COMP 2402 W24 Lab 4 Specifications

Trees & HashSets & Maps (& Lists)

Prelab: due on brightspace by Friday March 8th, 3:00pm (no lates)

Programming: due on gradescope by Wednesday March 13th, 3:00pm (24h late ok)

Postlab: due on brightspace.ca by Wednesday March 20th, 3:00pm (no lates)

Topic focus: Lec 1 - 15

Feedback on Lab 2 Feedback

Thank you very much to the 32 of you that took the time to fill out Lab 2 feedback. I was planning on writing up a report like last time but Life got in the way. But I've carefully read over your feedback and there were some common themes:

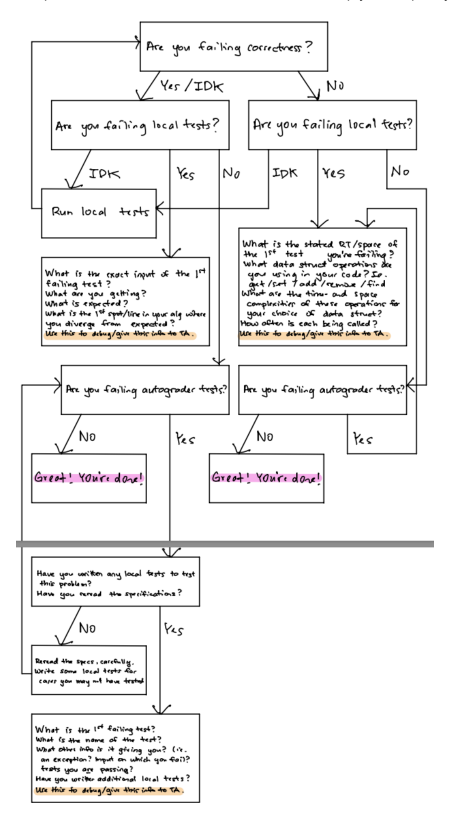
- Talk about iterators in lecture (will do next time! I mentioned their benefits in the in-person section but maybe my videos do not. Note to self to add that in.)
- For any local test written after I read the Lab 2 Feedback, you'll notice that the "Expect" comment is now in the output text as well. I hope this reduces some of your pattern matching time. Thanks for the suggestion, mystery student :
- Lab 2 took longer than Lab 1 (which is natural, but maybe some warning could be given.) I can/should also emphasize how you should ask for help when you are stuck. Our TAs are on piazza all. the. time. While during "deadline week" you might not get a response right away, our average response time is 43 minutes. We've had over 600 posts in 6 weeks so we've been busy! You can't see all the private posts we get, but if you ask a question during "forum monitoring" times and you don't get a response asap it *could* be that the TA is busy responding to a bunch of other private posts.
- autograder: some issues remain although many of you appreciate getting feedback before the deadline. Without the autograder you wouldn't have any hints as to whether you had a good data structure choice and since this course is about that choice, you would have no idea whether your choice is good. But I know it can be frustrating to not know where your bugs are. Sadly, this frustration will persist throughout your computer science career in various ways. I will continue to work on naming the tests better.

Changes from Labs 1-3

The autograders have more structured feedback, so that it's a bit better at being a "checkpoint." For example, many of the PhyloTree tests are for k=1, or when the Strings are not null, etc. You can focus on these special cases first and then worry about more general cases, if you wish.

There are still no hidden tests. In a sense, the autograder variability serves as a "hidden test" in that you sometimes have to look at your code and determine its complexity (gasp!) and decide whether it matches the desired complexity rather than relying on the occasionally flaky tests.

Finally, I made this new flow chart to help you determine where your problems are and how to communicate those problems to the course staff so that we can help you as quickly as possible!



Lab Objectives

The labs in this course are meant to give you the opportunity to practice with the topics of this course in a way that is challenging yet also manageable. At times you may struggle and at others it may seem more straight-forward; keep trying and practicing and you will improve.

Specifically, Lab 4 aims to improve your

- 1. Implementation Skills
 - Implement iterative operations for tree-based data structures similar to the BinaryTree and BinarySearchTree (add, find, remove, iterate, traverse, rotations).
- Design Skills
 - Use object-oriented programming concepts such as abstraction and polymorphism to provide flexibility in your choices of which interface implementation to use.
- 3. Critical Thinking
 - Demonstrate a solid understanding of the pros and cons of various implementations of the SSet (randomized, expected, worst-case) and of the situations when an unsorted set or map might be a better choice than a sorted set or list. This includes considering the time- and space-complexity of various operations in different data structures.
- 4. Algorithmic Thinking
 - Apply algorithmic thinking to design efficient algorithms for common tree-based operations, such as find(x), iteration over nodes using pointers, careful pointer manipulation in tree rotations, and removing recursion to avoid a big call stack.
- 5. Error Handling
 - Implement appropriate error handling and boundary checks to ensure the robustness of the data structures.
- 6. Testing
 - Determine the necessary tests to ensure your algorithms are correct and efficient.
- 7. Planning
 - Maintain or adopt good academic and programming habits through the pre-lab.
- 8. Reflection
 - Reflect on your choice of data structures and algorithms through the post-lab.

