

Algorithms - Class Notes

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Introduction To Algorithms

Introduction To Algorithms

1.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 1.1 Algorithms
- Chapter 1.2 Algorithms As A Technology
- Chapter 2.1 Insertion Sort
- Chapter 2.2 Analyzing Algorithms
- Chapter 2.3 Designing Algorithms
- Chapter 3.1 Asymptotic Algorithms
- Chapter 3.2 Standard Notations And Common Functions

1.0.2 Lectures

The lecture videos for this week are:

- Introduction To Algorithms ≈ 29 min.
- Insertion Sort ≈ 44 min.
- Time And Space Complexity ≈ 31 min.
- Asymptotic Notation ≈ 31 min.
- Pitfalls: Logarithms And Exponentials ≈ 16 min.

1.0.3 Assignments

The assignment for this week is:

• Problem Set 1 - Introduction To Algorithms

1.0.4 Quiz

The quizzes for this week are:

• Quiz 1

1.0.5 Chapter Summary

The first chapter that is being covered this week is **Chapter 1: The Role Of Algorithms In Computing**. The first section from this chapter is **Section 1.1: Algorithms**.

Section 1.1: Algorithms

Overview

At its core, an algorithm is a step-by-step procedure or a set of rules designed to perform a specific task or solve a particular problem. In computer science, algorithms are fundamental as they provide the methodical instructions that a computer follows to execute various tasks.

Key Aspects

- Step-by-Step Procedure: An algorithm is like a recipe in a cookbook. It contains steps that are performed in a sequence. Each step is clear and unambiguous. For example, an algorithm to add two numbers will have steps like: 'take the first number', 'take the second number', and 'add them together'.
- Specific Task or Problem: Algorithms are designed with a goal in mind. This can range from simple tasks like sorting a list of numbers to more complex operations like compressing data or finding the shortest path in a network.
- Efficiency: An important aspect of algorithms is their efficiency, which is generally measured in terms of time (how fast the algorithm runs) and space (how much memory it uses). Efficient algorithms can handle large data sets or complex computations more effectively.
- **Determinism and Correctness**: Typically, for a given input, an algorithm should produce a predictable and correct output. This consistency is crucial for reliability in computer programs.
- Pseudocode and Implementation: Algorithms are often initially described in pseudocode, a language-like notation that outlines the algorithm steps in human-readable form. They are then implemented in programming languages like Python, Java, or C++.

Algorithms form the backbone of all computer programs and applications. Understanding them is key to being an effective computer scientist, as they help you to understand how to approach problems systematically and efficiently.

The next section from this chapter is Section 1.2: Algorithms As A Technology.

Section 1.2: Algorithms As A Technology

Overview

When we talk about algorithms as a technology, we're essentially considering them as tools or solutions that can be applied to solve real-world problems using computers. The efficiency of these algorithms is a critical aspect, as it determines how effectively and quickly these problems can be solved.

Key Aspects

- Algorithms as Tools: Just like a hammer is a tool for a carpenter, algorithms are tools for computer scientists. They are applied to process data, solve complex calculations, make decisions based on input, and more. In the modern world, algorithms are behind everything from simple web searches to complex machine learning models.
- Measuring Efficiency: The efficiency of an algorithm is measured mainly in two ways:
 - **Time Complexity**: This refers to the amount of time an algorithm takes to complete its task as a function of the size of the input data. It's often expressed using Big \mathcal{O} notation (like $\mathcal{O}(n)$, $\mathcal{O}(\log(n))$, etc.), which gives an upper bound on the time. A faster algorithm has lower time complexity.
 - **Space Complexity**: This measures the amount of memory space required by an algorithm to run. An efficient algorithm uses space judiciously, especially important when dealing with large data sets.
- Importance of Efficiency: Efficient algorithms can handle larger data sets and more complex problems without requiring excessive computational resources. This is especially important in a world where data is growing exponentially. For example, an inefficient sorting algorithm might work fine for a small list but becomes impractically slow for a list with millions of elements.

- Trade-offs: There's often a trade-off between time and space complexity. Sometimes, an algorithm that's faster will use more memory, and vice versa. The choice of algorithm often depends on the specific constraints and requirements of the problem you're trying to solve.
- Real-World Applications: Efficient algorithms power everything from Google's search engine to the routing of data over networks. Inefficient algorithms can lead to slow performance, high energy consumption, and poor scalability.

Understanding the efficiency of algorithms is key to designing effective solutions in computer science. It's about finding the right balance between speed and resource usage, which can vary greatly depending on the problem at hand.

The next chapter that is being covered this week is **Chapter 2: Getting Started**. The first section from this chapter is **Section 2.1: Insertion Sort**.

Section 2.1: Insertion Sort

Overview

Insertion sort is a simple sorting algorithm that builds the final sorted array (or list) one item at a time. It's much less efficient on large lists than more advanced algorithms like quicksort, heapsort, or merge sort. However, its simplicity makes it easy to understand and implement, especially for small datasets or as a teaching tool.

Key Aspects

• Basic Concept: Imagine you are playing cards. When you're dealt a card, you insert it into the correct position to keep your hand sorted. Insertion sort works similarly. It iterates through an input list and removes one element per iteration, finding the location it belongs to in the already-sorted part of the list, and inserts it there.

• How It Works:

- Start with the second element of the list (the first element is considered already sorted).
- Compare this element with the elements before it, moving backward.
- If the current element is smaller than the compared element, swap them.
- Continue moving backward and swapping as long as the current element is smaller than the compared elements.
- Once the correct position is found where the current element is no longer smaller than the compared element, or you reach the beginning of the array, the current element is placed, and the algorithm moves to the next element in the unsorted part of the list.
- Repeat this process until the entire list is sorted.

• Time Complexity:

- Best Case: $\mathcal{O}(n)$ This happens when the list is already sorted, and only minimal comparisons are needed.
- Average and Worst Case: $\mathcal{O}(n^2)$ When the list is in reverse order, or elements are randomly distributed, each element may need to be compared with all other sorted elements.
- Space Complexity: $\mathcal{O}(1)$ Insertion sort is a space-efficient algorithm because it only requires a constant amount of additional storage space.

• Use Cases:

- Small Lists: It's efficient for small data sets.
- Partially Sorted Lists: It's efficient if the list is already partially sorted.
- Online Algorithms: It can sort the list as it receives it, making it useful in situations where the complete list of data isn't available initially (online algorithms).

• Advantages:

- Simple to understand and implement.
- Efficient for small data sets.
- More efficient than other simple algorithms like bubble sort.
- Stable: Maintains the relative order of equal elements.
- Can sort a list as it receives it.

· Disadvantages:

- Inefficient for large data sets due to its $\mathcal{O}(n^2)$ time complexity.
- Many shifts and comparisons are required.

Insertion sort is a great starting point for understanding sorting algorithms and concepts like in-place sorting and algorithm efficiency. Here is the insertion sort in C++.

```
void insertionSort(int arr[], int length) {
   int i, key, j;
   for (i = 1; i < length; i++) {
        key = arr[i]; // The element to be inserted in the sorted sequence
        j = i - 1;

        // Move elements of arr[0..i-1], that are greater than key,
        // to one position ahead of their current position
        while (j >= 0 && arr[j] > key) {
        arr[j + 1] = arr[j];
        j = j - 1;
        }
        arr[j + 1] = key; // Place key at after the element just smaller than it.
    }
}
```

The next section from this chapter is **Section 2.2:** Analyzing Algorithms.

Section 2.2: Analyzing Algorithms

Overview

Analyzing algorithms is an essential part of computer science, as it helps you understand how efficient an algorithm is in terms of time and space usage. This analysis is crucial for choosing the right algorithm for a particular problem, especially when dealing with large datasets or resource-constrained environments.

Key Aspects

- **Time Complexity**: This measures how the runtime of an algorithm increases with the size of the input. It's a way to estimate the time taken for an algorithm to run, relative to the size of its input.
 - Big O Notation: The most common way to express time complexity. It provides an upper bound on the time complexity, ensuring that the algorithm will not take more time than the given complexity in the worst case.
 - Common Time Complexities: Constant $\mathcal{O}(1)$, logarithmic $\mathcal{O}(\log(n))$, linear $\mathcal{O}(n)$, linearithmic $\mathcal{O}(n\log(n))$, quadratic $\mathcal{O}(n^2)$, cubic $\mathcal{O}(n^3)$, exponential $\mathcal{O}(2^n)$, etc.
- Space Complexity: This refers to the amount of memory space required by an algorithm in its execution. Space complexity becomes crucial for applications that run on devices with limited memory.
 - Auxiliary Space: Apart from space complexity, sometimes you look at the extra space or temporary space used by an algorithm.
- Worst, Average, and Best Case Analysis:
 - Worst Case: The maximum time or space required by the algorithm for any input of size n. It's a guarantee that the algorithm will not take more time or space than this.

- Average Case: It's more practical as it provides a more realistic measure of the algorithm's performance.
 It's the average time or space taken by the algorithm over all possible inputs.
- Best Case: The minimum time or space taken by the algorithm for any input of size n. It's usually not as useful as worst or average cases but sometimes relevant for comparison.
- Amortized Analysis: In some cases, an algorithm might have a very high time complexity for a particular operation but generally runs faster. Amortized analysis gives the average time per operation, taken over a sequence of operations.
- Empirical Analysis: This involves running the algorithm with different inputs and measuring the time and space it uses. This approach gives practical insights but is influenced by hardware and software environments.
- Theoretical vs. Practical: While theoretical analysis gives you complexity in terms of n, practical performance can be influenced by factors like constant factors, lower-order terms, the architecture of the machine, and so on.
- Algorithmic Paradigms: Sometimes, the analysis is also done based on the approach used by the algorithm, like Divide and Conquer, Dynamic Programming, Greedy Techniques, etc.

Understanding these aspects of algorithm analysis can help you predict how an algorithm will perform in various situations, which is crucial for making informed decisions in software development and system design.

The last section from this chapter is **Section 2.3: Designing Algorithms**.

Section 2.3: Designing Algorithms

Overview

Understanding these aspects of algorithm analysis can help you predict how an algorithm will perform in various situations, which is crucial for making informed decisions in software development and system design.

Procedure

• Divide:

- The first step is to divide the original problem into smaller sub-problems. These sub-problems should ideally be similar to the original problem but smaller in scale.
- The division continues recursively until the sub-problems are small enough to be solved straightforwardly (base case).

• Conquer:

- Solve each sub-problem. As you've broken the problem down into smaller pieces, these can often be solved more easily and efficiently.
- If the sub-problems are still complex, apply the divide and conquer strategy recursively to them.

• Combine:

- Finally, combine the solutions of the sub-problems to form a solution to the original problem.
- The method of combining may vary depending on the problem.

Characteristics and Advantages

- Recursive Nature: Divide and conquer is inherently recursive, as it involves solving smaller instances of the same problem.
- Efficiency: It can significantly reduce the time complexity for many problems, especially those where a direct approach would be inefficient.
- Parallelism: Sub-problems can often be solved in parallel, making divide and conquer algorithms well-suited for parallel processing and multi-threading.

Classic Examples

- Merge Sort: An array is divided into halves, each half is sorted (conquer), and then the sorted halves are merged back together (combine).
- Quick Sort: The array is partitioned around a 'pivot' element, and then the sub-arrays are sorted independently.
- **Binary Search**: The problem of searching for an element in a sorted array is divided by comparing the target with the middle element, effectively halving the search space each time.
- Strassen's Algorithm for Matrix Multiplication: This approach improves the efficiency of matrix multiplication by dividing matrices and reducing the number of multiplications needed.

Considerations

- Overheads: The overhead of recursion and combining solutions can sometimes make a divide and conquer algorithm less efficient for small datasets.
- Stack Space: Recursive methods can use significant stack space, which might be a limiting factor for very deep recursive calls.

Understanding Divide and Conquer is crucial for computer science students, as it forms the basis for many efficient and widely-used algorithms. It's a great example of how a problem can be simplified and made more manageable through recursion and systematic approach.

The next chapter that is being covered this week is **Chapter 3: Growth Of Functions**. The first section from this chapter is **Section 3.1: Asymptotic Notation**.

Section 3.1: Asymptotic Notation

Overview

Asymptotic notation in computer science is a mathematical tool used to describe the behavior of algorithms, particularly in terms of their time and space complexity. It provides a way to analyze an algorithm's efficiency by considering the limits of its performance as the input size grows towards infinity. There are three primary types of asymptotic notation:

Notations

- Big \mathcal{O} Notation (\mathcal{O} -notation):
 - **Description**: Big \mathcal{O} notation describes the upper bound of the complexity. It gives the worst-case scenario of an algorithm's growth rate.
 - **Usage**: It's used to describe the maximum amount of time (or space) an algorithm requires for any input of size n.
 - **Example**: If an algorithm has a time complexity of $\mathcal{O}(n^2)$, it means that the time taken will increase at most quadratically with the size of the input.
- Omega Notation (Ω -notation):
 - Description: Omega notation is used to describe the lower bound of the complexity. It provides a
 guarantee of at least this amount of resources the algorithm needs.
 - Usage: It shows the best-case scenario (minimum performance) for an algorithm.
 - **Example**: If an algorithm has a time complexity of $\Omega(n)$, it means that the algorithm will take at least linear time in the best case.
- Theta Notation (Θ -notation):

- **Description**: Theta notation tightly bounds the complexity from above and below, meaning it defines both the upper and lower limits of the time or space complexity.
- **Usage**: It's used when we want to indicate that an algorithm has both upper and lower bounds that are asymptotically the same.
- **Example**: An algorithm with time complexity $\Theta(n \log(n))$ will have its running time increase logarithmically in the best case and linearithmically in the worst case.

Key Points

- Asymptotic Behavior: These notations are not about providing exact running times or space requirements but rather describing the growth rate relative to the input size as it becomes very large.
- Simplification: Constants and lower-order terms are usually ignored in asymptotic analysis. For instance, $3n^2 + 2n + 1$ is simply $\mathcal{O}(n^2)$.
- Widely Used in Algorithm Analysis: Asymptotic notations are crucial for comparing algorithms, especially when deciding which algorithm to use for a given problem.

Understanding these notations is fundamental in computer science as it allows you to abstract away from machine-specific details and focus on the algorithm's inherent efficiency. It's an essential part of algorithm design and analysis, providing a common language to compare and discuss the performance of different algorithms.



Divide And Conquer

Divide And Conquer

2.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 4.1 The Maximum-Subarray Problem
- Chapter 4.2 Strassen's Algorithm For Matrix Multiplication
- Chapter 4.3 The Substitution Method For Solving Recurrences
- Chapter 4.4 The Recursion-Tree Method For Solving Recurrences

2.0.2 Lectures

The lecture videos for this week are:

- Divide And Conquer: Mergesort ≈ 38 min.
- Analysis Of Mergesort ≈ 24 min.
- Max Subarray Problem ≈ 24 min.
- Karatsuba Multiplication Algorithm ≈ 40 min.
- Solving Recurrences: Master Method ≈ 28 min.

2.0.3 Assignments

The assignment for this week is:

• Problem Set 2 - Divide And Conquer

2.0.4 Quiz

The quizzes for this week are:

• Quiz 2

2.0.5 Chapter Summary

The chapter for this week is **Chapter 4: Divide And Conquer**. The first section from this chapter is **Section 4.1: The Maximum Sub-Array Problem**.

Section 4.1: The Maximum Sub-Array Problem

Overview

The Maximum Sub-Array Problem involves finding the contiguous sub-array within a one-dimensional array of numbers which has the largest sum. It's a classic example of a divide-and-conquer approach in algorithm design.

Problem Statement

Given an array of integers (both positive and negative), the goal is to find a contiguous sub-array that maximizes the sum of its elements.

Divide and Conquer Strategy

- Divide: The array is divided into two halves.
- Conquer: Recursively find the maximum sub-array in each half.
- Combine: Determine the maximum sub-array that crosses the midpoint and combine the results of the three cases (left, right, and crossing the midpoint) to find the overall maximum.

