## Problem Set 2

This problem set is due October 11 at 11:59PM.

Solutions should be turned in through the course website in PS or PDF form using LaTeX. The course website has links to a number of editors that are useful for writing in LaTeX. It is recommended that you download the LaTeX solution template for this problem set which includes placeholders for solutions.

Exercises are for extra practice and should not be turned in. **Exercises:** 

- 1. CLRS 11.2-1
- 2. CLRS 11.2-2
- 3. CLRS 11.3-1
- 4. CLRS 11.3-3
- 1. Rotating Binary Search Trees

In this problem we'll explore the rotation operation on binary search trees. As discussed in class, this operation changes the structure of a binary search tree without affecting the inorder of the underlying nodes. See CLRS section 13.2 (page 277) for a description of the operation.

- (a) (6 points) Let l(v) denote the number of nodes in v's left subtree. Let L(T) be the sum of l(v) over all nodes of the tree T. Show that a right rotation decreases L(T). Deduce that it is impossible to do more than  $O(n^2)$  consecutive right rotations in an n node tree, i.e., with NO left rotations mixed in.
- (b) (6 points) Show that on an *n* node left path (a tree where all children are left children) it is possible to do  $\Omega(n^2)$  consecutive right rotations. (Again, no left rotations allowed.)
- 2. How Good is this Hash?

The following hash functions are used to hash strings of characters. Describe at least one strength and one weakness of each of the hash functions. (Don't try to do a rigorous mathematical analysis of the properties of the functions.)

(a) (6 points) Direct addressing, using the first two bytes of the hash key as an address into a 65,536-entry hashtable. (Assume all keys are at least two bytes long.)

- (b) (6 points) Add the values of all bytes in the hash key into a single-byte counter, ignoring overflow. Use the result as an index into a 256-element hash table.
- (c) (6 points) Start with a fixed prime number constant  $p_0$  and a second small number s. Set  $p := p_0$ . For each byte  $b_i$  in the hash key, set  $p := p + (p << s) + b_i$ , where p << s means "shift p left by s bit positions". Mod the final value of p by a prime number to produce the hash code. (Just describe what could be good or bad about this scheme depending on the values of  $p_0$  and s.)
- 3. Longest Common Substring

Humans have 23 pairs of chromosomes, while other primates like chimpanzees have 24 pairs. Biologists claim that human chromosome #2 is a fusion of two primate chromosomes that they call 2a and 2b. We wish to verify this claim by locating long nucleotide chains shared between the human and primate chromosomes.

We define the *longest common substring* of two strings to be the longest contiguous string that is a substring of both strings e.g. the longest common substring of DEADBEEF and EA7BEEF is BEEF. <sup>1</sup> If there is a tie for longest common substring, we just want to find one of them.

(a) (2 points) Ben Bitdiddle wrote a python program to find the longest common substring of two strings. What is the asymptotic running time of his code? Assume |s| = |t| = n.

- (b) (2 points) Describe a simple algorithm that finds the longest common substring in  $O(n^3)$  time.
- (c) (2 points) Describe a simple algorithm that finds the longest common substring in  $O(n^2 \log n)$  time.

<sup>&</sup>lt;sup>1</sup>Do not confuse this with the *longest common subsequence*, in which the characters do not need to be contiguous. The longest common subsequence of DEADBEEF and EA7BEEF is EABEEF.

- (d) (12 points) Describe an algorithm that finds the longest common substring in  $O(n \log n)$  time. If you use Rabin-Karp hashing, be sure to pick a specific hash function, describe why it works well for this problem, and discuss how you will handle collisions.
- (e) (**12 points**) Implement your algorithm from part (d). Be sure to thoroughly comment your code so the course staff can read it. Some simple test cases are available for download on the class website.

The human chromosome 2 and the chimp chromosomes 2a and 2b are quite large (over 100,000,000 nucleotides each) so we took the first and last million nucleotides of each chromosome and put them in separate files. These files are available on the class website, and also in /mit/6.006/dna/

chr2\_first\_1000000 contains the first million nucleotides of human chromosome 2, and chr2a\_first\_1000000 contains the first million nucleotides of chimpanzee chromosome 2a. Note: these files contain both uppercase and lowercase letters that are used by biologists to distinguish between parts of the chromosomes called introns and extrons.

- i. How long is the longest common substring of chr2\_first\_1000000 and chr2a\_first\_1000000?
- ii. How long is the longest common substring of chr2\_first\_1000000 and chr2b\_first\_1000000?
- iii. How long is the longest common substring of chr2\_last\_1000000 and chr2a\_last\_1000000?
- iv. How long is the longest common substring of chr2\_last\_1000000 and chr2b\_last\_1000000?

When you are finished, submit your code through the class website.

- (f) **Optional:** Make your algorithm run in  $O(n \log k)$  time, where k is the length of the longest common substring.
- (g) **Optional:** Run your algorithm on the entire chromosomes. They are available in /mit/6.006/dna in compressed form.