Assignment Components

Details of each component follow later in the specifications.

- 1. (6.7 points) Prelab (complete on brightspace)
- 2. (80 points) Programming portion (submit on gradescope.ca)
 - a. (50 points) PhyloTree Implementation
 - b. (10 points) Genes implementation
 - c. (10 points) GenesOrder Implementation
 - d. (10 points) GenesCount implementation
- 3. (13.3 points) Postlab (complete on brightspace)

Grading Criteria & Submission Guidelines

See the <u>Grading Criteria</u> and <u>Submission Guidelines</u> of Lab 1; they are the same.

Collaboration & Academic Integrity

- 1. Individual work is expected. Any collaboration should be explicitly mentioned and acknowledged at the top of each file. It is okay to discuss high-level approaches with your peers, or low-level syntax-type questions, but you must construct your solution on your own (as in: you have to formulate the code of your solution on your own.) Consider the analogy of writing an essay. You might talk with a peer about the high-level concepts of your thesis, or you might ask them about grammar or even phrasing of individual sentences. But you should not be writing the essay sentence-by-sentence with someone else's help; in the end you have to sit down and write that thing on your own.
- 2. Plagiarism will result in severe consequences. Ensure that all code and documentation are your own work. Do not send code to or receive code from any source except for course staff or the textbook, even if you change a thing here or there. It helps to keep the analogy of an essay in mind; it is not okay to take a paragraph from a friend and then rearrange some of the words or replace some with a thesaurus. It's not even okay to paraphrase each sentence. It is not okay to send your essay to a peer. Similarly here, you cannot start with code that is not yours and then "make it your own" with minor edits. Automated tools for detecting plagiarism will be employed in this course.
- 3. The same restrictions apply to **Al programmers** (such as chatGPT, copilot). You can use them to help with basic syntax (e.g. "spelling" and "grammar") or to understand broad concepts (e.g. getting feedback on a thesis) but you have to formulate your solution in code on your own (e.g. you have to write that essay yourself.)
- 4. Note that **contract cheating sites** are known, unauthorized, and regularly monitored. Some of these services employ misleading advertising practices and have a high risk of blackmail and extortion.
- 5. Every student should be familiar with the Carleton University student academic integrity policy. Academic integrity is upheld in this course to the best of Prof Alexa's abilities, as it protects the students that put in the effort to work on coursework within the allowable parameters. Potential violations must be reported to the Dean of Academic Integrity. If you ever have questions about what is or is not allowable regarding academic integrity, please do not hesitate to reach out to course staff. We are happy to answer.

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Workflow

In a perfect world, this is how you would complete Lab 4:

- 1. Attend or watch the relevant lectures that are listed in the heading of this document.
- 2. Read the <u>lab objectives</u> listed on the second page of this document. For each data structure listed there, review its important algorithms as well as the time- and space-complexity of its methods. This should give you some pros and cons for each.
- 3. Carefully read each problem detailed in the Programming section of this document.
 - a. Make sure you understand the problem.
 - b. Try the given examples by hand to get a better understanding of the problem.
 - c. Pay special attention to any special cases or edge cases.
 - d. Try more examples of your own devising if you need them.
 - e. Do not start programming yet!
 - f. Consider attending or watching the lab's workshop video, posted on brightspace.
- 4. Once you have completed steps 1-3, you are ready to do the <u>prelab</u> (brightspace).
- 5. Complete the programming portion of the lab.
 - a. Take this one problem at a time. Any order should be okay.
 - b. Remember what you learned in Steps 1-4 as you brainstorm solutions.
 - c. Test locally at frequent intervals. Do not write a whole program then test it afterwards. See the section on Local Tests to help you here.
 - d. Submit to gradescope.ca whenever you have made good progress, but do not use gradescope.ca as your only tests. Gradescope keeps your most recent score unless you select a different submission to be active. See the section on the Gradescope Autograder to help you debug here.
 - e. If you're stuck on a problem for more than 30 minutes, ask for help using the <u>How to Get Help</u> section. Move on to something else until help has arrived.
- 6. Once the late programming deadline has passed, complete the post-lab (brightspace).
 - a. If you did well on the programming portion, this is not meant to take too long.
 - b. There are resources available to help you posted on brightspace under the Lab 2 module. There is a solutions walk-through video for each part, and also a debrief document where I walk through the problem solving process and learning engagement I was hoping you would experience. You might consider looking at these before completing the prelab if you had trouble with any of the programming parts.
 - c. You do not have to have completed the programming parts in order to do the postlab. If you were stuck on a problem, this is an opportunity to look at the sample solution videos, to figure out what went wrong for you, and to still learn what you were meant to learn.

Coding Environment Setup

Lab 1's Coding Environment Setup will work with Lab 4 (replace Lab1/11 with Lab4/14). The file structure and many of the files will be the same, with new and different files as well.