Key Components

- Crossing Sub-Array: A crucial step is finding the maximum sub-array that crosses the midpoint. This is done by looking for the maximum sub-array ending at the midpoint from the left side and starting from the midpoint on the right side, then combining these two.
- Recursive Nature: The problem is recursively solved for the left and right halves, and the solutions are merged.

Algorithmic Steps

- 1. Base Case: If the array has only one element, return this element as the maximum sum.
- 2. Divide the Array: Find the midpoint of the array.
- 3. Recursive Calls:
 - Find the maximum sub-array in the left half.
 - Find the maximum sub-array in the right half.
- 4. Find the Crossing Maximum Sub-Array: Calculate the maximum sub-array that crosses the midpoint.
- 5. Combine Results: The maximum of these three values (left, right, crossing) is the solution.

Complexity Analysis

The time complexity of this algorithm is $\mathcal{O}(n \log(n))$, where n is the number of elements in the array. This is more efficient than the $\mathcal{O}(n^2)$ complexity of the brute-force approach.

The next section from this chapter is Section 4.2: Strassen's Algorithm For Matrix Multiplication.

Section 4.2: Strassen's Algorithm For Matrix Multiplication

Overview

Strassen's Algorithm is an innovative approach to matrix multiplication. It's known for reducing the computational complexity compared to the standard matrix multiplication technique.

Problem Statement

- Standard matrix multiplication of two $n \times n$ matrices has a time complexity of $\mathcal{O}(n^3)$.
- Strassen's Algorithm improves this, especially for large matrices.

Divide And Conquer Approach

- **Divide**: The algorithm divides each matrix into four $\frac{n}{2} \times \frac{n}{2}$ sub-matrices.
- Conquer: It recursively multiplies these smaller matrices.
- Combine: Utilizes fewer multiplications (7 instead of 8 in standard approach) to combine these sub-matrices into the final product.

Strassen's Formulae

- The algorithm uses 7 multiplication operations (M1 to M7) on combined elements of the sub-matrices, reducing the number of multiplications needed.
- These 7 products are then used to calculate the entries of the final $n \times n$ matrices.

Key Concepts

- Sub-Matrix Multiplication: Instead of multiplying matrices directly, the algorithm operates on smaller matrices.
- Addition and Subtraction of Matrices: Employed extensively to prepare the matrices for the 7 multiplication operations.

Complexity Analysis

- Time Complexity: Approximately $\mathcal{O}(n^{2.81})$, which is lower than the $\mathcal{O}(n^3)$ of the standard method.
- This is due to the master theorem in the analysis of divide-and-conquer algorithms.

Limitations And Extensions

- Overhead for smaller matrices can outweigh its theoretical efficiency.
- The algorithm has paved the way for further research into even more efficient matrix multiplication algorithms.

The next section from this chapter is Section 4.3: The Substitution Method For Solving Recurrences.

Section 4.3: The Substitution Method For Solving Recurrences

Overview

The Substitution Method, also known as the 'Guess-and-Check' method, is used to solve recurrence relations, which often arise in the analysis of recursive algorithms. It involves making a guess about the solution's form and then proving it by induction.

Problem Statement

- Recurrence relations define a sequence of values using recursion.
- The method helps in determining the time complexity of recursive algorithms by solving these relations.

Basic Steps

- Guess the Form: Start with a hypothesis about the form of the solution (usually based on experience or pattern observation).
- Verify by Induction: Use mathematical induction to prove that the guess is correct.
- Refine if Necessary: If the initial guess doesn't hold, refine it and attempt the proof again.

Process Of Induction

- Base Case: Verify that the solution holds for the initial term(s) of the sequence.
- Inductive Step: Assume the solution holds for a general term (say, the n-th term) and then prove it for the next term (n+1).

Common Recurrence Types

- Linear Recurrences (e.g. T(n) = T(n-1) + n).
- Divide-and-Conquer Recurrences (e.g. T(n) = 2T(n/2) + n).

Advantages And Challenges

- Advantage: Allows for a tailored approach for different recurrences.
- Challenge: Finding the correct guess can be non-trivial and requires insight.

Example

For a recurrence like $T(n) = 2T(\lfloor n/2 \rfloor) + n$, one might guess that $T(n) = \mathcal{O}(n \log(n))$ and then use induction to prove it.

The next section from this chapter is Section 4.4: The Recursion-Tree Method For Solving Recurrences.

Section 4.4: The Recursion-Tree Method For Solving Recurrences

Overview

The Recursion-Tree Method is a visual and intuitive approach to solving recurrence relations. It involves drawing a tree to represent the recursive calls and their costs, helping to visualize the structure of the recursion.

Purpose And Application

- Used to determine the time complexity of recursive algorithms.
- Particularly useful for recurrences arising from divide-and-conquer algorithms.

Steps In The Method

- **Draw the Recursion Tree**: Each node represents a recursive call, and the tree shows how the problem is divided and conquered.
- Calculate Cost at Each Level: Assign costs to each level of the tree based on the recurrence relation.
- Summarize the Total Cost: Add up the costs across all levels of the tree to find the total cost of the algorithm.

Tree Structure

- The root represents the initial problem.
- Child nodes represent sub-problems created by recursive calls.
- The depth of the tree correlates with the number of recursive calls.

Analyzing The Tree

- Determine the number of levels in the tree.
- Calculate the total number of nodes or the total cost at each level.
- Sum these costs to estimate the overall complexity of the recurrence.

Example Of Analysis

- For a recurrence like T(n) = 2T(n/2) + n, the tree will have 2 branches at each level, and the cost at each level will be proportional to n.
- The depth of the tree is typically $\log(n)$, leading to a total cost of $\mathcal{O}(n\log(n))$.

Advantages And Limitations

- Advantages: Intuitive and visual; helpful in understanding the pattern of recursive calls.
- Limitations: Can be complex for some recurrences; less precise than mathematical methods.

The last section from this chapter is Section 4.5: The Master Method For Solving Recurrences.

Section 4.5: The Master Method For Solving Recurrences

Overview

The Master Method provides a cookbook-style solution for solving recurrences, particularly those arising from divide-and-conquer algorithms. It's a formulaic approach that, when applicable, quickly gives the asymptotic behavior of the recurrence.

Applicability

Best suited for recurrences of the form T(n) = aT(n/b) + f(n), where:

- a is the number of sub-problems in the recursion.
- n/b is the size of each sub-problem.
- f(n) is the cost of the work done outside the recursive calls.

The Three Cases Of The Master Method

- Case 1: If $f(n) = \mathcal{O}(n^c)$ where $c < \log_b(a)$, then $T(n) = \Theta(n^{\log_b(a)})$.
- Case 2: If $f(n) = \Theta(n^c)$ where $c = \log_b(a)$, then $T(n) = \Theta(n^c \log(n))$.
- Case 3: If $f(n) = \Omega(n^c)$ where $c > \log_b(a)$ and if $af(n/b) \le kf(n)$ for some constant k < 1 and sufficiently large n, then $T(n) = \Theta(f(n))$.

How to Use the Master Method

- Identify a, b, and f(n) in the given recurrence.
- Determine which of the three cases applies.
- Apply the corresponding formula to find the solution.

Examples

- Merge Sort: For T(n) = 2T(n/2) + n, it falls under Case 2 (since f(n) = n and $c = \log_2(2) = 1$), leading to $T(n) = \Theta(n \log(n))$.
- Binary Search: For T(n) = T(n/2) + 1, it falls under Case 2, giving $T(n) = \Theta(\log(n))$.

Advantages And Limitations

- Advantages: Simplifies the process of solving recurrences; provides quick and direct answers.
- Limitations: Not applicable to all types of recurrences; specific conditions must be met.



Heaps And Quicksort

Heaps And Quicksort

3.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 5.1 The Hiring Problem
- Chapter 5.2 Indicator Random Variables
- Chapter 5.3 Randomized Algorithms
- Chapter 5.4 Probabilistic Analysis And Further Uses Of Indicator Random Variables
- Chapter 6.1 Heaps
- Chapter 6.2 Maintaining The Heap Property
- Chapter 6.3 Building A Heap
- Chapter 6.4 The Heapsort Algorithm
- Chapter 6.5 Priority Queues
- Chapter 7.1 Description Of Quicksort
- Chapter 7.2 Performance Of Quicksort
- Chapter 7.3 A Randomized Version Of Quicksort
- Chapter 7.4 Analysis Of Quicksort

3.0.2 Lectures

The lecture videos for this week are:

- Heaps: Introduction And Basic Properties ≈ 24 min.
- Heap Operations: Bubble Up/Down ≈ 29 min.
- Heap Operations: Insert, Delete And Heapsort ≈ 29 min.
- Overview of Quicksort ≈ 14 min.
- Quicksort: Lomuto's Partitioning Algorithm ≈ 25 min.
- Quicksort: Worst Case And Average Case Analysis ≈ 28 min.

3.0.3 Assignments

The assignment for this week is:

- Problem Set 3 Heaps And Quicksort
- Programming Assignment 1 Heaps And Quicksort

3.0.4 Quiz

The quizzes for this week are:

• Quiz 3

3.0.5 Chapter Summary

The first chapter that we are covering this week is **Chapter 5: Probabilistic Analysis and Randomized Algorithms**. The first section that I we are covering from this section is **Section 5.1: The hiring problem**.

Section 5.1: The hiring problem

The Hiring Problem

The hiring problem is a classic example in the study of probabilistic analysis and randomized algorithms, illustrating the challenges and strategies involved in decision-making under uncertainty. It models a situation where an employer must decide whether to hire a candidate immediately after an interview, without the ability to recall previous candidates. The decision is based solely on the relative ranking of the current candidate compared to those interviewed so far. The employer aims to minimize the cost associated with hiring while maximizing the quality of the hire. This problem showcases the use of expected values to determine the best strategy and introduces randomized algorithms as a means to optimize hiring decisions in an uncertain environment.

• Problem Description

- In the hiring problem, an employer interviews n applicants one by one in a random sequence and must decide immediately after each interview whether to hire the candidate. The goal is to hire the best candidate with minimal costs, assuming each hiring incurs a fixed cost. The challenge arises because the decision must be made without knowledge of future candidates.
- Example: Consider interviewing 5 candidates randomly, where the decision to hire is based on whether a candidate is superior to all previously interviewed ones, potentially leading to multiple hires and associated costs.

• Probabilistic Analysis

- Probabilistic analysis of the hiring problem assumes equal likelihood for any order of candidate quality, aiming to calculate the expected number of hires. This analysis reveals that, depending on the hiring strategy, the expected number of hires can significantly vary, with an optimal strategy minimizing this expectation.
- Example: While the worst-case scenario results in hiring after every interview (n hires), an optimal strategy could significantly reduce the expected number of hires, often related to the logarithm of the number of candidates ($\log n$).

• Randomized Algorithms

- To improve decision-making under uncertainty, randomized algorithms may be employed in the hiring process. These algorithms can optimize the expected outcome by introducing randomness into the hiring criteria, thereby potentially reducing the total hiring cost while maintaining a high probability of hiring the best candidate.
- Example: Implementing a randomized threshold for hiring decisions that adjusts based on the proportion
 of candidates already interviewed to those remaining can optimize hiring costs and outcomes.

The next section that will be covered from this chapter this week is **Section 5.2: Indicator Random Variables**.

Section 5.2: Indicator Random Variables

Indicator Random Variables

Indicator random variables are a fundamental concept in probability theory and statistical analysis, particularly useful in the study of algorithms. They simplify the analysis of complex random processes by breaking them down into simpler, binary outcomes. An indicator random variable, denoted as I_A , takes the value 1 if event A occurs and 0 otherwise. This binary nature makes it incredibly powerful for probabilistic analysis and expected value calculations, as it allows for the straightforward aggregation of probabilities across multiple events.

Definition

- An indicator random variable I_A is defined for an event A, such that $I_A = 1$ if A occurs, and $I_A = 0$ if A does not occur. This simple mechanism provides a clear way to quantify the occurrence of events in probabilistic terms.
- Example: In a card game, let A be the event that a drawn card is an ace. The indicator random variable I_A would be 1 if the drawn card is an ace, and 0 otherwise.

• Expected Value of an Indicator Random Variable

- The expected value, $E[I_A]$, of an indicator random variable is equal to the probability of the event A occurring, denoted by P(A). This is because $E[I_A] = 1 \cdot P(A) + 0 \cdot P(\neg A) = P(A)$, where $\neg A$ represents the complement of A.
- Example: If the probability of drawing an ace from a standard deck of cards is $\frac{1}{13}$, then $E[I_A] = \frac{1}{13}$.

• Using Indicator Random Variables in Algorithms

- Indicator random variables can significantly simplify the analysis of algorithms, especially when calculating the expected number of occurrences of an event. By summing the indicator variables for each instance of the event, we can easily compute the overall expected number of occurrences.
- Example: Consider an algorithm that processes n items, where each item has a probability p of triggering a specific action (event A). The expected number of triggered actions is the sum of n indicator random variables, each representing the occurrence of A for an item, resulting in an expected total of $n \cdot p$.

The next section that will be covered in this chapter Section 5.3: Randomized Algorithms.

Section 5.3: Randomized Algorithms

Randomized Algorithms

Randomized algorithms incorporate randomness as a part of their logic, making decisions not solely based on input but also on random values. This approach can lead to simpler, faster, or more efficient algorithms compared to their deterministic counterparts. The use of randomness can help in overcoming obstacles such as worst-case scenarios or unknown inputs. Randomized algorithms are widely applied in fields such as cryptography, algorithm design, and computational complexity.

• Advantages of Randomized Algorithms

- Randomized algorithms often have simpler implementations and can achieve better performance on average compared to deterministic algorithms. They are particularly useful in dealing with adversarial inputs or in scenarios where the input distribution is unknown.
- Example: QuickSort algorithm, where the pivot is chosen randomly. This randomization ensures that the average time complexity remains $O(n \log n)$, even in the worst-case input scenarios.

• Types of Randomized Algorithms

- Monte Carlo algorithms provide a probabilistic solution that may be incorrect with a certain probability.
 These algorithms are useful when a fast approximate solution is acceptable.
- Las Vegas algorithms always produce a correct solution, but their running time is variable and depends
 on the random choices made during execution. The algorithm terminates once a correct solution is found.
- Example: The ZPP (Zero-error Probabilistic Polynomial time) class includes algorithms that are Las Vegas type, ensuring correctness with variable execution time.

Sample Python Code for Randomized QuickSort

The Randomized QuickSort algorithm selects a pivot element randomly from the array to be sorted, partitioning the array around this pivot, and recursively sorting the subarrays. This random selection helps to ensure that the algorithm achieves good average-case performance.

```
import random
    def randomized_partition(arr, low, high):
        pivot_index = random.randint(low, high)
        arr[pivot_index], arr[high] = arr[high], arr[pivot_index]
        pivot = arr[high]
        i = low - 1
        for j in range(low, high):
             if arr[j] < pivot:</pre>
                 arr[i], arr[j] = arr[j], arr[i]
        arr[i+1], arr[high] = arr[high], arr[i+1]
        return i+1
    def randomized_quicksort(arr, low, high):
        if low < high:
    pi = randomized_partition(arr, low, high)</pre>
16
17
             randomized_quicksort(arr, low, pi-1)
             randomized_quicksort(arr, pi+1, high)
    # Example usage
    arr = [10, 7, 8, 9, 1, 5]
    randomized_quicksort(arr, 0, len(arr)-1)
    print("Sorted array:", arr)
```

This Python code snippet demonstrates how Randomized QuickSort works by selecting a pivot element randomly and using it to partition the array. The use of randomness in selecting the pivot reduces the likelihood of encountering the worst-case performance scenario, which is typical in the standard QuickSort when the smallest or largest element is always chosen as the pivot.

The last section from this chapter is **Section 5.4: Probabilistic Analysis And Further Uses Of Indicator Random Variables**.

