CSCI 102 assignment 6 – BSTs for counting k-mers

Oct 27

In this assignment you will be implementing a binary search tree (BST) to count the occurrences of strings of length k in DNA. As an example, we may want to take the sequence AGGAGAGG, break it into sequences of length k=3 – in biology we call these 3-mers – AGG, GGA, GAG, AGA, GAG, AGG, and count the occurrences of these strings in a map {AGG:2, GGA:1, GAG:2, AGA:1}. A classical option for implementing such a map is a hash table. Hopwever, sometimes we want k to be large and the DNA sequences we analyze can get very large; in these cases we can expect a large number of collisions, making hash tables inefficient. Instead, we can use BST to store entries with keys that are k-mers and values that are integers. Here you will implement a BST to store counts of k-mers in long DNA sequences.

To build a BST with keys that are Strings we will need a way to decide if one string is "greater or less than" another. To compare strings as such, we decide strings that are alphabetically later are "greater than". If you have two Strings, string1 and string2, Java allows you to compare them alphabetically by writing string1.compareTo(string2).

- Define a class KmerCounter which implements Map<String, Integer>. It will have private class Node. Node will have attributes Integer count, Node parent, Node left, Node right, String kmer with get and set methods for each. You can think of the node as storing an entry with key kmer and value count. KmerCounter will have two private attributes, the size int size and the root of the BST Node root. Feel free to reuse our BST code from class.
- Create a method void printPositions that prints the k-mers and their counts at each node by performing an in-order traversal.
- Create a method int get(String kmer) that returns the count corresponding to a k-mer stored in the tree (return 0 if the k-mer is not in the tree).
- Create a method int addCount(String kmer) that adds one to the count of the count of kmer. Use the regular BST rules (no need for AVL). If kmer is not in the BST create a new node for storing its count.
- Create a method int remove(String kmer) that removes and returns the count of kmer. If kmer is not in the BST return 0.
- Create a main method where you take the DNA sequence of human insulin below and build a KmerCounter counting all of its 4-mers; do this by looping through each position of the sequence and adding 1 to the count of the 4-mer at that position with the method addCount. Remove the 4-mer TGAG from your tree and run printPositions.

Insulin DNA sequence:

TGTTTCCATGCTCTGTGTACGTGCCGGACGACTGGGAGGTGTCTCGAGAGAAGATCACCCTCCTTCGAGAGC
TGGGGCAGGGCTCCTTCGGCATGGTGTATGAGGGCAATGCCAGGGACATCATCAAGGGTGAGGCAGAGACCC
GCGTGGCGGTGAAGACGGTCAACGAGTCAGCCAGTCTCCGAGAGCCGATTGAGTTCCTCAATGAGGCCTCGG
TCATGAAGGGCTTCACCTGCCATCACGTGGTGCGCCTCCTGGGAGTGGTGTCCCAAGGGCCAGCCCACGCTGG
TGGTGATGGAGCTGATGGCTCACGGAGACCTGAAGAGCTACCTCCGTTCTCTGCGGCCAGAGGCTGAG

Please submit your code and answers to the questions in a zipped folder on Brightspace by Nov 5.