Programming Components

Programming Notes

- 1. You may use <u>java.util.HashSet</u> instead of ods.MultiplicativeHashSet if you want the fastest implementation of a HashSet. Note that HashSet has an iterator.
- 2. You may use <u>java.util.HashMap</u> as well, for a fast implementation of a HashMap. Note that HashMap has an iterator over its keys or values.
- 3. If you want to use Collections.sort on an array-based Collection, you can!

```
e.g. ArrayDeque<String> ad = ...  // initialize ad etc
   Collections.sort(ad);  // now ad is sorted!
```

This takes O(n log n) time, where n is the number of elements in ad.

You can also sort a Collection using a new comparison rule (a Comparator), eg

```
e.g. Collections.sort(ad,MyComp); // sort ad using MyComp
where you may have defined MyComp elsewhere like this
class MyComp implements Comparator<String> {
    public int compare(String a, String b) {
        // Fill in compare
    }
}
```

Here are some examples on <u>geeksforgeeks</u>. Note that sorting a linked-list-based Collection can be done, but java first dumps the linked list into a structure with fast random access, and then sorts on that better structure, then dumps the sorted thing back into your linked-list-based structure. Keep that in mind, I suppose.

4. If you want to store objects of a class (of your own definition) in a sorted set, you need to ensure your class implements Comparable. For example, you could have

```
class MyClass implements Comparable<MyClass> {
    int x;
    int y;
    public int compareTo(MyClass b) {
        // Fill in compareTo
    }
}
```

Here are some examples on geeksforgeeks.

5. Note that you should not need recursion for any of the problems. You can/should run all tests with the -Xss144k flag (or, the smallest heap size your local machine will allow.) Sometimes you'll pass such tests even with recursion.

Implementation Practice [50 marks]

PhyloTree [50 marks]

PhyloTree represents a tree of distinct Strings (where a String can ostensibly represent a DNA sequence; no biology knowledge necessary for the lab). Your task is to implement the described methods according to the specifications below, using the starter code and lecture to guide you. The provided implementations are incomplete and incorrect, but should be enough to compile and run the (failing) local and autograder tests.

The data structure is a modified BinaryTree of Strings of some fixed length k, where each String is either null or of some length k>0 that is fixed for the given tree. The tree represents a "phylogenetic tree," which is essentially a hypothesized family tree for DNA. When a node has value null it means we haven't determined that ancestor in the tree yet. Each (non-null) String of the tree should be distinct and of length k.

This is the general idea. Let's get to some details.

Inner Class

Node (Represents a Node in our tree.)

String s
Node parent, left, right
boolean mark
boolean[] set

The String of length k in this Node, or null. The parent, left child and right child of this Node. Helpful node-specific boolean, used in LCA. Helpful node-specific set, used in computeSet.

Fields

Node r
 The root Node for the tree.

• int n The number of nodes in the tree.

• int k The length of non-null Strings in the tree, k>0.

 HashMap<String, Node> stringsToNodes A map from each non-null String stored

in the Tree to the node that contains that String. Used to speed up various methods.

Implemented Methods & Constructor

PhyloTree()
 Initializes the fields so that this tree is empty and k = 1.

- PhyloTree(int k)
 Initializes the fields so that this tree is empty and initializes k to the parameter.
- public int size()

 Returns the number of nodes in the tree.
- public void clear()
 Clears the tree so that it contains no elements.
- public String toString()
 Returns a list-based String representation of the tree in in-order that is helpful for debugging purposes.
- public String prettyPrint()
 Returns a tree-like String representation of the tree that is helpful for debugging purposes. This is only useful for smaller trees; for larger trees with missing children it can misrepresent the tree so use it cautiously.
- public int height()
 Returns the first Node in an in-order traversal. Note that his is a recursive method so use it with caution.
- protected Node firstNode()
 Returns the first Node in an in-order traversal.
- protected Node nextNode (Node w)
 Returns the Node that follows w in an in-order traversal.
- public Iterator<String> iterator()
 Returns an <u>Iterator</u> that iterates over the Strings of the tree in in-order order. Note that the Strings are not being stored in sorted order, so it may not appear sorted.

Constructor & Methods For You to Implement

Implement all the following methods without recursion.

Don't forget to update stringsToNodes when you add a non-null String to the PhyloTree!

Use the examples from lecture (e.g. traverse2, size2, findLastNode) to guide you here.

 public boolean addChild(String parent, String child) throws IllegalArgumentException
 Adds a Node containing child as the child of the existing parent.
 If parent is null, adds child as new root, with existing root as its left child. If parent has no children, add child as a left child.

If parent has left child, add child as a right child.

Throws an IllegalArgumentException if parent already has 2 children.

Throws an IllegalArgumentException if parent is not null and doesn't exist.

Throws an IllegalArgumentException if child is not null and already exists

Throws an IllegalArgumentException if child is not null and its length is not k.

If you've implemented addChild, it is a very good idea to test this locally and on the autograder before proceeding. If your addChild isn't working, none of the other tests are likely to work.

• public PhyloTree (ArrayStack<String> a) // Constructor Initializes the tree to contain the elements of a, where a contains the elements of the tree in "level order" (i.e. a[0] is the root, a[1] and a[2] are the root's children, and so forth.) Throws an IllegalArgumentException if non-null duplicates exist. Throws an IllegalArgumentException if any non-null String has length!= k, where k is the length of the (non-null) Strings in a (they should all have same length.)