Section 5.4: Probabilistic Analysis And Further Uses Of Indicator Random Variables

Probabilistic Analysis and Further Uses of Indicator Random Variables

Probabilistic analysis leverages the concept of probability to analyze the behavior of algorithms under the assumption that the input is determined randomly. This analysis often uses indicator random variables to simplify the computation of expected values, particularly in complex algorithms. Indicator random variables serve as a powerful tool for breaking down the analysis into manageable parts, each representing a basic event that can either occur or not.

• Basics of Probabilistic Analysis

- Probabilistic analysis calculates the average-case complexity of algorithms, assuming a probabilistic distribution of all possible inputs. It helps in understanding the behavior of algorithms under typical conditions.
- Example: Analyzing the average number of comparisons in QuickSort or the average-case running time of a hashing algorithm.

• Further Uses of Indicator Random Variables

- Indicator random variables are extensively used to simplify the analysis of the expected value of complex random variables by breaking the problem down into simpler, binary outcomes.
- Example: The number of collisions in a hashing algorithm can be analyzed using indicator random variables representing whether each pair of elements collides.

Mathematical Formulae For Expected Value Calculation

The expected value of a sum of indicator random variables can be calculated using the linearity of expectation. If $X = \sum_{i=1}^{n} I_{A_i}$, where I_{A_i} is the indicator random variable for event A_i , then the expected value of X, E[X], is the sum of the expected values of the indicator random variables:

$$E[X] = E\left[\sum_{i=1}^{n} I_{A_i}\right] = \sum_{i=1}^{n} E[I_{A_i}] = \sum_{i=1}^{n} P(A_i)$$

This formula is particularly useful because it allows the expected value of X to be computed directly from the probabilities of the individual events A_i without needing to consider the dependencies between them.

This section highlights the significance of probabilistic analysis in understanding the average-case performance of algorithms and how indicator random variables facilitate this analysis by enabling the straightforward calculation of expected values. Mathematical formulae within the highlight environment elucidate the method for calculating the expected value of a sum of indicator random variables, showcasing the elegance and power of this approach in algorithmic analysis.

The next chapter that we will cover this week is **Chapter 6: Heapsort**. The first section that will be covered from this section is **Section 6.1: Heaps**.

Section 6.1: Heaps

Heaps

Heaps are a specialized tree-based data structure that satisfies the heap property. In a heap, for any given node C with parent P, the key (the value) of P is either greater than or equal to C for a max heap, or less than or equal to C for a min heap. This property makes heaps useful for implementing priority queues and for the heapsort algorithm.

• Heap Properties

- A heap is a complete binary tree, meaning it is completely filled at all levels except possibly the lowest, which is filled from the left up to a point.
- In a max heap, for every node i other than the root, the value of node i is less than or equal to the value of its parent. Conversely, in a min heap, every node i's value is greater than or equal to the value of its parent. These properties can be succinctly represented as $A[\operatorname{parent}(i)] \geq A[i]$ for a max heap and $A[\operatorname{parent}(i)] \leq A[i]$ for a min heap, where A is the array representation of the heap.

• Basic Heap Operations

- Insertion Adding a new element to the heap while maintaining the heap property. This operation is typically $O(\log n)$ due to the need for potentially "bubbling up" the new element.
- Deletion Removing the root element from the heap (the maximum element in a max heap or the minimum in a min heap) and then restructuring the heap to maintain the heap property, usually through a process called "heapify," which is also $O(\log n)$.

Sample Python Code for Heapsort

Heapsort is an efficient comparison-based sorting algorithm that builds a heap from the input data and then repeatedly extracts the maximum element (in a max heap) or the minimum element (in a min heap) to build the sorted list.

```
1  def heapify(arr, n, i):
2     largest = i
3     left = 2 * i + 1
4     right = 2 * i + 2
5
6     if left < n and arr[largest] < arr[left]:
7          largest = left
8     if right < n and arr[largest] < arr[right]:
9          largest = right
10     if largest != i:</pre>
```

```
arr[i], arr[largest] = arr[largest], arr[i]
heapify(arr, n, largest)

def heapsort(arr):
    n = len(arr)
    for i in range(n // 2 - 1, -1, -1):
        heapify(arr, n, i)

for i in range(n-1, 0, -1):
        arr[i], arr[0] = arr[0], arr[i]
heapify(arr, i, 0)

# Example usage
arr = [12, 11, 13, 5, 6, 7]
heapsort(arr)
print("Sorted array is:", arr)
```

This code first builds a max heap from the input array. Then, it repeatedly removes the largest element from the heap (the root of the heap), placing it at the end of the array, and calls heapify to restore the max heap property for the remaining elements.

The next section that will be covered this week is **Section 6.2: Maintaining The Heap Property**.

Section 6.2: Maintaining The Heap Property

Maintaining the Heap Property

Maintaining the heap property is crucial in heap-based data structures, particularly in operations such as insertions, deletions, and the Heapsort algorithm. The Max-Heapify operation is a key procedure used to maintain the max heap property in a heap after an element has been added or removed.

• Max-Heapify Operation

- The Max-Heapify operation ensures that the heap property is maintained by comparing a node with its children and swapping it with one of them if necessary so that the node's value is not less than the values of the children. This process is applied recursively down the heap.
- For a node at index i, its left child is at index 2i+1 and its right child at 2i+2 in the array representation of the heap. The Max-Heapify operation is applied when the subtree rooted at i violates the heap property.

Sample Python Code for Max-Heapify

The Max-Heapify function is a cornerstone of heap operations, ensuring the max heap property is satisfied throughout the heap. This function is most commonly used in the Heapsort algorithm and when inserting or removing elements from a heap.

```
def max_heapify(arr, n, i):
    largest = i
    left = 2 * i + 1
    right = 2 * i + 2

if left < n and arr[left] > arr[largest]:
    largest = left
    if right < n and arr[right] > arr[largest]:
    largest = right
    if largest != i:
        arr[i], arr[largest] = arr[largest], arr[i]
        max_heapify(arr, n, largest)

# Example usage
    arr = [3, 2, 15, 5, 4, 45]
    n = len(arr)
    max_heapify(arr, n, 0)
    print("Max-Heapified array:", arr)
```

This Python code snippet demonstrates the Max-Heapify operation applied to an array that represents a heap. The function ensures that, for any given node, if the children's values are greater than the node's value, the largest value among the node and its children becomes the parent node, thus maintaining the max heap property.

The next section that will be covered this week is **Section 6.3: Building A Heap**.

Section 6.3: Building A Heap

Building a Heap

Building a heap from an unsorted array is an essential step in heap operations, particularly before performing the Heapsort algorithm. This process transforms an arbitrary array into a heap by applying the Max-Heapify operation in a bottom-up manner. The algorithm efficiently produces a max heap (or min heap, accordingly) by ensuring that all subtrees satisfy the heap property.

• Heap Construction Algorithm

- The process starts from the lowest non-leaf nodes and applies the Max-Heapify operation to each, moving upwards to the root of the heap. Since leaves are trivially heaps, starting from the first non-leaf node ensures that each application of Max-Heapify makes the subtree rooted at the current node a valid heap.
- This bottom-up approach guarantees that the heap property is maintained for all nodes. The efficiency of this method lies in its O(n) complexity, contrary to the intuitive $O(n \log n)$ if Max-Heapify were applied naively to each node starting from the root.

Sample Python Code for Building a Max Heap

Building a max heap involves organizing the elements of an array so that the value of each parent node is greater than or equal to the values of its children, satisfying the max heap property for the entire array.

```
def heapify(arr, n, i):
    largest = i
left = 2 * i + 1
    right = 2 * i + 2
    if left < n and arr[left] > arr[largest]:
         largest = left
    if right < n and arr[right] > arr[largest]:
         largest = right
    if largest != i:
         arr[i], arr[largest] = arr[largest], arr[i]
         heapify(arr, n, largest)
def build_max_heap(arr):
    n = len(arr)
        i in range(n // 2 - 1, -1, -1):
        heapify(arr, n, i)
# Example usage
arr = [1, 12, 9, 5, 6, 10]
build_max_heap(arr)
print("Max Heap:", arr)
```

This code snippet demonstrates how to transform an unsorted array into a max heap. The 'build_max_heap' function iterates over each non-leaf node, applying 'heapify' to ensure that the subtree rooted at each node satisfies the max heap property, ultimately building a max heap from the entire array.

The next section that is covered from this chapter is **Section 6.4: The Heapsort Algorithm**.

Section 6.4: The Heapsort Algorithm

The Heapsort Algorithm

The Heapsort algorithm is a comparison-based sorting technique based on the binary heap data structure. It's similar to selection sort where we first find the maximum element and place the maximum at the end. We repeat the same process for the remaining elements. Heapsort is particularly efficient for its ability to sort in-place, requiring no additional storage beyond what is needed for the list.

• Heapsort Procedure

- The algorithm begins by building a max heap from the input array.
- It then repeatedly removes the maximum element from the heap (the root of the heap), and moves it to the end of the array. After removing the maximum element, it must re-heapify the remaining elements to ensure the heap property is maintained.
- This process continues until all elements are removed from the heap and placed into the array in sorted order.
- The beauty of Heapsort lies in its $O(n \log n)$ time complexity for all cases: worst, average, and best.

Sample Python Code for Heapsort

Heapsort utilizes the heap data structure to sort an array in an efficient manner. The key process in Heapsort is to first transform the list of elements into a heap and then sort the elements using the heap properties.

```
def heapify(arr, n, i):
    largest = i
left = 2 * i + 1
    right = 2 * i + 2
    if left < n and arr[left] > arr[largest]:
         largest = left
    if right < n and arr[right] > arr[largest]:
         largest = right
    if largest != i:
         arr[i], arr[largest] = arr[largest], arr[i]
         heapify(arr, n, largest)
def heapsort(arr):
        len(arr)
    for i in range(n // 2 - 1, -1, -1):
    heapify(arr, n, i)
for i in range(n-1, 0, -1):
    arr[i], arr[0] = arr[0], arr[i] # swap
         heapify(arr, i, 0)
# Example usage
arr = [12, 11, 13, 5, 6, 7]
heapsort (arr)
print("Sorted array is:", arr)
```

This code showcases the Heapsort algorithm in action. The initial step is to build a max heap from the unsorted input array. Following this, the algorithm repeatedly swaps the first element of the array (the largest value in the heap) with the last element of the heap, reduces the heap size by one, and then heapifies the root element to ensure the max heap property. This process is repeated until the array is sorted.

The last section from this chapter is **Section 6.5**: **Priority Queues**.

Section 6.5: Priority Queues

Priority Queues

Priority queues are an abstract data type that operate similarly to regular queues or stacks, but with an added feature: each element has a "priority" associated with it. In a priority queue, an element with high priority is served before an element with low priority. If two elements have the same priority, they are served according to their order in the queue. Priority queues are typically implemented using heaps, as heaps provide an efficient way to maintain the order of the elements by their priority, allowing for quick insertion and removal of the highest or lowest priority item.

• Applications of Priority Queues

 Priority queues are used in many areas of computer science, including scheduling processes in operating systems, pathfinding algorithms like A* for AI and games, and in the implementation of algorithms like Dijkstra's shortest path algorithm.

Operations

- The primary operations of a priority queue include *insert* (or *enqueue*), which adds an element to the queue with a given priority; *peek* (or *find-min / find-max*), which returns the highest (or lowest) priority element without removing it; and *remove* (or *dequeue*), which removes and returns the element with the highest (or lowest) priority.

Sample Python Code for Priority Queue using Heap

This Python code example demonstrates a simple priority queue implemented with a binary heap. The heapq module in Python provides an easy way to maintain a heap, allowing the priority queue to efficiently perform its operations.

```
import heapq
class PriorityQueue:
    def __init__(self):
        self.heap = []

def insert(self, item, priority):
        heapq.heappush(self.heap, (priority, item))

def peek(self):
        return self.heap[0][1] if self.heap else None

def remove(self):
        return heapq.heappop(self.heap)[1] if self.heap else None

# Example usage
pq = PriorityQueue()
pq insert("Task 1", 3)
pq.insert("Task 2", 1)
pq.insert("Task 3", 2)
print("Peek at highest priority item:", pq.peek())
print("Remove highest priority item:", pq.remove())
print("Remove next highest priority item:", pq.remove())

print("Remove next highest priority item:", pq.remove())
```

In this implementation, the 'PriorityQueue' class utilizes a min heap to keep track of items by priority, where a smaller number indicates a higher priority. The 'insert' method adds a new item with its priority to the heap, the 'peek' method retrieves the highest priority item without removing it, and the 'remove' method removes and returns the highest priority item from the queue.

The last chapter this week is **Chapter 7: Quicksort**. The first section from this chapter is **Section 7.1: Description Of Quicksort**.

Section 7.1: Description Of Quicksort

Description of Quicksort

Quicksort is a highly efficient sorting algorithm that employs a divide-and-conquer approach to sort elements in an array. It was developed by Tony Hoare in 1959 and has since become one of the most widely utilized sorting algorithms. Quicksort works by selecting a 'pivot' element from the array and partitioning the other elements into two sub-arrays, based on whether they are less than or greater than the pivot. These sub-arrays are then sorted recursively, leading to a fully sorted array.

• Algorithm Steps

- Choosing a Pivot: The pivot can be selected through various strategies, such as picking the first element, the last element, the median, or a random element. The choice of pivot significantly affects the algorithm's performance.

- Partitioning: The array is rearranged so all elements less than the pivot come before it, while all elements greater than the pivot come after it. The pivot then occupies its correct position in the sorted array.
- Recursively Sorting Sub-arrays: The process is applied recursively to the sub-array of elements with smaller values and the sub-array of elements with larger values.

• Performance

- The average and best-case time complexity of Quicksort is $O(n \log n)$, making it highly efficient for large datasets. In the worst case, its time complexity can degrade to $O(n^2)$, especially when the smallest or largest element is consistently chosen as the pivot.
- Its practical efficiency comes from the low overhead and cache efficiency, making it faster in practice than other $O(n \log n)$ sorting algorithms.

• In-place Sorting

 Quicksort sorts the array in place, requiring no additional storage space except for the stack space used by recursion, enhancing its memory efficiency.

Sample Python Code for Quicksort

The following Python code illustrates the Quicksort algorithm, demonstrating the process of pivot selection, partitioning, and recursive sorting.

This implementation selects the last element as the pivot and uses the partitioning method to place the pivot element in its correct position in the sorted array. It then recursively sorts the sub-arrays before and after the pivot to achieve the sorted array.

The next section from this chapter is Section 7.2: Performance Of Quicksort.

Section 7.2: Performance Of Quicksort

Performance of Quicksort

The performance of Quicksort is one of the key reasons for its popularity in practical applications. It is known for its efficiency in sorting large datasets, with a performance that can vary significantly based on the choice of pivot and the arrangement of the input data.

• Time Complexity

- The average-case time complexity of Quicksort is $O(n \log n)$, where n is the number of elements in the array. This optimal performance is achieved through the effective division of the array into partitions and the efficient sorting of these partitions.
- The worst-case time complexity occurs when the smallest or largest element is consistently chosen as the pivot, leading to partitions of sizes n-1 and 1, resulting in a time complexity of $O(n^2)$. However, this scenario is rare, especially with good pivot selection strategies.

- The best-case scenario, with a time complexity of $O(n \log n)$, occurs when the partition process divides the array into two equal parts, thereby minimizing the depth of the recursion tree.

• Space Complexity

- Quicksort is an in-place sorting algorithm, but it requires additional space for the recursive function calls. The space complexity is $O(\log n)$ in the best case, which corresponds to the height of a balanced recursion tree.
- In the worst case, with unbalanced partitions, the space complexity can degrade to O(n) due to the depth of the recursion stack.

