# Lecture 5: Scheduling and Binary Search Trees

## Lecture Overview

- Runway reservation system
  - Definition
  - How to solve with lists
- Binary Search Trees
  - Operations

## Readings

CLRS Chapter 10, 12.1-3

## Runway Reservation System

- Airport with single (very busy) runway (Boston  $6 \rightarrow 1$ )
- "Reservations" for future landings
- When plane lands, it is removed from set of pending events
- Reserve req specify "requested landing time" t
- Add t to the set if no other landings are scheduled within k minutes either way. Assume that k can vary.
  - else error, don't schedule

## Example



Figure 1: Runway Reservation System Example

Let R denote the reserved landing times: R = (41, 46, 49, 56) and k = 3

```
Request for time: 44 not allowed (46 \in R) 53 OK 20 not allowed (already past) \mid R \mid = n
```

Goal: Run this system efficiently in  $O(\lg n)$  time

#### Algorithm

Keep R as a sorted list.

```
init: R = []
req(t): if t < now: return "error"
for i in range (len(R)):
    if abs(t-R[i]) < k: return "error"
R.append(t)
R = sorted(R)
land: t = R[0]
if (t != now) return error
R = R[1: ] (drop R[0] from R)</pre>
```

#### Can we do better?

- Sorted list: Appending and sorting takes  $\Theta(n \lg n)$  time. However, it is possible to insert new time/plane rather than append and sort but insertion takes  $\Theta(n)$  time. A k minute check can be done in O(1) once the insertion point is found.
- Sorted array: It is possible to do binary search to find place to insert in  $O(\lg n)$  time. Using binary search, we find the smallest i such that  $R[i] \geq t$ , i.e., the next larger element. We then compare R[i] and R[i-1] against t. Actual insertion however requires shifting elements which requires  $\Theta(n)$  time.
- Unsorted list/array: k minute check takes O(n) time.
- Min-Heap: It is possible to insert in  $O(\lg n)$  time. However, the k minute check will require O(n) time.
- Dictionary or Python Set: Insertion is O(1) time. k minute check takes  $\Omega(n)$  time

What if times are in whole minutes?

Large array indexed by time does the trick. This will not work for arbitrary precision time or verifying width slots for landing.

**Key Lesson**: Need fast insertion into sorted list.

# Binary Search Trees (BST)

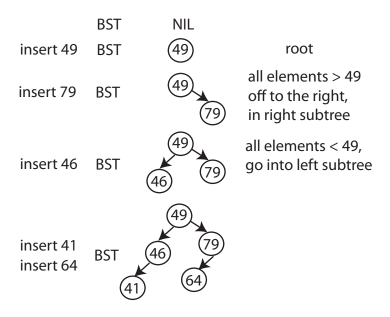


Figure 2: Binary Search Tree

## Properties

Each node x in the binary tree has a key key(x). Nodes other than the root have a parent p(x). Nodes may have a left child left(x) and/or a right child right(x). These are pointers unlike in a heap.

The invariant is: for any node x, for all nodes y in the left subtree of x,  $key(y) \le key(x)$ . For all nodes y in the right subtree of x  $key(y) \ge key(x)$ .

## Insertion: insert(val)

Follow left and right pointers till you find the position (or see the value), as illustrated in Figure 2. We can do the "within k=3" check for runway reservation during insertion. If you see on the path from the root an element that is within k=3 of what you are inserting, then you interrupt the procedure, and do not insert.

#### Finding a value in the BST if it exists: find(val)

Follow left and right pointers until you find it or hit NIL.

## Finding the minimum element in a BST: findmin()

Key is to just go left till you cannot go left anymore.

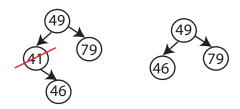


Figure 3: Delete-Min: finds minimum and eliminates it

## Complexity

All operations are O(h) where h is height of the BST.

Finding the next larger element: next-larger(x)

```
Note that x is a node in the BST, not a value.
next-larger(x)

if right child not NIL, return minimum(right)
else y = parent(x)
while y not NIL and x = right(y)
    x = y; y = parent(y)
return(y);
```

See Fig. 4 for an example. What would next-larger(find(46)) return?

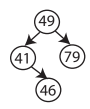


Figure 4: next-larger(x)

## New Requirement

Rank(t): How many planes are scheduled to land at times  $\leq t$ ? The new requirement necessitates a design amendment.

Cannot solve it efficiently with what we have but can augment the BST structure.

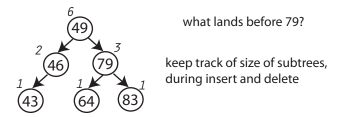


Figure 5: Augmenting the BST Structure

Summarizing from Fig. 5, the algorithm for augmentation is as follows:

- 1. Walk down tree to find desired time
- 2. Add in nodes that are smaller
- 3. Add in subtree sizes to the left

In total, this takes O(h) time.

subtree
$$49$$
 $46$ 
 $1 + 2 + 1 + 1 = 5$ 
 $79$ 
 $64$ 
subtree

Figure 6: Augmentation Algorithm Example

All the Python code for the Binary Search Trees discussed here are available at this link.