If you've implemented PhyloTree, it is a very good idea to test this locally and on the autograder before proceeding. If this constructor isn't working, many of the tests that depend on this constructor will also fail.

public String LCA(String s, String t)
 Returns the "least common ancestor" (LCA) of the nodes containing s and t.
 The LCA is the first ancestor that s and t have in common (the one farthest from root r).
 Throws an IllegalArgumentException if s or t are null or aren't in the tree.

Note: The \mathtt{Node} class has a \mathtt{mark} field that can help you keep track of, say, nodes you've seen. Just remember that if you "mark" a node, you should "unmark" it before you return from LCA otherwise it will be "marked" the next time you call it.

Hint: break your code into manageable parts. Perhaps separate out the logic of checking for exceptions from finding the nodes that contain s and t from the logic that find the LCA of those nodes.

public void fixUp(String s)
 Performs a series of single rotations involving s until s is in a "good spot."
 In particular, if the node containing s is not in "binary search tree" order relative to its parent (e.g. it's a left child greater than its parent, or a right child less than its parent), use the appropriate single rotation to swap the parent-child relationship.
 Proceed up the tree as necessary until s is "good" relative to its parent.

If s's parent is null, it is in a "good spot".

Note that the rest of the tree may be completely disordered; that's okay.

Throws an IllegalArgumentException if s is null or is not in the tree.

Hint: single rotations come up in Treaps and BinaryHeaps; look there for guidance.

• public void computeSets(int index)

For this problem, each Node can hold a String of length k of DNABases (or is null).

(You can use the provided class field DNABases = $\{A, C, G, T\}$ here.)

Initially, all the leaves will be non-null; the internal nodes may or may not be null.

The given index indicates which index of the length-k String is relevant.

This problem asks you to compute each node's set of possible bases for the given index. You'll use the node's set field for this:

```
i.e. if {\tt DNABases[b]} is a possible base for the given index of node u, you'll set
```

```
u.set[b] = true
```

(ex: if (index index) of u can be A, G then u.set=[true, false, true, false])

Throws an IllegalArgumentException if the index<0 or index>=k.

Throws an IllegalArgumentException if any of the leaves are null.

Use the following algorithm to compute the sets:

```
for each node v in post-order (working from leaves upwards)
   if v is a leaf
        v's base is v's character at given index
   else if v has one child c
        v's set of bases is c's set of bases
   else
        if v's left and right children have overlapping sets
            v's set= the overlap/intersection of left &
        right
        else (the sets do not overlap)
        v's set = the combination/union of left & right
```

Note: there are some methods for debugging and testing that will be helpful: setToBases (Node u) - returns a String representation of node u's set

allSets() - returns an ArrayStack of all node's u's sets, in post-order

Desired Complexity

In the following table, let

- n denote the number of nodes in our tree;
- d(s,t) denote the length of the path between nodes s and t
- rotations denotes the maximum number of rotations needed to fix u (which in the worst-case could be the length of the root → u path, but in best case could be O(1))

marks	method	time complexity	(extra) space complexity
10	add(p,c)	O(1) ^E	0(1)
10	PhyloTree(a)	O(n)	0(n)
10	LCA(s,t)	O(d(s,t))	0(1)
10	fixUp(u)	O(rotations)	0(1)
10	computeSets(index)	O(n)	0(1)

Examples

addChild

Tree (pre)	k	method	Tree (post)
А	1	addChild(A,B)	A B
A B	1	addChild(A,C)	A B C
A B	1	addChild(B,C)	A B C
AA BB CC	2	addChild(CC,DD)	AA BB CC DD
AA BB CC	2	addChild(CC,null)	AA BB CC null
A B C	1	addChild(null,D)	D A B C
A B C	1	addChild(A,D)	IllegalArgumentException (parent already has 2 children)
A B C	1	addChild(B,C)	IllegalArgumentException (child already exists)
A B C	1	addChild(D,E)	IllegalArgumentException (parent not found)

A	1	addChild(B,DD)	IllegalArgumentException
ВС			(child is not of length k)

PhyloTree

method	Tree (post)	k	n
PhyloTree(A,B)	A B	1	2
PhyloTree(B,A)	B A	1	2
PhyloTree(A,B,C,D)	A B C D	1	4
PhyloTree(null,null,AA, BB, CC)	null null AA BB CC	2	5
PhyloTree(A,B,C,D,E,F,G,H,null)	B C D E F G H null	1	9
PhyloTree(A,A)	IllegalArgumentEx (duplicate elemen		n
PhyloTree (A, BB)	IllegalArgumentEx (A and BB have di		

LCA

Tree	k	method	output
A	1	LCA(A,A)	A
A B	1	LCA(A,B)	A
A B	1	LCA(B,B)	В
AA BB CC	2	LCA(BB,CC)	AA
A B <mark>C</mark>	1	LCA(D,C)	A