• Optimizations and Practical Performance

- Several strategies can optimize Quicksort's performance, including choosing a pivot using methods like
 the median-of-three, which selects the pivot as the median of the first, middle, and last elements of the
 partition.
- The use of a hybrid algorithm, such as switching to insertion sort for small partitions, can significantly reduce the overhead of recursive calls, thereby improving performance.
- The dual-pivot Quicksort, where two pivots are used to divide the array into three parts, has also been shown to offer improvements in sorting efficiency.

In summary, the performance of Quicksort is generally very good and often surpasses other sorting algorithms, especially for large arrays. With intelligent pivot selection and optimizations for small arrays, Quicksort can achieve near-optimal efficiency, making it a versatile choice for a wide range of sorting applications.

The next section for this chapter that is covered this week is Section 7.3: A Randomized Version Of Quicksort.

Section 7.3: A Randomized Version Of Quicksort

A Randomized Version of Quicksort

A randomized version of Quicksort enhances the traditional Quicksort algorithm by selecting the pivot in a random manner. This approach helps to optimize the performance of Quicksort by reducing the likelihood of encountering the worst-case time complexity of $O(n^2)$, regardless of the input distribution. The randomized selection of pivots ensures that the algorithm's average-case time complexity of $O(n \log n)$ is achieved more consistently.

Advantages of Randomization

- Randomization in Quicksort mitigates the risk of poor performance on already sorted or nearly sorted arrays, which are problematic for deterministic pivot selection methods.
- It provides a probabilistic guarantee on the algorithm's performance, making its time complexity predictable and generally close to the average-case scenario.

• Implementation Considerations

- The primary change in a randomized Quicksort algorithm is the method of pivot selection. Instead of choosing a fixed position, the pivot is chosen randomly from the sub-array that is currently being sorted.
- This random pivot selection can be efficiently implemented by swapping the randomly chosen element with the first or last element of the sub-array and then proceeding with the standard partitioning process.

Sample Python Code for Randomized Quicksort

The following Python code demonstrates the implementation of a randomized version of Quicksort, highlighting the randomized pivot selection.

```
import random

def randomized_partition(arr, low, high):
    pivot_index = random.randint(low, high)
    arr[pivot_index], arr[high] = arr[high], arr[pivot_index] # Swap pivot with the end element
    return partition(arr, low, high)
```

```
def partition(arr, low, high):
    pivot = arr[high]
    i = low - 1
for j in range(low, high):
        if arr[j] < pivot:</pre>
            i += 1
            arr[i], arr[j] = arr[j], arr[i]
    arr[i+1], arr[high] = arr[high], arr[i+1]
def randomized_quicksort(arr, low, high):
    if low < high:
        pi = randomized_partition(arr, low, high)
        randomized_quicksort(arr, low, pi-1)
        randomized_quicksort(arr, pi+1, high)
# Example usage
arr = [10, 7, 8, 9, 1, 5]
randomized_quicksort(arr, 0, len(arr)-1)
print("Sorted array using randomized Quicksort:", arr)
```

This implementation introduces randomness to the pivot selection process in Quicksort, effectively randomizing the algorithm. By selecting a random pivot, the algorithm minimizes the chance of experiencing the worst-case scenario, thus maintaining efficient performance across various input arrays.

The last section from this chapter is **Section 7.4: Analysis Of Quicksort**.

Section 7.4: Analysis Of Quicksort

Analysis of Quicksort

The analysis of Quicksort focuses on understanding its time complexity in various scenarios and the factors that influence its performance. Despite its worst-case time complexity of $O(n^2)$, Quicksort is often faster in practice than other sorting algorithms with the same average-case complexity of $O(n \log n)$. This efficiency is largely due to the constant factors hidden in the big-O notation, the way it benefits from cache memory, and its ability to sort in place with minimal additional memory requirements.

• Average-Case Analysis

- The average-case time complexity of Quicksort is $O(n \log n)$, which is derived under the assumption that all permutations of the input array are equally probable. This scenario occurs when the pivot divides the array into parts of reasonably proportional sizes, allowing the depth of the recursion to be logarithmic relative to the size of the array.

• Worst-Case Analysis

- The worst-case scenario happens when the partitioning routine produces one sub-array with n-1 elements and one with 0 elements, leading to a recursion depth of n. This scenario typically occurs when the smallest or largest element is always chosen as the pivot. The time complexity in this case is $O(n^2)$.

• Best-Case Analysis

- The best-case scenario for Quicksort occurs when the partition process divides the array into two equal parts at each step. This ideal situation also results in a time complexity of $O(n \log n)$, but with a smaller constant factor than in the average case.

Space Complexity

 $O(\log n)$

- The space complexity of Quicksort is $O(\log n)$ in the best case, corresponding to the height of a balanced recursion tree. In the worst case, the space complexity can increase to O(n) due to the stack space required for the recursive calls.

Optimizations

- Various optimizations can improve Quicksort's performance and reduce the likelihood of hitting the worst-case scenario. These include choosing the pivot via more sophisticated methods (such as the median of three or random selection), switching to a different sorting algorithm like insertion sort for small arrays, and using tail recursion optimization.

In summary, the efficiency of Quicksort in practice is attributed to its $O(n \log n)$ average-case time complexity and the various optimizations that can be applied to minimize the chances of encountering the worst-case scenario. Despite the theoretical $O(n^2)$ worst-case time complexity, these optimizations ensure that Quicksort remains one of the fastest sorting algorithms for practical applications.



Exam 1

Exam 1

4.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 8.1 Lower Bounds For Sorting
- Chapter 9.1 Minimum And Maximum
- Chapter 9.2 Selection In Expected Linear Time
- Chapter 9.3 Selection In Worst-Case Linear Time

4.0.2 Lectures

The lecture videos for this week are:

- Quickselect Algorithm ≈ 18 min.
- Deterministic Partitioning: Median of Medians Trick ≈ 27 min.
- Linear Time Sorting Algorithms ≈ 19 min.
- Binary Search Trees: Introduction ≈ 22 min.
- Binary Search Tree: Operations ≈ 31 min.

4.0.3 Assignments

The assignment for this week is:

• Problem Set 4 - Quick Select And Binary Search Trees

4.0.4 Quiz

The quizzes for this week are:

• Quiz 4

4.0.5 Exam

The exam for this week is:

- Spot Exam 1 Notes
- Spot Exam 1

4.0.6 Chapter Summary

The first chapter that is being discussed this week is **Chapter 8: Sorting in Linear Time**. The first section that we are going to cover from this chapter this week is **Section 8.1: Lower Bounds For Sorting**.

Section 8.1: Lower Bounds For Sorting

Lower Bounds for Sorting

The concept of lower bounds for sorting is fundamental in understanding the efficiency of comparison-based sorting algorithms. It establishes a theoretical minimum for the number of comparisons needed to sort a list of elements, providing a benchmark for evaluating sorting algorithm performances.

• Comparison-based Sorting Algorithms

- These algorithms sort elements by comparing them. Examples include QuickSort, MergeSort, and HeapSort.
- The efficiency of these algorithms is often measured by the number of comparisons they make.

• Decision Tree Model

- The decision tree is a conceptual tool used to represent the comparisons made by a sorting algorithm for a given number of elements.
- Each node represents a comparison between two elements, and each path from the root to a leaf represents
 a possible sequence of comparisons needed to sort the elements.

• Lower Bound on Comparisons

- In the decision tree model, the depth of the tree (the longest path from the root to a leaf) represents the worst-case number of comparisons.
- For a list of n distinct elements, there are n! (factorial of n) possible permutations, each corresponding to a unique leaf in the decision tree.
- The lower bound for any comparison-based sorting algorithm is $\Omega(n \log n)$ comparisons in the worst case. This is derived from the fact that the height of a balanced binary tree with n! leaves is $\log_2(n!)$, which is asymptotically equivalent to $n \log n$.

This theoretical limit is crucial because it implies that no comparison-based sorting algorithm can be faster than $\Omega(n \log n)$ in the worst case. Thus, algorithms achieving this bound, such as MergeSort and HeapSort, are considered asymptotically optimal among comparison-based sorts.

The last section that will be covered from this week is **Section 8.2**: **Counting Sort**.

Section 8.2: Counting Sort

Counting Sort

Counting Sort is a non-comparison-based sorting algorithm that sorts the elements of an array by counting the number of occurrences of each unique element. Its efficiency comes from directly determining the position of each element in the output sequence, making it particularly effective for sorting integers or other items where the range of potential values is known and not too large.

• Algorithm Overview

- Counting Sort works by creating an auxiliary array that counts the occurrences of each element in the input.
- The counts are then used to determine the positions of each element in the sorted array.

• Steps of Counting Sort

- Count each element in the input array and store the count in an auxiliary array.
- Accumulate the counts to get the starting index of each element.
- Place the elements in the output array based on their starting indexes and update the counts.

• Time Complexity

- The time complexity of Counting Sort is O(n+k), where n is the number of elements in the input array and k is the range of the input.
- This makes Counting Sort very efficient when k is not significantly larger than n.

Sample Python Code for Counting Sort

The following Python code demonstrates the implementation of the Counting Sort algorithm, showcasing its efficiency in sorting integers.

```
def counting_sort(arr):
    max_element = int(max(arr))
min_element = int(min(arr))
    range_of_elements = max_element - min_element + 1
    # Create count array to store the count of individual elements
    count_array = [0] * range_of_elements
output_array = [0] * len(arr)
    # Store the count of each number
    for i in range(0, len(arr)):
        count_array[arr[i]-min_element] += 1
    # Change count_array so that count_array[i] now contains actual
    # position of this element in output array
for i in range(1, len(count_array)):
        count array[i] += count array[i-1]
    # Build the output character array
    for i in range(len(arr)-1, -1, -1)
        output_array[count_array[arr[i] - min_element] - 1] = arr[i]
        count_array[arr[i] - min_element] -= 1
    # Copy the output array to arr, so that arr now
    # contains sorted characters
    for i in range(0, len(arr)):
        arr[i] = output_array[i]
    return arr
```

This implementation of Counting Sort demonstrates its non-comparative approach by using counting and index calculations to sort the array efficiently, especially useful for sorting integers within a known range.

The next chapter that is being covered this week is **Chapter 9: Medians And Order Statistics**. The first section that is being covered from this chapter for this week is **Section 9.1: Minimum And Maximum**.

Section 9.1: Minimum And Maximum

Minimum and Maximum

The problem of finding the minimum and maximum elements in an array is a fundamental task in computer science, often serving as a precursor to more complex operations like sorting or searching. Efficiently determining the minimum and maximum values can be done through various algorithms, with the goal of minimizing the number of comparisons required.

• Linear Search Approach

- The simplest method involves iterating through the array, comparing each element to keep track of the current minimum and maximum values found so far.
- This approach requires 2(n-1) comparisons in the worst case, where n is the number of elements in the array.

• Pairwise Comparison

- A more efficient method involves comparing elements in pairs, reducing the total number of comparisons.
- This method can find both the minimum and maximum with approximately $\frac{3n}{2}$ comparisons, significantly fewer than the linear search approach for large arrays.

Sample Python Code for Finding Minimum and Maximum

The following Python code snippets demonstrate the implementation of both the linear search approach and the pairwise comparison method to find the minimum and maximum values in an array.

```
1 # Linear Search Approach
```

```
def find min max linear(arr):
    if not arr:
         return None, None
    min_val = max_val = arr[0]
         val in arr[1:]:
         if val < min_val:</pre>
         min_val = val
elif val > max_val:
              max_val = val
     return min_val, max_val
# Pairwise Comparison Approach
def find_min_max_pairwise(arr):
    if not arr:
         return None, None
     if len(arr) % 2 == 0:
         min_val = min(arr[0], arr[1])
         max_val = max(arr[0], arr[1])
         start = 2
         min_val = max_val = arr[0]
         start = 1
    for i in range(start, len(arr), 2):
         local_min = min(arr[i], arr[i + 1])
local_max = max(arr[i], arr[i + 1])
         min_val = min(min_val, local_min)
max_val = max(max_val, local_max)
    return min_val, max_val
```

These implementations illustrate different strategies for finding the minimum and maximum values, showcasing the trade-offs in terms of complexity and number of comparisons.

The last section that will be covered from this chapter this week is **Section 9.2: Selection In Expected Linear Time**.

Section 9.2: Selection In Expected Linear Time

Selection in Expected Linear Time

The task of selecting the i^{th} smallest element from an unsorted array can be efficiently solved using the Randomized-Select algorithm. This algorithm employs a randomized partitioning strategy, similar to that used in the randomized version of Quicksort, to ensure that its expected time complexity remains linear, or O(n), despite the worst-case scenario potentially being quadratic.

• Algorithm Overview

- Randomized-Select utilizes the divide-and-conquer strategy by randomly partitioning the array around a pivot. It then recursively processes one of the partitions, based on the position of the pivot relative to the i^{th} smallest element's desired position.
- The randomization aspect ensures that the algorithm, on average, splits the array into relatively equal parts, thereby maintaining linear time complexity on average.

• Expected Time Complexity

- The expected time complexity of Randomized-Select is O(n), where n is the number of elements in the array. This efficiency is attributed to the random selection of the pivot, which, on average, leads to balanced partitions.

Sample Python Code for Randomized-Select

The following Python code demonstrates the implementation of the Randomized-Select algorithm, show-casing its recursive nature and the randomized partitioning technique.

```
import random

def randomized_partition(arr, low, high):
    pivot_index = random.randint(low, high)
    arr[pivot_index], arr[high] = arr[high], arr[pivot_index] # Swap pivot with the end element
    pivot = arr[high]
```

```
i = low - 1
for j in range(low, high):
    if arr[j] <= pivot:
        i += 1
        arr[i+1], arr[j] = arr[j], arr[i]
        arr[i+1], arr[high] = arr[high], arr[i+1]
    return i + 1

def randomized_select(arr, low, high, i):
    if low == high:
        return arr[low]

pivot_index = randomized_partition(arr, low, high)
    k = pivot_index - low + 1  # Number of elements on the left
if i == k: # The pivot value is the answer
    return arr[pivot_index]
elif i < k:
    return randomized_select(arr, low, pivot_index - 1, i)
else:
    return randomized_select(arr, pivot_index + 1, high, i - k)</pre>
```

This implementation of Randomized-Select effectively demonstrates how a randomized pivot selection can be used to select the i^{th} smallest element in an unsorted array in expected linear time.



Red-Black Trees, Augmented Data Structures And Hashtables

Red-Black Trees, Augmented Data Structures And Hashtables

5.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 11.1 Direct-Address Tables
- Chapter 11.2 Hash Tables
- Chapter 11.3 Hash Functions
- Chapter 13.1 Properties Of Red-Black Trees
- Chapter 13.2 Rotations
- Chapter 13.3 Insertion
- Chapter 14.1 Dynamic Order Statistics
- Chapter 14.2 How To Augment A Data Structure

5.0.2 Lectures

The lecture videos for this week are:

- Red Black Trees: Basics ≈ 33 min.
- Red Black Trees: Operations ≈ 30 min.
- Augmenting Trees: Dynamic Order Statistics ≈ 15 min.
- Treaps ≈ 29 min.
- Hashtables: Basics ≈ 25 min.
- Universal Hash Functions ≈ 21 min.

5.0.3 Assignments

The assignment for this week is:

• Problem Set 5 - Red-Black Trees, Augmented Data Structures And Hashtables

5.0.4 Quiz

The quizzes for this week are:

• Quiz 5

5.0.5 Chapter Summary

The first chapter that is being covered this week is **Chapter 11: Hash Tables**. The first section that is covered from this chapter this week is **Section 11.1: Direct-Address Tables**.

Section 11.1: Direct-Address Tables

Direct-Address Tables

Direct-Address Tables represent a straightforward yet efficient method for storing and retrieving data when the universe U of keys is reasonably small. This data structure allows for constant-time operations for many common tasks, such as search, insert, and delete, by directly using the keys as indices in an array.

• Concept and Structure

- In a direct-address table, each position or slot of the table corresponds to a key in the universe of possible keys. If there is an element with key k, it is placed in slot k.
- This one-to-one correspondence between keys and table positions ensures that operations can be performed in O(1) time.