## Have we accomplished anything?

Height h of the tree should be  $O(\lg n)$ .

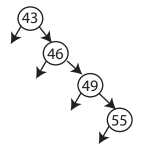


Figure 7: Insert into BST in sorted order

The tree in Fig. 7 looks like a linked list. We have achieved O(n) not  $O(\lg n)!!$ 

Balanced BSTs to the rescue in the next lecture!

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## **Problem Set 4**

## Both theory and programming questions are due Friday, 14 October at 11:59PM.

Remember that for the written response question, your goal is to communicate. Full credit will be given only to a correct solution which is described clearly. Convoluted and obtuse descriptions might receive low marks, even when they are correct. Also, aim for concise solutions, as it will save you time spent on write-ups, and also help you conceptualize the key idea of the problem.

We will provide the solutions to the problem set 10 hours after the problem set is due, which you will use to find any errors in the proof that you submitted. You will need to submit a critique of your solutions by **Thursday, October 20th, 11:59PM**. Your grade will be based on both your solutions and your critique of the solutions.

## **Problem 4-1.** [35 points] **Hash Functions and Load**

- (a) Imagine that an algorithm requires us to hash strings containing English phrases. Knowing that strings are stored as sequences of characters, Alyssa P. Hacker decides to simply use the sum of those character values (modulo the size of her hash table) as the string's hash. Will the performance of her implementation match the expected value shown in lecture?
  - 1. Yes, the sum operation will space strings out nicely by length.
  - 2. Yes, the sum operation will space strings out nicely by the characters they contain.
  - 3. No, because reordering the words in a string will not produce a different hash.
  - 4. No, because the independence condition of the simple uniform hashing assumption is violated.

**Solution:** No, because reordering the words in a string will not produce a different hash.

- (b) Alyssa decides to implement both collision resolution and dynamic resizing for her hash table. However, she doesn't want to do more work than necessary, so she wonders if she needs both to maintain the correctness and performance she expects. After all, if she has dynamic resizing, she can resize to avoid collisions; and if she has collision resolution, collisions don't cause correctness issues. Which statement about these two properties true?
  - 1. Dynamic resizing alone will preserve both properties.
  - 2. Dynamic resizing alone will preserve correctness, but not performance.

- 3. Collision resolution alone will preserve performance, but not correctness.
- 4. Both are necessary to maintain performance and correctness.

**Solution:** Both are necessary to maintain performance and correctness. Without collision resolution, no correctness: could have an actual hash collision, and then no amount of resizing will let both be entered into the table. Without dynamic resizing, the load factor will get large, and everything will turn into a linear-time lookup (assuming chaining).

- (c) Suppose that Alyssa decides to implement resizing. If Alyssa is enlarging a table of size m into a table of size m', and the table contains n elements, what is the best time complexity she can achieve?
  - 1.  $\Theta(m)$
  - 2.  $\Theta(m')$
  - 3.  $\Theta(n)$
  - 4.  $\Theta(nm')$
  - 5.  $\Theta(m+m')$
  - 6.  $\Theta(m+n)$
  - 7.  $\Theta(m'+n)$

**Solution:**  $\Theta(m'+n)$ . It takes O(m') time to create a new hash table (allocating the memory can take constant time, but it then needs to be initialized). It takes O(m+n) time to go through each slot in the old table and copy each item. In total, it comes out to  $\Theta(m'+m+n)$ , but since m < m', the answer is just  $\Theta(m'+n)$ .

(d) In lecture, we discussed doubling the size of our hash table. Ivy H. Crimson begins to implement this approach (that is, she lets m'=2m) but stops when it occurs to her that she might be able to avoid wasting half of the memory the table occupies on empty space by letting m'=m+k instead, where k is some constant. Does this work? If so, why do you think we don't do it? There is a good theoretical reason as well as several additional practical concerns; a complete answer will touch on both points.

**Solution:** Theoretically, our cost will now be O(n) even after amortization. Loosely speaking, we were able to achieve O(1) amortized cost because we performed an O(n) time operation every O(n) step. Now, however, we're performing this O(n) operation every O(1) steps. Practically, the computer will play more nicely with operations based around doubling (doubling is a fast operation, allocating memory blocks of sizes that are powers of two has plenty of advantages, etc).

## **Problem 4-2.** [10 points] **Python Dictionaries**

We're going to get started by checking out a file from Python's Subversion repository at svn.python.org. The Python project operates a web frontend to their version control system, so we'll be able to do this using a browser.

#### Problem Set 4

Visit http://svn.python.org/projects/python/trunk/Objects/dictnotes.txt.

These are actual notes prepared by contributors to the Python project, as they currently exist in the Python source tree. (Cool! Actually, this document is a fascinating read—and you should be able to understand most of it.) Read over the seven use cases identified at the top of this document.