D E			
A B C D E	1	LCA(D,E)	В
A B C D E	1	LCA(E,A)	A
null B C D E	1	LCA(E,C)	null
A B C	1	LCA(A, <mark>D</mark>)	IllegalArgumentException (input D not in the tree)
A B C	1	LCA(D, null)	IllegalArgumentException (input cannot be null)
A B C	1	LCA (B, <mark>DD</mark>)	IllegalArgumentException (input DD is not in the tree)

fixUp

Tree (pre)	method	Tree (post)
A B	fixUp(A)	А В
null B	fixUp(B)	null B
A B	fixUp(B)	B A
A B C	fixUp(B)	B A C
A B C	fixUp(C)	C A B
C B A	fixUp(A)	A C B
D A B C	fixUp(C)	C D B A

A B C	fixUp(null)	IllegalArgumentException (input is null)
A B C	fixUp(D)	IllegalArgumentException (input not in tree)
A B C	addChild(D,E)	IllegalArgumentException (parent not found)
A B C	addChild(B,DD)	IllegalArgumentException (child is not of length k)

computeSets

Tree (pre)	k	method	Tree with Sets
AT	2	computeSets(0)	AT[A]
AT	2	computeSets(1)	A <mark>T</mark> [T]
null <mark>A</mark> C AG	2	computeSets(0)	<pre>null[A] (intersection) AC[A] AG[A]</pre>
null A C	1	computeSets(0)	null[A,C] (union) A[A] C[C]
TT CC GG AC CA AT GT	2	computeSets(0)	TT[A] CC[A,C] GG[A,G] AC[A] CA[C] AT[A] GT[G]
A C	1	computeSets(0)	A[C] C[C]
A G T	1	computeSets(1)	<pre>IllegalArgumentException (index >= k)</pre>
A G null	1	computeSets(0)	IllegalArgumentException (leaf is null)

Interface Practice [30 marks]

Genes, Revisited

In lab 3 the Genes problem asked you to compute the number of different strands of length k encountered in the input. This problem is similar in many ways, so I recommend you first review a sample solution (yours or ours) or the debrief before tackling this problem.

Method Signature

```
public static int genes(InputGenerator<Character> gen, int k)
```

Method Behaviour

Decodes each gene (substrand) of length $\, \mathrm{k} \, > \, 0$ in the sequence generated by a given InputGenerator as follows, then returns the number of different decodings:

```
For gene consisting of input characters c_i c_{i+1} c_{i+2}. ... c_{i+k-1} we decode it as i \cdot c_i + (i+1) \cdot c_{i+1} + (i+2) \cdot c_{i+2} + \ldots + (i+k-1) \cdot c_{i+k-1}
```

Desired Complexity & Notes

 $O(n)^E$ time, where n is the number of characters generated. $O(k+d)^E$ space, where k is the integer input parameter, and d is the solution.

- Since there are 4 bases, $d \le 4^k$ and thus log d = O(k).
- For reference, in Java, A=65, C=67, G=71, U=85.
- For large n and k, it's conceivable your decoding will have integer overflow. That's okay, let it overflow. The tests will overflow as well, which is simpler than describing ways to prevent overflow.

Examples

characters generated by gen	k	decoding of individual characters	different decodings	output (d)	n
AAAA	3	[A·0,A·1,A·2,A·3] [0,65,130,195]	[<mark>195</mark> , 390]	2	4
AAAAU	3	[0, <mark>65,130,195</mark> ,340	[195, <mark>390</mark> , 665]	3	5
UGCGAAUAUA	3	[0,71,134,213,260,325,510,455,680,58	[205,418,607, 798,1095,1290, 1645,1720]	8	10
UUUUCCCCAAAA	12	[0,85,170,255,268,335,402,469,520,585,650,715]	[4454]	1	12

UUUUCCCCAAA	12	[0,85,170,255,268,335,402,469,520,585,650]	[]	0	11
c ₆₄ CGAc ₆₈	1	[, C·65, G·66, A·67,	[, 4355,]	<n< td=""><td>n</td></n<>	n
(no chars)	> 0		[]	0	0

Testing & Autograder

There are limited local tests in Genes.main and tests/GenesTest.java; see <u>Lab 1</u> for testing instructions. Submit Genes.java to gradescope; see <u>Lab 1</u> for submission instructions.

GenesOrder, Revisited

Method Signature

public static AbstractList<String> genesOrder(InputGenerator<Character>
gen, int k)

Method Behaviour & Notes

Returns a list of the different genes (substrands) of length k>0 in the sequence generated by a given InputGenerator, in sorted order.

An <u>AbstractList</u> is an abstract java interface that is implemented by all the ods List implementations (e.g. ArrayStack, ArrayDeque, DLList, SkiplistList, etc). Using AbstractList<String> as the return type means you have the flexibility to choose which of these List implementations best suits your code and the desired complexity requirements.

Desired Complexity & Notes

 $O(k^2n)^A$ time, where n is the number of characters generated.

0 (kd) space, where k is the integer input parameter, and d is the solution.

Since there are 4 bases, $d \le 4^k$ and thus log d = O(k).