• Operations

- **Search**: To find an element with a specific key k, directly access the slot k in the table.
- **Insert**: To insert an element with key k, place it in slot k.
- **Delete**: To remove an element with key k, clear the slot k.

• Limitations

- The main limitation of direct-address tables is their space complexity. For a large universe of keys, where only a small subset of keys are actually used, direct-address tables can be impractically large.
- They are most effective when the range of key values is not significantly larger than the number of elements to be stored.

Sample Python Code for Direct-Address Tables

The Python code below demonstrates a basic implementation of a direct-address table for storing and retrieving integer keys.

```
class DirectAddressTable:
    def __init__(self, size):
        self.table = [None] * size

def insert(self, key, data):
        self.table[key] = data

def delete(self, key):
        self.table[key] = None

def search(self, key):
    return self.table[key]
```

This example illustrates the simplicity and efficiency of direct-address tables for scenarios where the keys are integers and the universe of keys is small.

The next section that is covered from this chapter this week is **Section 11.2: Hash Tables**.

Section 11.2: Hash Tables

Hash Tables

Hash tables are a powerful data structure used to implement associative arrays, which map keys to values. They achieve high efficiency for searching, insertion, and deletion operations by using a hash function to compute an index into an array of buckets or slots, where the desired value can be stored or found.

• Hash Function

- The hash function takes a key and maps it to an integer, which is used as the index at which the data associated with the key is stored. A well-designed hash function reduces collisions and distributes keys uniformly across the table.

• Handling Collisions

- Chaining: This method involves storing all elements that hash to the same index in a linked list attached to each bucket. While simple, its worst-case search time can degrade to O(n) if many elements hash to the same value.
- Open Addressing: This method finds another slot for the colliding element using strategies like linear probing, quadratic probing, or double hashing. It can suffer from clustering, which affects performance, but maintains a constant O(1) average time complexity for search, insert, and delete under good conditions.

• Load Factor and Resizing

- The load factor, defined as the number of stored entries divided by the number of buckets, affects the performance of the hash table. A high load factor increases the likelihood of collisions.
- Hash tables resize and rehash their contents to a larger array once the load factor exceeds a certain threshold, ensuring that the average time complexities for insertions, deletions, and searches remain at O(1) under the assumption of good hashing.

• Runtime Complexity

- The average-case time complexity for search, insert, and delete operations in a well-implemented hash table is O(1), assuming a good hash function and low load factor.
- In the worst case, particularly with poor hash function choice or high load factor, the time complexity can degrade to O(n), where n is the number of entries in the table. This scenario is more common with chaining when many keys hash to the same index.

Sample Python Code for a Simple Hash Table Using Chaining

This Python code provides a basic implementation of a hash table using chaining to handle collisions, with functions for insertion, searching, and deletion.

```
class HashTable:
    def __init__(self, size):
        self.size = size
        self.table = [[] for _ in range(self.size)]
    def hash_function(self, key):
        return key % self.size
    def insert(self, key, value):
        hash_index = self.hash_function(key)
for i, (k, v) in enumerate(self.table[hash_index]):
             if k == key:
                 self.table[hash_index][i] = (key, value) # Update existing key
        self.table[hash_index].append((key, value))  # Insert new key
    def search(self, key):
        hash index = self.hash function(kev)
        for k, v in self.table[hash_index]:
             if k == key:
        return None
    def delete(self, key):
        hash_index =
                      self.hash_function(key)
        for i, (k, v) in enumerate(self.table[hash_index]):
             if k == key:
                 del self.table[hash_index][i]
                 return
```

This implementation illustrates the basic principles and efficiency of hash tables, highlighting the constant average-case time complexity for key-based operations.

The last section that is covered from this chapter this week is **Section 11.3: Hash Functions**.

Section 11.3: Hash Functions

Hash Functions

Hash functions are a critical component of hash tables, responsible for mapping keys to indices in the table. A good hash function is essential for distributing keys uniformly across the hash table, minimizing collisions, and ensuring efficient data retrieval and storage operations.

Characteristics of a Good Hash Function

- Uniform Distribution: The hash function should distribute keys as evenly as possible over the hash table to minimize collisions.
- **Deterministic**: Given a particular key, the hash function should always produce the same index.
- Efficient to Compute: The hash function should be quick to calculate, allowing for fast data retrieval
 and insertion.
- Minimizes Collisions: While collisions are inevitable, a good hash function should minimize their occurrence.

• Types of Hash Functions

- Division Method: This method involves taking the remainder of the key divided by the table size. It
 is simple and effective for certain key distributions.
- Multiplication Method: The key is multiplied by a constant fractional part, and the fractional part of
 the result is used to determine the index. This method offers good performance for a wide range of key
 values.
- Universal Hashing: A set of hash functions is used, and one is chosen at random for each instance
 of the hash table. This approach reduces the likelihood of collisions and provides better worst-case
 guarantees.

Collision Resolution Techniques

- When two keys hash to the same index, collision resolution techniques such as chaining or open addressing are employed to handle the collision.

• Runtime Complexity

- The efficiency of hash table operations significantly depends on the quality of the hash function. A well-designed hash function can maintain the average-case time complexity of hash table operations at O(1).
- Poor hash function design can lead to clustering or many collisions, degrading performance to O(n) in the worst case.

Considerations in Designing Hash Functions

When designing or choosing a hash function for a particular application, it is crucial to consider the types of keys to be used, the expected distribution of those keys, and the size of the hash table. The goal is to achieve a balance between computational efficiency and the uniform distribution of keys across the table, minimizing the risk of collisions and the subsequent need for collision resolution.

The next chapter that is being covered this week is **Chapter 13: Red-Black Trees**. The first section that is being covered from this chapter this week is **Section 13.1: Properties Of Red-Black Trees**.

Section 13.1: Properties Of Red-Black Trees

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Properties of Red-Black Trees

Red-black trees are a type of self-balancing binary search tree, where each node contains an extra bit for denoting the color of the node, either red or black. This structure ensures that the tree remains approximately balanced, resulting in improved performance for search, insertion, and deletion operations. The properties of red-black trees are crucial for maintaining this balance and guaranteeing a worst-case time complexity of $O(\log{(n)})$ for these operations.

• Properties

- Property 1: Every node is either red or black.
- Property 2: The root is always black.
- Property 3: All leaves (NIL nodes) are black.
- Property 4: If a node is red, then both its children are black. This property prevents the formation of a red node having a red parent, which helps in maintaining the balance of the tree.
- Property 5: For each node, all simple paths from the node to descendant leaves contain the same number of black nodes. This property, also known as the "black-height" property, ensures that the path from the root to the farthest leaf is no more than twice as long as the path from the root to the nearest leaf, maintaining the tree's balanced height.

• Implications of the Properties

- These properties together enforce a critical balance across the tree, ensuring that the longest path from the root to a leaf is no more than twice as long as the shortest path from the root to a leaf. This balance is key to the $O(\log(n))$ search time.
- The requirements for coloration and black-height contribute to a self-balancing nature, where operations such as insertions and deletions may require a series of color changes and rotations to restore red-black properties, but the resulting tree remains balanced.

• Runtime Complexity

- The balancing properties of red-black trees ensure that the height of the tree is always $O(\log(n))$ where n is the number of nodes in the tree. Consequently, search, insertion, and deletion operations all have $O(\log(n))$ worst-case time complexity.

Maintaining Red-Black Tree Properties

Operations on red-black trees, such as insertion and deletion, require careful manipulation to maintain the five properties. This often involves recoloring nodes and performing tree rotations. Despite the complexity of these operations, the payoff is a highly efficient, self-balancing tree ideal for applications requiring frequent insertions and deletions, where balanced search times are critical.

The next section that is being covered from this chapter this week is **Section 13.2: Rotations**.

Section 13.2: Rotations

Rotations in Red-Black Trees

Rotations are fundamental operations in red-black trees, used to maintain the tree's balancing properties after insertions and deletions. These operations are pivotal for preserving the red-black properties, ensuring the tree remains balanced and maintains its $O(\log{(n)})$ search, insertion, and deletion complexities. There are two primary types of rotations: left rotations and right rotations.

• Left Rotation

- A left rotation around a node x involves moving x to its left child's position, while x's right child moves up to x's original position. The left child of x's original right child becomes x's new right child.
- The purpose of a left rotation is to decrease the height of the right subtree and increase the height of the left subtree, helping to maintain the tree's balanced property.

• Right Rotation

- A right rotation is the inverse of a left rotation. It involves moving a node x to its right child's position, while x's left child moves up to x's original position. The right child of x's original left child becomes x's new left child.
- Right rotations are used to decrease the height of the left subtree and increase the height of the right subtree, contributing to the balance of the tree.

• Application of Rotations

- Rotations are used during the insertion and deletion processes in red-black trees. After a new node is
 inserted or a node is deleted, the tree may violate the red-black properties. Rotations, along with color
 changes, are used to restore these properties.
- The choice between performing a left or right rotation depends on the structure of the tree at the point where the red-black properties are violated.

• Runtime Complexity

- Each rotation operation can be performed in O(1) time, as it involves only a fixed number of pointer changes.
- Despite the potential for multiple rotations during an insertion or deletion operation, the overall time complexity of these operations remains $O(\log(n))$, as the height of the tree is logarithmic in the number of nodes.

Significance of Rotations

Rotations are critical for maintaining the balance of red-black trees. By carefully applying these operations, it's possible to ensure that the tree satisfies all red-black properties after each insertion and deletion, preserving the $O(\log(n))$ complexity for fundamental operations and ensuring efficient performance.

The last section that is being covered from this chapter this week is **Section 13.2: Insertion**.

Section 13.2: Insertion

Insertion in Red-Black Trees

Insertion in red-black trees is a process that adds a new node to the tree while maintaining the red-black properties. The insertion process involves placing the new node in the correct location in the binary search tree and then restoring the red-black properties through a series of rotations and recolorings.

• Insertion Steps

- Step 1: Insertion as in a Binary Search Tree (BST): Initially, the new node is inserted using the standard BST insertion procedure. The new node is inserted as a leaf node and colored red.
- Step 2: Restoration of Red-Black Properties: After the initial insertion, the tree may violate the red-black properties. A series of rotations and recolorings are performed to restore these properties. The specific actions depend on the relationship between the newly inserted node, its parent, and its uncle (the sibling of its parent).

• Case Analysis for Restoration

- Case 1: The parent of the newly inserted node is black. In this case, the insertion does not violate any red-black properties, and no further actions are necessary.
- Case 2: The parent and uncle of the newly inserted node are both red. Recolor the parent and uncle to black and the grandparent to red, then continue to fix any further violations up the tree.
- Case 3: The parent is red but the uncle is black, leading to two subcases based on the node's position relative to its parent and grandparent. Rotations and possibly recoloring are used to restore properties.

• Runtime Complexity

- The insertion operation, including the restoration of red-black properties, has a worst-case time complexity of $O(\log(n))$, where n is the number of nodes in the tree. This is because the height of the tree is logarithmic in the number of nodes, and the restoration process involves a constant number of rotations and recolorings.

Importance of Insertion Procedure

The careful design of the insertion procedure ensures that red-black trees maintain their balancing properties, allowing them to provide efficient search, insertion, and deletion operations. The restoration step is crucial for maintaining the tree's balanced height, which is key to achieving $O(\log(n))$ performance for dynamic set operations.

The last chapter that is being covered this **Chapter 14: Augmenting Data Structures**. The first section that is being covered from this chapter this week is **Section 14.1: Dynamic Order Statistics**.

Section 14.1: Dynamic Order Statistics

Dynamic Order Statistics

Dynamic Order Statistics refers to the problem of efficiently maintaining and querying the k^{th} smallest (or largest) element in a dynamically changing set. This set may undergo insertions and deletions of elements, and the goal is to be able to quickly find the k^{th} order statistic after each such modification. Augmented data structures, particularly augmented red-black trees, are commonly used to solve this problem.

• Augmented Red-Black Trees for Dynamic Order Statistics

- To support dynamic order statistics, red-black trees are augmented with additional information. Specifically, each node in the tree stores the size of the subtree rooted at that node (including itself).
- This size attribute allows the tree to support a *Select* operation, which returns the node containing the k^{th} smallest element, and a *Rank* operation, which returns the position of a given element in the linear order of elements maintained by the tree.

• Select Operation

- The Select operation recursively traverses the tree, using the size of the left subtree of each node to determine whether the k^{th} smallest element lies in the left subtree, the right subtree, or is the current node itself.

• Rank Operation

- The *Rank* operation determines the position of a node (the rank) by traversing the path from the node to the root. It adds the sizes of left subtrees plus one (for the node itself) whenever the path moves from a node to its right child.

• Maintaining the Size Attribute

- The size attribute of each node is maintained through insertions and deletions by updating the sizes of all nodes along the path from the inserted or deleted node to the root.

• Runtime Complexity

- Both the *Select* and *Rank* operations can be performed in $O(\log n)$ time on an augmented red-black tree, where n is the number of elements in the tree. This efficiency is due to the balanced nature of red-black trees and the direct access provided by the size attribute.

Significance of Dynamic Order Statistics

Dynamic order statistics are crucial for applications that require efficient, real-time access to order-related information in a dynamic dataset, such as databases, information retrieval systems, and online analytics platforms. The augmentation of red-black trees for this purpose exemplifies a powerful technique for extending the functionality of basic data structures to meet specific computational needs.

The last section that is being covered from this chapter this week is **Section 14.2: How To Augment A Data Structure**.

Section 14.2: How To Augment A Data Structure

How To Augment A Data Structure

Augmenting a data structure involves enhancing a standard data structure with additional information or capabilities to support operations beyond its original scope. This process typically involves adding extra fields to the data structure's nodes and updating these fields during the data structure's modification operations (such as insertions and deletions) to maintain the integrity of the augmented information.

- Steps for Augmenting a Data Structure
 - Identify the Additional Information Needed: Determine what extra information is necessary to support the new operations.
 - Modify the Node Structure: Add fields to the nodes of the data structure to store the additional information.
 - Update the Modification Operations: Modify the insert, delete, and any other operations that alter the data structure to maintain and update the new fields appropriately.
 - Implement the New Operations: Develop the new operations that utilize the augmented information
 to perform their tasks efficiently.
- Example: Augmenting a Binary Search Tree to Support Size Information
 - The goal is to augment a binary search tree (BST) to keep track of the size of each subtree, allowing for efficient calculation of the rank of elements and selection of the k^{th} smallest element.

Sample Python Code for an Augmented BST

This Python code demonstrates how to augment a BST to include size information in each node, supporting efficient size-based selections and rankings.

```
class TreeNode:
    def __init__(self, key, left=None, right=None):
        self.key = key
        self.left = left
        self.right = right
self.size = 1 # S
                        # Size of the subtree rooted at this node
class AugmentedBST:
    def __init__(self):
        self.root = None
    def insert(self, key):
        self.root = self._insert(self.root, key)
        _insert(self, node, key):
        if node is None:
            return TreeNode(key)
        if key < node.key:</pre>
            node.left = self._insert(node.left, key)
        else:
            node.right = self._insert(node.right, key)
        node.size = 1 + (node.left.size if node.left else 0) + (node.right.size if node.right else 0)
        return node
    def select(self, k):
        # Returns the kth smallest element in the BST
        return self._select(self.root, k)
```

```
def _select(self, node, k):

if node is None:

return None

left_size = node.left.size if node.left else 0

if k == left_size + 1:

return node.key

elif k <= left_size:

return self._select(node.left, k)

else:

return self._select(node.right, k - left_size - 1)
```

This augmented BST includes a 'size' attribute in each node, representing the size of the subtree rooted at that node. The 'insert' method updates this attribute to maintain the correct subtree sizes, enabling efficient implementations of operations like 'select', which finds the k^{th} smallest element.



