "Thread" the text through the keyword tree

t = "mr and mrs durster of number 4 privetdrive were proud to say that they were perfectlynormal thank you very much"





## Keyword Trees: Threading

- Threading is "complete" when we reach a leaf in the keyword tree
- When threading is "complete," we've found a pattern in the text

t = " mr and mrs dursley of number 4 p rivetdrive were proud to say that they were perfectlynormal thank you very much"



#### Suffix Trees=Collapsed Keyword Trees

- All suffixes of a given sequence
- Similar to keyword trees, except vertices of degree 2 are removed and the strings on either side are merged
  - Each edge is labeled
     with a *substring* of a text
  - All internal vertices have at least three edges
  - Terminal vertices, leaves, are labeled by the index of the pattern.



#### Suffix Tree of a Text

- Construct a keyword tree from all suffixes of a text
- Collapse non-branching paths into an edge (path compression)



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## Suffix Trees: Advantages

- With careful bookkeeping a test's suffix tree can be constructed in a single pass of the text
- Thus, suffix trees can be built faster than keyword trees of suffixes and transforming them



### Suffix Tree Construction

- Few books, including ours, delve into the details of suffix tree construction algorithms due to its reputation for being overly complicated.
- Weiner's and McCreight's original linear algorithms for constructing a suffix trees had some disadvantages.
- Principle among them was the requirement that the tree be built in reverse order, meaning that the tree was grown incrementally by adding characters from the end of the input.
- This ruled it out for on-line processing



## Ukkonen's Clever Bookkeeping

- Esko Ukkonen's construction works from left to right.
- It's incremental. Each step transforms the Suffix Tree of the prefix ending at the i<sup>th</sup> character to the Suffix Tree ending at the i+1<sup>th</sup>.



## Tree Properties

- Extensions are done by threading each new prefix through the tree and visiting each of the suffixes of the current tree.
- At each step we start at the longest suffix (BOOK), and work our way down to the shortest (empty string)
- Each ends at a node of three types:
  - A leaf node (1,2,4,5)
  - An explicit node (0, 3)
  - An implicit node (Between characters of a substring labeling an edge, such as BO, BOO, and OO).



#### Observations

- There are 5 suffixes in the tree (including the empty string) after adding BOOK
- They are represented by the root and 4 leaves
- Adding the next letter, another 'K', requires visiting each of the suffixes in the existing tree, in order of decreasing length, and adding letter 'K' to its end.
- Adding a character to a leaf node never creates a new explicit node, regardless of the letter
- If the root already has an edge labeled 'K' we just extend it



#### Split and Add Update

- The next step is to add an 'E' to our tree
- As before, add 'E' to each suffix in order of decreasing lengths BOOKK, OOKK, OKK, KK, K
- The first suffix that does not terminate at a leaf is called the "active point" of the suffix tree



# Updating an Explicit Node

• After updating suffix K, we still have to update the next shorter suffix, which is the empty string.





## Generalizing

- Once a leaf node, always a leaf node
- Additional characters only extends the edge leading to the leaf (leaves are easy)
- When adding a new leaf, its edge will represent all characters from the i<sup>th</sup> suffix's starting point to the i+1<sup>st</sup> text's end. Because of this once a leaf is created, we can just forget about it. If the edge is later split, its start may change but it will extend to the end.
- This means that we only need to keep track of the active point in each tree, and update from there.



#### One Last Detail

- The algorithm sketch so far glosses over one detail. At each step of an update we need to keep track of the next smaller suffix from the i<sup>th</sup> update
- To do this a suffix pointer is kept at each internal node
- For Pseudo code
  - Mark Nelson, "Fast String Searching with Suffix Trees"
     <u>Dr. Dobb's Journal</u> August, 1996
- For proofs of linear space/time performance
  - E. Ukkonen. <u>"On-line</u> <u>construction of suffix trees</u>. Algorithmica, 14(3):249-260, September 1995.



The suffix tree for ABABABC with suffix pointers shown as dashed lines

### Use of Suffix Trees

- Suffix trees hold all suffixes of a text, T
  - i.e., ATCGC: ATCGC, TCGC, CGC, GC, C
  - Builds in O(m) time for text of length m
- To find any pattern *P* in a text:
  - Build suffix tree for text, O(m), m = |T|
  - Thread the pattern through the suffix tree
  - Can find pattern in O(n) time! (n = |P|)
- O(|T| + |P|) time for "Pattern Matching Problem" (better than Naïve O(|P||T|)
  - Build suffix tree and lookup pattern
- Multiple Pattern Matching in O(|T| + k |P|)

#### Pattern Matching with Suffix Trees

#### SuffixTreePatternMatching(p,t)

- 1 Build **suffix tree** for text **t**
- 2 Thread pattern **p** through **suffix tree**
- **3** if threading is complete
- 4 traverse all paths from the threading's endpoint to leaves and **output** their positions
- 5 else
- 6 **output** "Pattern does not appear in text"

#### Suffix Trees: Example



#### Multiple Pattern Matching: Summary

- Keyword and suffix trees are useful data structures supporting various pattern finding problems
- *Keyword trees*:
  - Build keyword tree of patterns, and *thread text* through it
- Suffix trees:
  - Build suffix tree of text, and thread patterns through it



#### Suffix Trees: Theory vs. Practice

- In concept, suffix trees are extremely powerful for making a variety of queries concerning a sequence
  - What is the shortest unique substring?
  - How many times does a given string appear in a text?
- Despite the existence of linear-time construction algorithms, and O(m) search times, suffix trees are still rarely used for genome scale searching
  - Large storage overhead
- Close cousins of the Suffix-Tree (Suffix Arrays and Burrows-Wheeler Transforms) are more common
- Next lecture