Examples

characters generated by gen	k	output	n
AAAA	3	AAA	4
AUAAA	3	AAA, AUA, UAA	5

UGCGAAUAUA	3	AAU, AUA, CGA, GAA, GCG, UAU, UGC	10
UUUUCCCCAAAA	12	UUUUCCCCAAAA	12
(no characters)	> 0	0	0

Testing & Autograder

There are limited local tests in <code>GenesOrder.main</code> and <code>tests/GenesOrderTest.java;</code> see <code>Lab 1</code> for testing instructions. Submit <code>GenesOrder.java</code> to gradescope; see <code>Lab 1</code> for submission instructions.

GenesCount

Method Signature

public static AbstractList<String> genesCount(InputGenerator<Character>
gen, int k)

Method Behaviour & Notes

Returns a list of the different genes (substrands) of length k>0 in the sequence generated by a given InputGenerator, in order from most frequent to least frequent. When multiple genes have the same frequency, order them alphabetically.

Desired Complexity & Notes

```
O(k(n+d log d))^{AE} = O(k(n+kd))^{AE} time.
 O(kd) space.
```

Note: n is the number of characters generated, k is the parameter, and d is the solution. Since there are 4 bases, $d \le 4^k$ and thus $\log d = O(k)$.

Examples

characters generated by gen	k	output (d)	n
AAAA	3	AAA	4
AUAAA	3	AAA, AUA, UAA	5
AUAAUAA	3	AUA, UAA, AAU	5
UGCGAAUAUA	3	AUA, AAU, CGA, GAA, GCG, UAU, UGC	10
UUUUCCCCAAAA	12	UUUUCCCCAAAA	12
(no characters)	> 0		0

Testing & Autograder

There are limited local tests in <code>GenesCount.main</code> and <code>tests/GenesCountTest.java;</code> see <code>Lab 1</code> for testing instructions. Submit <code>GenesCount.java</code> to gradescope; see <code>Lab 1</code> for submission instructions.

Local Tests

See the <u>Local Tests section of Lab 1</u>; be sure to replace comp2402w2411 with comp2402w2414

Gradescope Autograder

See the <u>Gradescope Autograder section of Lab 1</u>; be sure to replace Lab 1 with Lab 4 where necessary.

How to Get Help

See the <u>How to Get Help section of Lab 1</u>. It will be updated with any new resources rather than duplicating information here.

Common Errors & Fixes

See the <u>Common Errors & Fixes section of Lab 1</u>. It will be updated with any new errors rather than duplicating information here.

Glossary

There is a glossary at the top of the <u>Problem Solving & Programming Tips</u> document. If you can't find a term listed there, please post publicly on piazza so that everyone can benefit from the answer!

Autograder Runtimes for Sample Solutions

To give you a sense for the autograder runtimes for "perfect scores" you can see some of the screenshots that follow.

addChild

```
addChild correctness ~5 nodes k=1 (0/0)
Running test (1) addChild correctness ~5 nodes k=1
        Test passed.
        Time taken: 0:00:00.890379
addChild correctness ~10 nodes long skinny k=2 (0.5/0.5)
Running test (2) addChild correctness ~10 nodes long skinny k=2
       Test passed.
       Time taken: 0:00:01.003476
addChild correctness ~10 nodes balanced k=2 (0.5/0.5)
Running test (3) addChild correctness ~10 nodes balanced k=2
        Test passed.
       Time taken: 0:00:01.011344
addChild correctness ~10 nodes random structure k=2 (0.5/0.5)
Running test (4) addChild correctness ~10 nodes random structure k=2
       Test passed.
       Time taken: 0:00:00.923837
addChild exceptions (0.5/0.5)
Running test (5) addChild exceptions
       Test passed.
       Time taken: 0:00:01.034277
addChild correctness ~30 nodes, random structure k=3 (0.5/0.5)
Running test (6) addChild correctness \sim 30 nodes, random structure k=3
        Test passed.
        Time taken: 0:00:01.224481
addChild O(n) time (0.5/0.5)
Running test (7) addChild O(n) time
       Test passed.
        Time taken: 0:00:01.109479
addChild O(1) time (4.5/4.5)
Running test (8) addChild O(1) time
       Test passed.
        Time taken: 0:00:01.635391
addChild O(1) space and O(1) call stack (2.5/2.5)
Running test (9) addChild O(1) space and O(1) call stack
       Test passed.
       Time taken: 0:00:02.095690
```

PhyloTree

PhyloTree correctness, some null, k=1 (0.5/0.5)

Running test (11) PhyloTree correctness, some null, k=1
 Test passed.
 Time taken: 0:00:00.971501

PhyloTree correctness k=2 (0.5/0.5)

Running test (12) PhyloTree correctness k=2
Test passed.
Time taken: 0:00:01.011076

PhyloTree correctness k>=2 (0.5/0.5)

Running test (13) PhyloTree correctness k>=2
 Test passed.
 Time taken: 0:00:00.851926

PhyloTree exceptions (0.5/0.5)

Running test (14) PhyloTree exceptions
Test passed.
Time taken: 0:00:00.917263

PhyloTree O(n^2) time (0.5/0.5)

Running test (15) PhyloTree O(n^2) time
Test passed.
Time taken: 0:00:00.871402

PhyloTree O(n) time (4.5/4.5)