Dynamic Programming

Dynamic Programming

6.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 15.1 Rod Cutting
- Chapter 15.2 Matrix-Chain Multiplication
- Chapter 15.3 Elements Of Dynamic Programming
- Chapter 15.4 Longest Common Subsequence
- Chapter 15.5 Optimal Binary Search Trees

6.0.2 Lectures

The lecture videos for this week are:

- Rod Cutting Problem First Steps ≈ 31 min.
- Rod Cutting Problem: Memoization And Solution Recovery ≈ 24 min.
- Dynamic Programming: Coin Changing Problem ≈ 12 min.
- Knapsack Problem ≈ 29 min.
- Failure of Optimal Substructure ≈ 10 min.
- Longest Common Subsequence Problem ≈ 25 min.

6.0.3 Assignments

The assignment for this week is:

• Problem Set 6 - Dynamic Programming

6.0.4 Quiz

The quizzes for this week are:

• Quiz 6

6.0.5 Chapter Summary

The chapter that is being covered this week is **Chapter 15: Dynamic Programming**. The first section that is being covered this week is **Section 15.1: Rod Cutting**.

Section 15.1: Rod Cutting

Rod Cutting

The Rod Cutting problem is a classic example of dynamic programming, where the goal is to maximize profit from cutting a rod into smaller lengths and selling the pieces based on a given price table. This problem exemplifies the technique of solving complex problems by breaking them down into simpler subproblems, solving each subproblem just once, and storing their solutions - often in an array or hashtable - to avoid the computational cost of solving the same subproblem multiple times.

• Problem Statement

- Given a rod of length n and a table of prices p_i for i = 1, 2, ..., n, where p_i is the price of a rod of length i, determine the maximum revenue r_n obtainable by cutting up the rod and selling the pieces.

Approach

- The problem can be approached recursively, considering the revenue gained by making the first cut at different lengths k, and recursively solving the problem for a rod of length n k.
- To avoid the exponential time complexity due to the recursive structure of the problem, dynamic programming techniques are employed. These include:
 - 1. **Top-Down with Memoization**: Where the algorithm solves the problem by recursively breaking it down into smaller subproblems, caching the result of each subproblem to ensure that each subproblem is solved only once.
 - 2. **Bottom-Up Approach**: Where the algorithm iteratively solves all subproblems from the smallest to the largest, using the solutions of smaller problems to solve larger ones.

• Runtime Complexity

- The naive recursive solution has a runtime complexity of $O(2^n)$, which is significantly reduced to $O(n^2)$ with the use of dynamic programming, either top-down with memoization or bottom-up.

Sample Python Code for Rod Cutting using Dynamic Programming

This Python code snippet demonstrates the bottom-up dynamic programming approach to solve the Rod Cutting problem.

```
def rod_cutting(prices, n):
    # Initialize revenue array
    revenue = [0] * (n + 1)

for j in range(1, n + 1):
    max_rev = float('-inf')
    for i in range(1, j + 1):
        max_rev = max(max_rev, prices[i] + revenue[j - i])
    revenue[j] = max_rev

return revenue[n]
```

In this implementation, 'prices' is a dictionary where the keys are the lengths of the rod pieces and the values are the corresponding prices. The function 'rod_cutting' calculates the maximum revenue that can be obtained for a rod of length 'n' using the bottom-up dynamic programming approach.

The next section that is being covered from this chapter is **Section 15.2: Matrix-Chain Multiplication**.

Section 15.2: Matrix-Chain Multiplication

Matrix-Chain Multiplication

Matrix-Chain Multiplication is a classic optimization problem addressed by dynamic programming. It seeks to find the most efficient way to multiply a sequence of matrices. The challenge lies in the fact that matrix multiplication is associative, meaning the order in which the multiplications are performed can significantly affect the total number of scalar operations required, without changing the product. The goal is to determine the optimal parenthesization of the matrix product that minimizes the number of scalar multiplications.

• Problem Statement

- Given a chain $\langle A_1, A_2, ..., A_n \rangle$ of n matrices, where for i = 1, 2, ..., n, matrix A_i has dimension $p_{i-1} \times p_i$, fully parenthesize the product $A_1 A_2 ... A_n$ in a way that minimizes the number of scalar multiplications.

• Dynamic Programming Approach

- The solution involves a bottom-up approach that constructs a table m[i,j] for the minimum number of scalar multiplications needed to compute the matrix $A_iA_{i+1}...A_j$, for $1 \le i, j \le n$. The optimal solution to the entire problem is then found in m[1,n].
- The algorithm also maintains a table s[i, j] to record the index k at which the optimal split of the product sequence occurs, facilitating the construction of the optimal parenthesization.

• Runtime Complexity

– The time complexity of the dynamic programming solution for the Matrix-Chain Multiplication problem is $O(n^3)$, where n is the number of matrices in the chain. The space complexity for storing the m and s tables is $O(n^2)$.

Sample Python Code for Matrix-Chain Multiplication

The following Python code demonstrates the dynamic programming solution to the Matrix-Chain Multiplication problem.

```
def matrix_chain_order(p):
    n = len(p) - 1
    m = [[0 for _ in range(n)] for _ in range(n)]
    s = [[0 for _ in range(n)] for _ in range(n)]

for length in range(2, n + 1):
    for i in range(n - length + 1):
        j = i + length - 1
        m[i][j] = float('inf')
    for k in range(i, j):
        q = m[i][k] + m[k + 1][j] + p[i] * p[k + 1] * p[j + 1]
        if q < m[i][j]:
        m[i][j] = q
        s[i][j] = k

return m, s

def print_optimal_parens(s, i, j):
    if i = j:
        print(f"A{i+1}", end="")
    else:
        print_optimal_parens(s, i, s[i][j])
        print_optimal_parens(s, s[i][j] + 1, j)
        print_optimal_parens(s, s[i][j] + 1, j)
        print(")", end="")</pre>
```

This implementation calculates the minimum number of scalar multiplications needed to multiply a chain of matrices and prints the optimal way to parenthesize the product. The 'matrix_chain_order' function builds the m and s tables, and the 'print_optimal_parens' function prints the optimal parenthesization.

The next section that is being covered from this chapter this week is **Section 15.3: Elements Of Dynamic Programming**.

Section 15.3: Elements Of Dynamic Programming

Elements of Dynamic Programming

Dynamic Programming (DP) is a method for solving complex problems by breaking them down into simpler subproblems. It is particularly useful for problems exhibiting the properties of overlapping subproblems and optimal substructure. Understanding the key elements of dynamic programming is crucial for effectively applying it to a wide range of problems.

• Overlapping Subproblems

This property indicates that the space of subproblems is small, meaning that the same subproblems are solved repeatedly during the computation. Dynamic programming exploits this by solving each subproblem only once and storing its solution in a table, thereby avoiding the work of recomputing the answer every time the subproblem is encountered.

• Optimal Substructure

- A problem exhibits optimal substructure if an optimal solution to the problem contains within it optimal solutions to the subproblems. This property allows a problem to be solved by combining the solutions of its subproblems. Dynamic programming algorithms typically build up solutions to larger and larger subproblems using the solutions to smaller subproblems.

• Memoization vs. Tabulation

- **Memoization** (Top-Down Approach): This technique involves writing the recursive algorithm and using a table to store the results (memo) of subproblems to avoid recomputing them. The algorithm is typically expressed in a natural manner but modified to save the results of intermediate subproblems.
- Tabulation (Bottom-Up Approach): This technique involves filling up a DP table by solving subproblems in a specific order, ensuring that all subproblems needed to solve a problem are solved first. This approach iteratively builds up the solution to larger problems based on the solutions to smaller problems.

• Constructing Solutions

- Beyond calculating the optimal value that a dynamic programming solution seeks, many problems also require constructing the solution itself. This typically involves additional bookkeeping to trace back the decisions that led to the optimal value, often through a separate table or by augmenting the table used to calculate the optimal values.

• Runtime Complexity

The time complexity of a dynamic programming algorithm is typically determined by the number of subproblems multiplied by the time taken to solve each subproblem. Since each subproblem is solved only once and stored for future reference, dynamic programming can significantly reduce the time complexity from exponential to polynomial in many cases.

Applying Dynamic Programming

Successfully applying dynamic programming involves identifying that the problem has overlapping subproblems and optimal substructure, choosing between memoization and tabulation based on the problem and personal preference, and carefully defining the structure of the table(s) used for storing the solutions of subproblems. Understanding these elements is fundamental to designing efficient dynamic programming solutions.

The next section that is being covered from this chapter this week is **Section 15.4: Longest Common Subsequence**.

Section 15.4: Longest Common Subsequence

Longest Common Subsequence

The Longest Common Subsequence (LCS) problem is a classic example in the field of dynamic programming and computer science, focusing on finding the longest subsequence present in two sequences (not necessarily contiguous) in the same order. This problem is fundamental in the domains of bioinformatics, text processing, and diff tools, among others.

• Problem Statement

- Given two sequences $X = \{x_1, x_2, ..., x_m\}$ and $Y = \{y_1, y_2, ..., y_n\}$, find the length of their longest common subsequence, which is the longest sequence that can be derived from both X and Y by deleting some items without changing the order of the remaining elements.

• Dynamic Programming Formulation

- The problem can be solved by considering smaller instances of the problem, namely finding the LCS of prefixes of X and Y. Let c[i,j] be the length of the LCS of X_i and Y_j , where X_i and Y_j are the first i and j elements of X and Y, respectively.
- The solution is built up using the following recurrence:

$$c[i,j] = \begin{cases} 0 & \text{if } i = 0 \text{ or } j = 0\\ c[i-1,j-1] + 1 & \text{if } i,j > 0 \text{ and } x_i = y_j\\ \max(c[i,j-1],c[i-1,j]) & \text{if } i,j > 0 \text{ and } x_i \neq y_j \end{cases}$$

 This approach ensures that each subproblem is solved only once, with the solution being stored in a table from which the solution to the problem can be constructed.

• Runtime Complexity

- The runtime complexity of solving the LCS problem using dynamic programming is O(mn), where m and n are the lengths of the two sequences. This complexity arises from filling out a table of size $m \times n$ based on the recurrence relation.

Sample Python Code for Longest Common Subsequence

This Python code snippet demonstrates the dynamic programming solution to the LCS problem.

The 'lcs' function computes the length of the longest common subsequence between two strings 'X' and 'Y', illustrating the application of dynamic programming to efficiently solve this problem.

The next section that is being covered from this chapter this week is **Section 15.5: Optimal Binary Search Trees**.

Section 15.5: Optimal Binary Search Trees

Optimal Binary Search Trees

Optimal Binary Search Trees (BSTs) are a specialized form of binary search trees that are structured to minimize the cost of searching for a set of keys. The cost is typically measured in terms of the number of comparisons needed to find a key in the tree. The concept of optimality here relates to the minimization of the expected search cost, given a set of keys with known probabilities of access.

• Problem Statement

Given a set of n keys with their probabilities of being searched for, construct a binary search tree that
has the minimum expected search cost. The search cost of a key is proportional to the depth of the key
in the tree.

• Dynamic Programming Approach

— The solution to constructing an optimal BST involves a dynamic programming approach that examines all possible trees and selects the one with the lowest cost. This includes not only the given keys but also the probabilities of searching for values between these keys (often treated as "dummy keys" representing searches for values not in the set). - The algorithm calculates the cost of each subtree and uses these costs to construct the optimal solution for larger trees. It maintains a table e[i,j] for the cost of the optimal BST for keys k_i through k_j , and a table root[i,j] to store the root of the optimal subtree for k_i through k_j .

• Runtime Complexity

- The time complexity of constructing an optimal BST using dynamic programming is $O(n^3)$, where n is the number of keys. This complexity arises from the need to examine each possible subtree for each interval of keys.

Sample Python Code for Optimal Binary Search Trees

The following Python code snippet provides an implementation of the dynamic programming approach to construct optimal binary search trees.

```
def optimal_bst(keys, p, q, n):
    e = [[0 for j in range(n + 1)] for i in range(n + 2)]
    w = [[0 for j in range(n + 1)] for i in range(n + 2)]
    root = [[0 for j in range(n + 1)] for i in range(n + 1)]

for i in range(1, n + 2):
    e[i][i - 1] = q[i - 1]
    w[i][i - 1] = q[i - 1]

for l in range(1, n + 1):
    for i in range(1, n - 1 + 2):
        j = i + l - 1
        e[i][j] = float('inf')
    w[i][j] = w[i][j - 1] + p[j] + q[j]

for r in range(i, j + 1):
        t = e[i][r - 1] + e[r + 1][j] + w[i][j]

if t < e[i][j]:
    e[i][j] = t
    root[i][j] = r

return e, root</pre>
```

This implementation calculates the cost and structure of the optimal BST for a given set of keys and their search probabilities. The 'optimal_bst' function computes the cost e and the root table, which can be used to construct the tree itself.

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Exam 2

Exam 2

7.0.1 Exam

The exam for this week is:

- Spot Exam 2 Notes
- Spot Exam 2



Greedy Algorithms

Greedy Algorithms

8.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 16.1 An Activity-Selection Problem
- Chapter 16.2 Elements Of The Greedy Strategy
- Chapter 16.3 Huffman Codes

8.0.2 Lectures

The lecture videos for this week are:

- Introduction To Greed Algorithms ≈ 20 min.
- Greedy Interval Scheduling ≈ 23 min.
- Lossless Data Compression: PrefixCodes ≈ 30 min.
- Huffman Codes ≈ 28 min.

8.0.3 Assignments

The assignment for this week is:

• Programming Assignment 2 - Tries

8.0.4 Quiz

The quizzes for this week are:

• Quiz 7

8.0.5 Chapter Summary

The chapter that is being covered this week is **Chapter 16: Greedy Algorithms**. The first section that is being covered from this chapter this week is **Section 16.1: An Activity-Selection Problem**.

Section 16.1: An Activity-Selection Problem

An Activity-Selection Problem

The activity-selection problem is a classic example illustrating the power of greedy algorithms. The problem involves selecting the maximum number of activities that don't overlap from a set of activities, each with a start time and an end time. The goal is to make the optimal selection such that the maximum number of activities can be attended.

• Problem Statement

- Given n activities with their start times s_i and finish times f_i for each activity i, select the maximum number of activities that do not overlap. An activity i is said to overlap with activity j if their time intervals s_i , f_i and s_j , f_j intersect.

• Greedy-Choice Property

- The greedy choice is to always select the activity that finishes first. This choice is made because selecting an activity that finishes earlier leaves as much room as possible for the remaining activities.

• Greedy Algorithm Strategy

- Sort all activities by their finishing times.
- Select the first activity in the sorted list and mark it as the current activity.
- For each remaining activity in the list, if the start time of this activity is greater than or equal to the finish time of the current activity, select this activity and update the current activity to this one.

• Runtime Complexity

- The runtime complexity of the activity-selection problem using a greedy algorithm mainly depends on the sorting algorithm used. Assuming a sorting algorithm with $O(n \log n)$ complexity, the overall complexity of the activity-selection algorithm is $O(n \log n)$ due to the initial sorting step. The selection process itself is O(n), as it requires a single pass through the sorted list of activities.

Sample Python Code for Activity Selection Problem

This Python code snippet demonstrates the greedy algorithm solution to the activity-selection problem.

```
def activity_selection(start_times, finish_times):
    activities = sorted(zip(start_times, finish_times), key=lambda x: x[1])
    n = len(activities)
    selected = [activities[0]] # Select the first activity
    for i in range(1, n):
        if activities[i][0] >= selected[-1][1]: # If this activity doesn't overlap with the last selected
        selected.append(activities[i]) # Select it
    return selected
```

The 'activity_selection' function takes the start and finish times of the activities, sorts them by their finish times, and iteratively selects the maximum number of non-overlapping activities. This implementation highlights the effectiveness and simplicity of the greedy approach for solving the activity-selection problem.

The next section that is being covered from this chapter this week is **Section 16.2: Elements of The Greedy Strategy**.