- (a) Let's examine the "membership testing" use case. Which statement accurately describes this use case?
  - 1. Many insertions right after creation, and then mostly lookups.
  - 2. Many insertions right after creation, and then only lookups.
  - 3. A workload of evenly-mixed insertions/deletions and lookups.
  - 4. Alternating rounds of insertions/deletions and lookups.

**Solution:** Many insertions right after creation, and then mostly lookup.

- (b) Now imagine that you have to pick a hash function, size, collision resolution strategy, and so forth (all of the characteristics of a hash table we've seen so far) in order to make a hash table perfectly suited to this use case alone. Pick the statement that best describes the choices you might make.
  - 1. A large minimum size and a growth rate of 2.
  - 2. A small minimum size and a growth rate of 2.
  - 3. A large minimum size and a growth rate of 4.
  - 4. A small minimum size and a growth rate of 4.

**Solution:** Membership testing can benefit from highly sparse hash tables, so let's set growth factor to 4. We're going to be inserting a bunch of things up front, so let's start with a larger minimum size.

## **Problem 4-3.** [55 points] Matching DNA Sequences

The code and data used in this problem are available on the course website. Please take a peek at the README.txt for some instructions.

Ben Bitdiddle has recently moved into the Kendall Square area, which is full of biotechnology companies and their shiny, window-laden office buildings. While mocking their dorky lab coats makes him feel slightly better about himself, he is secretly jealous, and so he sets out to earn one of his very own. To pick up the necessary geek cred, he begins experimenting with DNA-matching technologies.

Ben would like to create mutants to do his bidding, and to get started, he'd like to know how closely related the creatures he's collected are. If two sequences contain mostly the same subsequences in mostly the same places, then they're likely closely related; if they don't, they probably aren't. (This is, of course, a gross oversimplification.)

For our purposes, we'll represent a DNA sample as a sequence of characters. (These characters will all be upper-case. You can look at the Wikipedia page on nucleotides for a list of code characters

and their meanings.) These sequences are very long, so comparing subsequences of them quickly is important. We've provided code in kfasta.py that reads the .fa files storing this data.

- (a) Let's start with subsequenceHashes, which returns all length-k subsequences and their hashes (and perhaps other information, if there's anything else you might find useful).
  - Hint: There will likely be many of these matches; the DNA sequences are tens of millions of nucleotides long. To avoid keeping them all in memory at once, implement your function as a generator. See the Python reference materials available online for details if you aren't familiar with this important language construct.
- (b) Implement Multidict and verify that your work passes the simple sanity tests provided.
  - Multidict should behave just like a Python dictionary, except that it can store multiple values per key. If no values exist for a key, it returns an empty list; otherwise, it returns the list of associated values. You may (and probably should) use the Python dictionary in your implementation.
- (c) Now it's time to implement <code>getExactSubmatches</code>. Ignore the parameter m for the time being; we'll get to that in the next part. Again, implementing this function as a generator is probably a good idea. (You will probably have many, many matches—think about the combinatorics of the situation briefly.) As a hint, consider that much of the work has already been done by <code>Multidict</code> and <code>subsequenceHashes</code>; also take a peek at the <code>RollingHash</code> implementation we've given you. With these building blocks, your solution probably does not need to be very complex (or more than a few lines).

This function should return pairs of offsets into the inputs. A tuple (x, y) being returned indicates that the k-length subsequence at position x in the first input matches the subsequence at position y in the second input.

- We've provided a simple sanity test; your solution should be correct at this point (that is, dnaseq.py will produce the right output) but it'll probably be too slow to be useful. If you like, you can try running it on the first portion of two inputs; we've provided two such prefixes (the short files in the data directory) that might be helpful.
- (d) The most significant reason why your solution is presently too slow to be useful is that you are hashing and inserting into your hash table tens of millions of elements, and then performing tens of millions of lookups into that hash table. Implement intervalSubsequenceHashes, which returns the same thing as subsequenceHashes except that it hashes only one in m subsequences. (A good implementation will not do more work than is necessary.) Modify your implementation of getExactSubmatches to honor m only for sequence A. Consider why we still see approximately the same result, and why we can't further improve performance by applying this technique to sequence B as well.

#### Problem Set 4

(e) Run comparisons between the two human samples (paternal and maternal) and between the paternal sample and each of the animal samples.

Feel free to take a peek at how the image-generation code works. Conceptually, what it's doing is keeping track of how many of your (x, y) match tuples land in each of a two-dimensional grid of bins, each of which corresponds to a pixel in the output image. At the end, it normalizes the counts so that the highest count observed is totally black and an empty bin is white.

Think for a second about what a perfect match (e.g., comparing a sequence to itself) should look like. Try comparing the two human samples you have (maternal and paternal), one of the humans against the chimp sample, and then against the dog sample. Make sure your results make sense!

We've posted what our reference solution produced for the human-human comparison, the human-chimp comparison, and the human-dog comparison.

Please submit the code that you wrote. (You should only have had to modify dnaseq.py, so that's all you need to submit.)

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