Running test (16) PhyloTree O(n) time
Test passed.
Time taken: 0:00:01.617529

PhyloTree O(n) space and O(1) call stack (2.5/2.5)

Running test (17) PhyloTree O(n) space and O(1) call stack
 Test passed.
 Time taken: 0:00:01.453938

```
LCA(sib, sib) correctness, k \ge 2 (0.5/0.5)
Running test (20) LCA(sib, sib) correctness, k>=2
        Test passed.
        Time taken: 0:00:00.917021
LCA(parent,child) correctness, k>=2 (0.5/0.5)
Running test (21) LCA(parent,child) correctness, k>=2
        Test passed.
        Time taken: 0:00:00.986909
LCA(s1, s2) for many combinations, k \ge 2 (0.5/0.5)
Running test (22) LCA(s1, s2) for many combinations, k>=2
        Test passed.
        Time taken: 0:00:01.028533
LCA exceptions (0.5/0.5)
Running test (23) LCA exceptions
        Test passed.
        Time taken: 0:00:00.958338
LCA O(n) time (0.5/0.5)
Running test (24) LCA O(n) time
        Test passed.
        Time taken: 0:00:01.061863
LCA O(1) time (4.5/4.5)
Running test (25) LCA O(1) time
        Test passed.
        Time taken: 0:00:01.535447
LCA O(n) space and O(1) call stack (2.5/2.5)
Running test (26) LCA O(n) space and O(1) call stack
        Test passed.
        Time taken: 0:00:01.526890
```

fixUp

```
fixUp correctness, many left rotations, k>=2 (0.5/0.5)
Running test (31) fixUp correctness, many left rotations, k>=2
        Test passed.
        Time taken: 0:00:00.924202
fixUp correctness, both rotations, k>=2 (0.5/0.5)
Running test (32) fixUp correctness, both rotations, k>=2
        Test passed.
        Time taken: 0:00:01.074886
fixUp exceptions (0.5/0.5)
Running test (33) fixUp exceptions
        Test passed.
        Time taken: 0:00:00.901181
fixUp O(n) time nodes near root (0.25/0.25)
Running test (34) fixUp O(n) time nodes near root
        Test passed.
        Time taken: 0:00:00.944982
fixUp O(1) time nodes near root (2.25/2.25)
Running test (35) fixUp O(1) time nodes near root
        Test passed.
        Time taken: 0:00:01.629240
fixUp O(n) time nodes far from root (0.25/0.25)
Running test (36) fixUp O(n) time nodes far from root
        Test passed.
        Time taken: 0:00:00.960360
fixUp O(1) time nodes far from root (2.25/2.25)
Running test (37) fixUp O(1) time nodes far from root
        Test passed.
        Time taken: 0:00:01.605332
fixUp O(1) space and O(1) call stack nodes near root (1.25/1.25)
Running test (38) fixUp O(1) space and O(1) call stack nodes near root
        Test passed.
        Time taken: 0:00:01.629480
fixUp O(1) space and O(1) call stack nodes far from root (1.25/1.25)
Running test (39) fixUp O(1) space and O(1) call stack nodes far from root
        Test passed.
        Time taken: 0:00:01.503864
```

computeSets

```
computeSets correctness all leaves same (0.25/0.25)
Running test (40) computeSets correctness all leaves same
        Test passed.
        Time taken: 0:00:01.245633
computeSets correctness all sets disjoint (0.25/0.25)
Running test (41) computeSets correctness all sets disjoint
        Test passed.
        Time taken: 0:00:01.106720
computeSets correctness sets partially overlap (0.5/0.5)
Running test (42) computeSets correctness sets partially overlap
        Test passed.
        Time taken: 0:00:01.217405
computeSets correctness all leaves same random structure (0.5/0.5)
Running test (43) computeSets correctness all leaves same random structure
        Test passed.
        Time taken: 0:00:01.263231
computeSets correctness random assignment (0.5/0.5)
Running test (44) computeSets correctness random assignment
        Test passed.
        Time taken: 0:00:01.263555
computeSets exceptions (0.5/0.5)
Running test (45) computeSets exceptions
        Test passed.
        Time taken: 0:00:00.981536
computeSets O(n^2) time (0.5/0.5)
Running test (46) computeSets O(n^2) time
       Test passed.
        Time taken: 0:00:01.188615
computeSets O(n) time (4.5/4.5)
Running test (47) computeSets O(n) time
        Test passed.
        Time taken: 0:00:02.246903
computeSets O(1) space and O(1) call stack (2.5/2.5)
Running test (48) computeSets O(1) space and O(1) call stack
        Test passed.
        Time taken: 0:00:02.836965
```