Section 16.2: Elements of The Greedy Strategy

Elements of The Greedy Strategy with a Python Example

The Greedy Strategy is a method for solving optimization problems by building a solution piece by piece, choosing the next piece with the most immediate benefit. Below, we detail the elements of the Greedy Strategy and illustrate them with a Python example for the problem of making change.

• Candidate Set

- The set of coins available to make change. For the making change problem, this set includes all denominations of coins available.

• Selection Function

 Chooses the largest denomination of coin that does not exceed the remaining amount of change to be given. This choice maximizes the immediate reduction in remaining change.

• Feasibility Function

- Ensures the selected coin does not exceed the amount of change remaining. This function maintains the constraint of not giving too much change.

• Objective Function

- Minimizes the number of coins given as change. The algorithm aims to reduce the total count of coins.

• Solution Function

- Determines if the total amount of change has been given. The algorithm terminates when the remaining change to be given is zero.

Sample Python Code for Making Change

This Python code snippet demonstrates the greedy algorithm solution to the making change problem.

```
def make_change(amount, denominations):
    denominations.sort(reverse=True) # Sort denominations in descending order
    result = []
    for coin in denominations:
        while amount >= coin:
        amount -= coin
        result.append(coin)
    return result
```

The 'make_change' function takes an amount and a list of coin denominations, then iteratively selects the largest denomination that does not exceed the remaining amount. This implementation showcases the Greedy Strategy elements at work in solving the making change problem.

Each element of the Greedy Strategy plays a critical role in the operation of the algorithm, guiding the selection of coins in a way that aims to minimize the total number of coins given, thereby achieving an efficient solution to the making change problem.

The last section that will be covered from this chapter this week is **Section 16.3: Huffman Codes**.

Section 16.3: Huffman Codes

Huffman Codes

Huffman codes are a type of prefix coding used for lossless data compression. They assign variable-length codes to input characters, with shorter codes assigned to more frequent characters, thereby achieving compression.

A Huffman tree is a binary tree used to encode characters. It is constructed based on the frequency of characters in the input data. Characters with higher frequencies are assigned shorter codes, while those with lower frequencies are assigned longer codes. The Huffman tree is built in a bottom-up manner, starting with individual characters as leaves and combining them to form higher nodes until the entire tree is formed.

Building A Huffman Tree

- Create a leaf node for each unique character in the input data, with the frequency of occurrence as the weight
 of the node.
- Place all leaf nodes in a priority queue, ordered by their frequencies.
- While there is more than one node in the priority queue:
 - Remove the two nodes with the lowest frequencies from the queue.
 - Create a new internal node with these two nodes as children. The frequency of the new node is the sum of the frequencies of its children.
 - Insert the new node back into the priority queue.
- The remaining node in the priority queue is the root of the Huffman tree.

Decoding A Huffman Tree

- Start at the root of the Huffman tree.
- For each bit in the encoded data:
 - If the bit is 0, move to the left child of the current node.
 - If the bit is 1, move to the right child of the current node.
 - If a leaf node is reached, output the corresponding character and return to the root of the tree.

Python Example: Huffman Coding

Below is an example how to use huffman coding in Python.

```
from heapq import heappop, heappush, heapify
from collections import defaultdict
    class Node:
         def __init__(self, char, freq):
              self.char = char
self.freq = freq
self.left = None
              self.right = None
    def build_huffman_tree(text):
         freq_map = defaultdict(int)
for char in text:
             freq_map[char] += 1
         priority_queue = [Node(char, freq) for char, freq in freq_map.items()]
         heapify(priority_queue)
         while len(priority_queue) > 1:
             left = heappop(priority_queue)
              right = heappop(priority_queue)
              internal_node = Node(None, left.freq + right.freq)
              internal_node.left = left
              internal_node.right = right
              heappush(priority_queue, internal_node)
         return priority_queue[0]
    def encode_huffman(root, code_dict, current_code=""):
30
         if root is not None:
             if root.char is not None:
    code_dict[root.char] = current_code
encode_huffman(root.left, code_dict, current_code + "0")
              encode_huffman(root.right, code_dict, current_code + "1")
36
37
    def huffman_encode(text):
         root = build_huffman_tree(text)
         code_dict = {}
         encode_huffman(root, code_dict)
         encoded_text = ''.join(code_dict[char] for char in text)
         return encoded_text, root
42
43
    def huffman_decode(encoded_text, root):
44
         decoded_text =
         current_node = root
         for bit in encoded_text:
             if bit == "0"
                  current_node = current_node.left
                  current_node = current_node.right
              if current_node.char is not None:
                 decoded_text += current_node.char
                  current_node = root
         return decoded_text
```

This Python code demonstrates the construction of a Huffman tree from input text, encoding of the text using Huffman coding, and decoding of the encoded text back to its original form.

Graphs And Basic Algorithms On Graphs

Graphs And Basic Algorithms On Graphs

9.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 22.1 Representation Of Graphs
- Chapter 22.2 Breadth-First Search
- Chapter 22.3 Depth-First Search
- Chapter 22.4 Topological Sort

9.0.2 Lectures

The lecture videos for this week are:

- Introduction To Graphs ≈ 14 min.
- Breadth First Search ≈ 17 min.
- Depth First Search ≈ 33 min.
- Topological Sorting ≈ 11 min.

9.0.3 Assignments

The assignment for this week is:

• Problem Set 7 - Graphs

9.0.4 Quiz

The quizzes for this week are:

• Quiz 8

9.0.5 Chapter Summary

The chapter that is being covered this week is **Chapter 22: Elementary Graph Algorithms**. The first section that is being covered from this chapter this week is **Section 22.1: Representations Of Graphs**.

Section 22.1: Representations Of Graphs

Representations of Graphs

Graphs can be represented in various ways, each with its own advantages and disadvantages. The two most common representations are adjacency lists and adjacency matrices. These representations are crucial for designing and analyzing algorithms that operate on graphs.

Adjacency Lists

An adjacency list represents a graph as an array of lists. Each list corresponds to a vertex in the graph and contains all vertices adjacent to it. This representation is efficient in terms of space, especially for sparse graphs where the number of edges is much less than the square of the number of vertices.

- Advantages: Space-efficient for sparse graphs; easy to iterate over the neighbors of a vertex.
- **Disadvantages:** Checking whether an edge exists between two vertices can be less efficient than with an adjacency matrix.

Adjacency Matrices

An adjacency matrix represents a graph as a 2D array of size $V \times V$ where V is the number of vertices in the graph. Each cell at position (i, j) in the matrix indicates whether there is an edge from vertex i to vertex j, typically with a 1 for the presence of an edge and a 0 for its absence.

- Advantages: Efficient for checking the existence of an edge between any two vertices; easy to add or remove edges.
- **Disadvantages:** Space-inefficient for sparse graphs, as it requires storing V^2 elements regardless of the number of edges.

Choosing the Right Representation

The choice between adjacency lists and matrices largely depends on the specific requirements of the algorithm and the nature of the graph being dealt with. Adjacency lists are generally preferred for sparse graphs due to their space efficiency, while adjacency matrices may be more suitable for dense graphs or when frequent edge existence checks are necessary.

The next section that is being covered from this chapter this week is Section 22.2: Breadth-First Search.

Section 22.2: Breadth-First Search

Breadth-first Search (BFS)

Breadth-first Search (BFS) is an algorithm for traversing or searching tree or graph data structures. It starts at the tree root (or some arbitrary node of a graph, sometimes referred to as a 'search key') and explores the neighbor nodes at the present depth prior to moving on to the nodes at the next depth level.

Algorithm Overview:

- BFS explores a graph in the broadest manner by visiting all neighbors of a starting vertex before moving to their children.
- It uses a queue to keep track of the next location to visit.
- BFS can be used to find the shortest path between two nodes in an unweighted graph.

Steps:

- 1. Start by putting any one of the graph's vertices at the back of a queue.
- 2. Take the front item of the queue and add it to the visited list.
- 3. Create a list of that vertex's adjacent nodes. Add the ones which aren't in the visited list to the back of the queue.
- 4. Keep repeating steps 2 and 3 until the queue is empty.

Applications:

- Finding shortest paths in unweighted graphs.
- Crawling web pages from a given URL.
- Social networking applications where you want to find people within a certain distance 'k' from a person.

Python Example: Breadth-first Search

Below is a simple Python example of BFS algorithm implementation to traverse graphs.

```
from collections import deque

def bfs(graph, start):
    visited = set()
    queue = deque([start])

while queue:
    vertex = queue.popleft()
    if vertex not in visited:
        visited.add(vertex)
    print(vertex, end=" ")

# Add all unvisited neighbours to the queue.
queue.extend(neighbour for neighbour in graph[vertex] if neighbour not in visited)
```

This Python function demonstrates performing a BFS on a graph represented as an adjacency list. Starting from a given node, it visits all nodes of the graph in the breadth-first manner.

The next section that is being covered from this chapter this week is Section 22.3: Depth-First Search.

Section 22.3: Depth-First Search

Depth-first Search (DFS)

Depth-first Search (DFS) is a fundamental algorithm used in graph theory to explore vertices and edges of a graph. It emphasizes diving as deep as possible into the graph's branches before backtracking. This strategy makes DFS an excellent tool for exploring maze structures, solving puzzles, and analyzing networks.

Algorithm Overview:

- DFS explores a graph by starting at the root (or any arbitrary node) and explores as far as possible along each branch before backtracking.
- It uses a stack (either the function call stack through recursion or an explicit stack data structure) to keep track of the path it's exploring.

Steps:

- 1. Mark the current vertex as visited and print or record it.
- 2. For each adjacent vertex connected to the current vertex, if it has not been visited, recursively visit that vertex.
- 3. Continue until all vertices that are reachable from the original source vertex have been explored.
- 4. If there are still unvisited vertices, select one as a new source and repeat the process, ensuring all vertices in the graph are explored.

Applications:

- Pathfinding and checking for connected components in a graph.
- Topological sorting.
- Finding strongly connected components.

Python Example: Depth-first Search

Below is a Python example demonstrating the DFS algorithm.

This Python function showcases how to perform a DFS on a graph represented as an adjacency list. It emphasizes a recursive approach to traverse all vertices of the graph, marking each as visited and exploring as far as possible before backtracking.

The last section that is being covered from this chapter this week is Section 22.4: Topological Sort.

Section 22.4: Topological Sort

Topological Sort

Topological sort is an algorithm for linearly ordering the vertices of a directed acyclic graph (DAG) such that for every directed edge uv from vertex u to vertex v, u comes before v in the ordering.

Algorithm Overview:

- Topological sorting is only possible if the graph has no directed cycles, i.e., it must be a DAG.
- A common approach to find a topological sort is to use Depth-first Search (DFS).

Steps:

- 1. Call DFS to compute the finishing times for each vertex.
- 2. As each vertex is finished, insert it onto the front of a linked list.
- 3. After all vertices have been processed, return the linked list of vertices.

Applications:

- Scheduling tasks under precedence constraints.
- Course scheduling.
- Determining the order of compilation tasks in programming projects.

Python Example: Topological Sort

Below is an example of how to implement topological sorting in Python using DFS.

```
def dfs(graph, node, visited, stack):
visited.add(node)
for neighbour in graph[node]:
if neighbour not in visited:
dfs(graph, neighbour, visited, stack)
stack.insert(0, node)
```

This Python function demonstrates a DFS-based approach to achieve a topological sort of a directed acyclic graph. The vertices are pushed onto a stack in the reverse order of their finishing times, ensuring that all directed edges point from a vertex earlier in the order to a vertex later in the order.



Exam 3 Study Week

Exam 3 Study Week

Study for Exam 3.



Strongly Connected Components and Spanning Trees

Strongly Connected Components and Spanning Trees

11.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 22.5 Strongly Connected Components
- Chapter 23.1 Growing A Minimum Spanning Tree
- Chapter 23.2 The Algorithms Of Kurskal And Prim

11.0.2 Lectures

The lecture videos for this week are:

- Strongly Connected Components: Introduction ≈ 15 min.
- Strongly Connected Components: Properties ≈ 16 min.
- Strongly Connected Components: Algorithms ≈ 16 min.
- Minimal Spanning Trees: Introduction ≈ 26 min.
- Minimal Spanning Trees: Prim's Algorithm ≈ 37 min.
- Minimal Spanning Tree: Kruskal's Algorithm ≈ 8 min.

11.0.3 Assignments

The assignment for this week is:

• Problem Set 8 - Strongly Connected Components and Spanning Trees

11.0.4 Quiz

The quizzes for this week are:

• Quiz 9

11.0.5 Exam

The exam for this week is:

- Spot Exam 3 Notes
- Spot Exam 3

11.0.6 Chapter Summary

The first chapter that is being covered this week is **Chapter 22: Elementary Graph Algorithms**. The section that is being covered from this chapter this week is **Section 22.5: Strongly Connected Components**.

Section 22.5: Strongly Connected Components

Strongly Connected Components (SCCs)

Strongly Connected Components (SCCs) refer to maximal subgraphs of a directed graph, where every vertex is reachable from every other vertex within the same subgraph. In essence, if there is a path from any vertex in a component to any other vertex in the same component, then that component is strongly connected.

Algorithm Overview:

- The concept of SCCs is fundamental in the analysis of directed graphs.
- Kosaraju's algorithm and Tarjan's algorithm are two well-known algorithms for finding SCCs in a directed graph.
- Both algorithms involve depth-first search (DFS) but differ in their approaches to processing the vertices.

Steps

Kosaraju's algorithm can be summarized in the following steps:

- 1. Perform a DFS on the original graph, recording the finish times for each vertex.
- 2. Compute the transpose of the graph (reverse the direction of every edge).
- 3. Perform a DFS on the transposed graph, in the order of decreasing finish times from the first DFS.
- 4. Each DFS in the transposed graph gives a strongly connected component.

Tarjan's algorithm, on the other hand, keeps track of low-link values and uses a stack to manage components during the DFS.

Applications:

- SCCs are used in theoretical computer science to understand the structure of complex directed graphs.
- They can identify closed loops or cycles within networks, which is useful in networking, the analysis of social networks, and in the study of biological systems.
- Understanding SCCs can also aid in optimizing navigational or routing strategies in directed graphs.

Python Example: Finding SCCs

Below is a simplified example of applying Kosaraju's algorithm to find SCCs in a directed graph.

```
def kosaraju(graph):
    stack = []
    visited = set()
    scc = []
    def dfs(vertex):
        if vertex not in visited:
             visited.add(vertex)
             for neighbour in graph[vertex]:
                 dfs(neighbour)
             stack.append(vertex)
    def reverse(graph):
        reversed_graph = {v: [] for v in graph}
        for vertex, edges in graph.items():
            for edge in edges:
    reversed_graph[edge].append(vertex)
        return reversed_graph
    def dfs_reverse(vertex, temp):
        if vertex not in visited:
             visited.add(vertex)
             temp.append(vertex)
             for neighbour in reversed_graph[vertex]:
                 dfs_reverse(neighbour, temp)
        vertex in graph:
        dfs(vertex)
    reversed graph = reverse(graph)
    visited.clear()
    while stack:
        vertex = stack.pop()
        if vertex not in visited:
    temp = []
             dfs reverse (vertex, temp)
```

```
38 scc.append(temp)
39
40 return scc
41
```

This function demonstrates the process of identifying SCCs in a directed graph using Kosaraju's algorithm, utilizing DFS, graph transposition, and stack operations.

The next chapter that is being covered this week is **Chapter 23: Minimum Spanning Trees** and the first section that is being a minimum covered is **Section 23.1: Growing A Minimum Spanning Tree**.

Section 23.1: Growing A Minimum Spanning Tree

Growing A Minimum Spanning Tree

A Minimum Spanning Tree (MST) of a weighted, undirected graph is a subset of the edges that connects all vertices together, without any cycles, and with the minimal possible total edge weight. The process of growing a minimum spanning tree involves incrementally adding edges to construct the MST, ensuring that the selected edges at each step contribute to the minimum overall weight without forming a cycle.