Genes

```
Genes correctness edge case empty strand (0.5/0.5)
Running test (4) Genes correctness edge case empty strand
        Test passed.
        Time taken: 0:00:00.774596
Genes correctness random (1/1)
Running test (5) Genes correctness random
       Test passed.
        Time taken: 0:00:00.755454
Genes O(n^2) time n~5,000, k~O(n), d~O(n) (0/0)
Running test (6) Genes O(n^2) time n-5,000, k-O(n), d-O(n)
       Test passed.
        Time taken: 0:00:00.841082
Genes O(nk^2) time n~100,000, k~100, d~O(n) (0.5/0.5)
Running test (7) Genes O(nk^2) time n~100,000, k~100, d~O(n)
       Test passed.
        Time taken: 0:00:01.145054
Genes O(nk) time n~15,000, k~O(n), d~O(n) (0.5/0.5)
Running test (8) Genes O(nk) time n-15,000, k-O(n), d-O(n)
       Test passed.
        Time taken: 0:00:00.846708
Genes O(n log d) time n~400,000, k~O(n), d~O(n) (2/2)
Running test (9) Genes O(n log d) time n~400,000, k~O(n), d~O(n)
       Test passed.
        Time taken: 0:00:01.627077
Genes O(n) time n~2,000,000, k~O(1), d~O(n) (2/2)
Running test (10) Genes O(n) time n~2,000,000, k~O(1), d~O(n)
       Test passed.
        Time taken: 0:00:02.152718
Genes O(n) space n~10,000, k, d=O(n) (0.5/0.5)
Running test (11) Genes O(n) space n~10,000, k, d=O(n)
        Test passed.
        Time taken: 0:00:00.891074
Genes O(k+d) space n=1,000,000, k, d=O(n) (2/2)
Running test (12) Genes O(k+d) space n=1,000,000, k, d=O(n)
       Test passed.
        Time taken: 0:00:03.287951
```

GenesOrder

```
GenesOrder correctness structured random (1/1)
Running test (3) GenesOrder correctness structured random
        Test passed.
        Time taken: 0:00:00.777766
GenesOrder correctness edge case empty strand (0.5/0.5)
Running test (4) GenesOrder correctness edge case empty strand
        Test passed.
        Time taken: 0:00:00.713601
GenesOrder correctness random (1/1)
Running test (5) GenesOrder correctness random
        Test passed.
        Time taken: 0:00:00.943812
GenesOrder O(k n^2) time n~5,000, k~O(n), d~O(1) (0/0)
Running test (6) GenesOrder O(k n^2) time n^5,000, k^0(n), d^0(1)
        Test passed.
        Time taken: 0:00:00.927420
GenesOrder O(k^2 n) time n~100,000, k~1,000, d~O(1) (1/1)
Running test (7) GenesOrder O(k^2 n) time n~100,000, k~1,000, d~O(1)
        Test passed.
        Time taken: 0:00:01.544293
GenesOrder O(k^2 d) time n~15,000, k~O(n), d~O(n) (2/2)
Running test (8) GenesOrder O(k^2 d) time n-15,000, k-O(n), d-O(n)
        Test passed.
        Time taken: 0:00:01.997045
GenesOrder O(k^2 d) time n~20,000, k~O(n), d~O(n) (2/2)
Running test (9) GenesOrder O(k^2 d) time n\sim20,000, k\sim0(n), d\sim0(n)
        Test passed.
        Time taken: 0:00:01.734473
GenesOrder O(n) space n=10,000, k, d=O(1) (0.5/0.5)
Running test (10) GenesOrder O(n) space n=10,000, k, d=O(1)
        Test passed.
        Time taken: 0:00:00.833136
GenesOrder O(kd) space n=1,000,000, k, d=O(1) (2/2)
Running test (11) GenesOrder O(kd) space n=1,000,000, k, d=O(1)
        Test passed.
        Time taken: 0:00:01.784532
```

GenesCount

```
GenesCount correctness random (1/1)
Running test (5) GenesCount correctness random
        Test passed.
        Time taken: 0:00:00.982121
GenesCount O(k n^2) time n~5,000, k~O(n), d~O(1) (0/0)
Running test (6) GenesCount O(k n^2) time n-5,000, k-O(n), d-O(1)
        Test passed.
        Time taken: 0:00:00.953249
GenesCount O(k^2 n) time n~60,000, k~1,000, d~O(1) (1/1)
Running test (7) GenesCount O(k^2 n) time n-60,000, k-1,000, d-O(1)
        Test passed.
        Time taken: 0:00:01.813724
GenesCount O(k^2 d) time n~8,000, k~O(n), d~O(n) (2/2)
Running test (8) GenesCount O(k^2 d) time n-8,000, k-O(n), d-O(n)
        Test passed.
        Time taken: 0:00:02.061771
GenesCount O(k^2 d) time n~10,000, k~O(n), d~O(n) (2/2)
Running test (9) GenesCount O(k^2 d) time n-10,000, k-O(n), d-O(n)
        Test passed.
        Time taken: 0:00:01.761667
GenesCount O(n) space n=10,000, k, d=O(1) (0.5/0.5)
Running test (10) GenesCount O(n) space n=10,000, k, d=O(1)
        Test passed.
        Time taken: 0:00:00.745329
GenesCount O(kd) space n=800,000, k, d=O(1) (2/2)
Running test (11) GenesCount O(kd) space n=800,000, k, d=O(1)
        Test passed.
        Time taken: 0:00:01.163159
```