Algorithm Overview:

- Prim's and Kruskal's algorithms are two classic approaches for growing a minimum spanning tree.
- Prim's algorithm builds the MST by starting with an arbitrary vertex and adding the cheapest edge from the tree to a vertex not yet in the tree at each step.
- Kruskal's algorithm builds the MST by sorting all the edges by their weights and adding them one by one, skipping those that would form a cycle.
- Both algorithms make use of greedy strategies to ensure that at each step, the choice made contributes to an overall optimal solution.

Steps:

For Prim's algorithm, the steps can be outlined as follows:

- 1. Initialize a tree with a single vertex, chosen arbitrarily from the graph.
- 2. Grow the tree by adding the cheapest edge from the tree to any vertex not yet in the tree.
- 3. Repeat step 2 until all vertices are included in the tree.

For Kruskal's algorithm, the steps are:

- 1. Sort all the edges in non-decreasing order of their weight.
- 2. Pick the smallest edge. Check if it forms a cycle with the spanning tree formed so far. If cycle is not formed, include this edge. Otherwise, discard it.
- 3. Repeat step 2 until there are V-1 edges in the tree, where V is the number of vertices in the graph.

Applications:

- Designing minimum cost networks like electrical grids, computer networks, and road networks.
- Approximating solutions for NP-hard problems, such as the traveling salesman problem.
- Cluster analysis in data science.

Python Example: Growing a Minimum Spanning Tree

Below is a simple implementation of Prim's algorithm for growing a minimum spanning tree.

This function uses Prim's algorithm to grow a minimum spanning tree from a given start vertex. The graph is represented as a dictionary of dictionaries, where the outer dictionary holds vertices, and each inner dictionary maps adjacent vertices to edge weights.

The last section that is being covered from this chapter this week is Section 23.2: The Algorithms Of Kruskal And Prim.

Section 23.2: The Algorithms Of Kruskal And Prim

The Algorithms of Kruskal and Prim

Kruskal's and Prim's algorithms are two efficient methods used to find a Minimum Spanning Tree (MST) of a connected, edge-weighted graph. The MST is a subset of the edges that connects all the vertices together, without any cycles, and with the minimal total edge weight.

Kruskal's Algorithm:

Kruskal's algorithm is a greedy algorithm that builds the MST by selecting the shortest edge that does not form a cycle and adding it to the growing spanning tree. The process is repeated until the tree spans all the vertices.

- 1. Sort all edges in the graph in increasing order of their weights.
- 2. Initialize a forest (a set of trees), where each vertex in the graph is a separate tree.
- 3. Repeat the following until the forest forms a single tree (the MST): Choose the smallest edge from the sorted edge list. If the edge connects two different trees, add it to the MST (and merge the two trees). Otherwise, discard the edge.
- 4. The resulting MST is the union of all selected edges.

Prim's Algorithm:

Prim's algorithm, another greedy approach, starts with a single vertex and grows the MST by adding the cheapest possible connection from the tree to another vertex not in the tree.

- 1. Start with a tree containing a single chosen vertex from the graph.
- 2. At each step, add the cheapest edge that connects a vertex in the tree to a vertex outside the tree.
- 3. Repeat step 2 until all vertices are included in the tree.

Applications:

Both algorithms are used in various applications, including:

- Designing network layouts (e.g., computer networks, telecommunications networks).
- Planning road systems to minimize cost.
- Solving certain approximation problems in algorithms.
- Generating mazes for computer games.

Comparison

While both Kruskal's and Prim's algorithms are efficient for finding the MST, their performance may vary depending on the structure of the graph and the specific use case. Kruskal's algorithm is generally simpler to implement and can be more efficient for sparse graphs. Prim's algorithm can be faster for dense graphs, especially when implemented with a priority queue.



Shortest Path Algorithms

Shortest Path Algorithms

12.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 24.1 The Bellman-Ford Algorithm
- Chapter 24.2 Single-Source Shortest Paths In Directed Acyclic Graphs
- Chapter 24.3 Dijkstra's Algorithm
- Chapter 24.4 Difference Constrains And Shortest Paths
- Chapter 24.5 Proofs Of Shortest-Paths Properties
- Chapter 25.1 Shortest Paths And Matrix Multiplication
- Chapter 25.2 The Floyd-Warshall Algorithm

12.0.2 Lectures

The lecture videos for this week are:

- Shortest Path Problems: Introduction ≈ 30 min.
- Bellman Ford Algorithm ≈ 19 min.
- Bellman Ford: Example And Analysis ≈ 27 min.
- Shortest Path Over DAGs ≈ 12 min.
- Dijkstra's Algorithm ≈ 20 min.
- Dijkstra's Algorithm Analysis ≈ 12 min.
- All Pairs Shortest Path ≈ 35 min.

12.0.3 Assignments

The assignment for this week is:

- Problem Set 9 Shortest Path Algorithms
- Programming Assignment 3 Shortest Path

12.0.4 Quiz

The quizzes for this week are:

• Quiz 10

12.0.5 Chapter Summary

The first chapter that is being covered this week is **Chapter 24: Single-Source Shortest Path Algorithms**. The first section that is being covered from this chapter this week is **Section 24.1: The Bellman-Ford Algorithm**.

Section 24.1: The Bellman-Ford Algorithm

The Bellman-Ford Algorithm

The Bellman-Ford algorithm is a graph search algorithm that calculates the shortest paths from a single source vertex to all other vertices in a weighted graph. It is capable of handling graphs with negative weight edges, unlike Dijkstra's algorithm, which only works with graphs that have non-negative edge weights. The Bellman-Ford algorithm can also detect negative cycles in a graph.

Algorithm Overview:

- The algorithm initializes the distance to the source vertex to 0 and all other vertices to infinity.
- It relaxes all the edges in the graph for (V-1) iterations, where V is the number of vertices. Relaxation means updating the distance to a vertex if a shorter path is found.
- After (V-1) iterations, it checks for negative weight cycles by trying to relax the edges once more. If a shorter path is found, a negative cycle exists.

Steps:

The Bellman-Ford algorithm can be summarized in the following steps:

- 1. Initialize distances from the source to all vertices as infinite and distance to the source itself as 0.
- 2. For each vertex, apply relaxation for all the edges.
- 3. Repeat step 2 for (V-1) times.
- 4. Check for negative-weight cycles by performing step 2 one more time. If a distance is updated, then report that a negative cycle exists.

Applications:

- The Bellman-Ford algorithm is used in network routing protocols, such as the Distance Vector Routing Protocol.
- It is beneficial for calculating shortest paths in graphs with negative edges or for detecting negative cycles.
- The algorithm is also useful in financial applications to detect arbitrage opportunities.

Python Example: The Bellman-Ford Algorithm

Below is a Python implementation of the Bellman-Ford algorithm.

```
bellman ford(graph, source):
distance = {vertex: float('infinity') for vertex in graph}
predecessor = {vertex: None for vertex in graph}
distance[source] = 0
      in range(len(graph) - 1):
    for vertex in graph:
        for neighbour, weight in graph[vertex].items():
             if distance[vertex] + weight < distance[neighbour]:
distance[neighbour] = distance[vertex] + weight
                 predecessor[neighbour] = vertex
# Check for negative weight cycles
for vertex in graph:
    for neighbour, weight in graph[vertex].items():
            distance[vertex] + weight < distance[neighbour]:</pre>
             print("Graph contains a negative-weight cycle")
             return None
return distance, predecessor
```

This function calculates the shortest paths from a single source to all other vertices and detects negative weight cycles if any exist.

The next topic that is being covered from this chapter this week is **Section 24.2: Single-Source Shortest**Paths In Directed Acyclic Graphs.

Section 24.2: Single-Source Shortest Paths In Directed Acyclic Graphs

Single-Source Shortest Paths in Directed Acyclic Graphs (DAGs)

Finding single-source shortest paths in Directed Acyclic Graphs (DAGs) involves calculating the shortest path from a given source vertex to all other vertices in a DAG. Since DAGs do not contain cycles, they allow for more efficient shortest path algorithms compared to general graphs.

Algorithm Overview:

- The algorithm exploits the DAG property that allows for topological sorting of its vertices.
- It starts by performing a topological sort of the graph to order the vertices linearly.
- The vertices are then processed in topological order, and for each vertex, the distances to its adjacent vertices are updated (relaxed) if a shorter path is found.

Steps:

The algorithm can be summarized in the following steps:

- 1. Perform a topological sort of the DAG.
- 2. Initialize distances from the source to all vertices as infinite, and the distance to the source itself as 0.
- 3. Process each vertex in the topologically sorted order. For each vertex, update the distances to its adjacent vertices
- 4. After all vertices are processed, the distance array contains the shortest distances from the source vertex to all other vertices.

Applications:

- This approach is used in scheduling problems where tasks depend on prior tasks (dependencies can be represented as a DAG).
- It is also useful in calculating shortest paths in networks that inherently do not contain cycles, such as dependency graphs in software projects.
- Finding shortest paths in DAGs can aid in optimization problems in various fields, including bioinformatics and compiler optimization.

Python Example: Single-Source Shortest Paths in DAGs

Below is a Python example demonstrating how to find single-source shortest paths in DAGs.

```
def shortest_path_dag(graph, source)
     top_order = topological_sort(graph)
distance = {vertex: float('infinity') for vertex in graph}
     distance[source] = 0
     for vertex in top_order:
          for neighbour, weight in graph[vertex]:
               if distance[vertex] + weight < distance[neighbour]:
    distance[neighbour] = distance[vertex] + weight</pre>
     return distance
def topological_sort(graph):
     visited = set()
stack = []
     def dfs(vertex):
          visited.add(vertex)
          for neighbour, _ in graph[vertex]:
    if neighbour not in visited:
                     dfs(neighbour)
          stack.insert(0, vertex)
     for vertex in graph:
           if vertex not in visited:
               dfs(vertex)
```

```
27
28 return stack
29
```

This function demonstrates calculating the shortest paths from a single source in a DAG by utilizing a topological sort followed by linear relaxation of edges.

The last section that is being covered in this chapter this week is **Section 24.3: Dijkstra's Algorithm**.

Section 24.3: Dijkstra's Algorithm

Dijkstra's Algorithm

Dijkstra's Algorithm is a popular method for finding the shortest paths from a single source vertex to all other vertices in a graph with non-negative edge weights. It is known for its efficiency in computing shortest paths on graphs without negative weight edges.

Algorithm Overview:

- The algorithm maintains a set of vertices whose shortest distance from the source is already determined and a set of vertices whose shortest distance is not yet determined.
- Initially, the distance to the source vertex is set to 0 and to infinity for all other vertices.
- At each step, the vertex with the minimum distance from the set of undetermined vertices is selected, its distance is finalized, and the distances to its adjacent vertices are updated.
- The process repeats until the shortest distances to all vertices are determined.

Steps:

The steps for Dijkstra's algorithm are as follows:

- 1. Initialize distances: Set the distance to the source vertex to 0 and all other vertices to infinity.
- 2. While there are vertices with undetermined shortest distances:
 - (a) Select the vertex with the minimum distance (let's call it the current vertex).
 - (b) Update the distances to the adjacent vertices of the current vertex.
 - (c) Finalize the distance of the current vertex.
- 3. The algorithm terminates when the shortest distances to all vertices are determined.

Applications:

- Dijkstra's Algorithm is widely used in network routing protocols to find the shortest path between nodes in a network.
- It is applied in geographical mapping to find the shortest path between geographical locations.
- This algorithm also finds its use in various fields such as telecommunications, transport, and game development for pathfinding and shortest path calculations.

Python Example: Dijkstra's Algorithm

Below is a Python implementation of Dijkstra's Algorithm.

```
import heapq
def dijkstra(graph, start):
distances = {vertex: float('infinity') for vertex in graph}
distances[start] = 0
```

```
priority_queue = [(0, start)]

while priority_queue:
current_distance, current_vertex = heapq.heappop(priority_queue)

if current_distance > distances[current_vertex]:
continue

for neighbour, weight in graph[current_vertex].items():
distance = current_distance + weight

if distance < distances[neighbour]:
distances[neighbour] = distance
heapq.heappush(priority_queue, (distance, neighbour))

return distances

return distances
```

This function calculates the shortest paths from a single source to all other vertices in a graph using Dijkstra's Algorithm, employing a priority queue to efficiently select the next vertex to process.

The next section that is being covered from this chapter this week is **Section 24.4: Difference Constraints**And Shortest Paths.

Section 24.4: Difference Constraints And Shortest Paths

Difference Constraints And Shortest Paths

Difference constraints are a form of constraints used in optimization problems that can be expressed as inequalities involving differences between variables. These constraints can be efficiently solved using shortest path algorithms in a graph, particularly by transforming the constraints into a graph and applying a shortest path algorithm such as Bellman-Ford, due to its ability to handle negative weights.

Algorithm Overview:

- The system of difference constraints is represented as a directed graph where each variable is a vertex.
- An inequality of the form $x_j x_i \leq b_k$ translates to an edge from vertex i to vertex j with weight b_k .
- A source vertex is added, connected to all other vertices with edges of weight 0, to facilitate finding shortest paths from this source to all vertices.
- The Bellman-Ford algorithm is then applied to find the shortest paths, which effectively solves the system of difference constraints.

Steps:

To solve a system of difference constraints, follow these steps:

- 1. Construct a directed graph based on the given constraints, adding a source vertex connected to all other vertices with zero-weight edges.
- 2. Apply the Bellman-Ford algorithm using the source vertex as the start. If the algorithm detects a negative cycle, the constraints are inconsistent.
- 3. If there are no negative cycles, the shortest path distances from the source to each vertex represent the solution to the system of constraints.

Applications:

- Difference constraints are widely used in scheduling problems where tasks have constraints relative to each
 other.
- They are also applied in circuit layout design, resource allocation, and various types of optimization problems where relationships between variables can be represented as inequalities.

Python Example: Solving Difference Constraints

While a direct Python example for solving a generic system of difference constraints is too specific to the system in question, the approach involves:

- 1. Modeling the constraints as a graph.
- 2. Applying the Bellman-Ford algorithm to find shortest paths.

This process finds feasible solutions to the difference constraints if they exist, based on the transformation of constraints into a shortest path problem.

The next chapter that is being covered this week is **Chapter 25: All-Pairs Shortest Paths**. The first section that is being covered from this chapter this week is **Section 25.1: Shortest Paths And Matrix Multiplication**.

Section 25.1: Shortest Paths And Matrix Multiplication

Shortest Paths And Matrix Multiplication

The concept of finding all-pairs shortest paths through matrix multiplication leverages the algebraic structure of matrices to efficiently compute the shortest paths between all pairs of vertices in a graph. This approach is particularly powerful for dense graphs and is based on the idea that the weight of the shortest path between two vertices can be computed through a series of matrix multiplications.

Algorithm Overview:

- The graph is represented as a weight matrix W, where W_{ij} is the weight of the edge from vertex i to vertex j. If there is no edge between i and j, W_{ij} is set to ∞ , except $W_{ii} = 0$ for all i.
- The algorithm iteratively computes matrix $D^{(k)}$, where $D^{(k)}_{ij}$ represents the shortest path from vertex i to vertex j using only vertices $\{1, 2, ..., k\}$ as intermediate vertices in the path.
- Initially, $D^{(0)} = W$. Then, for each k from 1 to n, $D^{(k)}$ is computed based on $D^{(k-1)}$ by considering whether a path through vertex k is shorter than the previously known paths.
- The final matrix $D^{(n)}$ contains the lengths of the shortest paths between all pairs of vertices.

Steps:

To compute all-pairs shortest paths using matrix multiplication, follow these steps:

- 1. Initialize $D^{(0)} = W$.
- 2. For k = 1 to n (where n is the number of vertices):
 - (a) Compute $D^{(k)}$ from $D^{(k-1)}$ where $D^{(k)}_{ij} = \min(D^{(k-1)}_{ij}, D^{(k-1)}_{ik} + D^{(k-1)}_{kj})$.
- 3. After completing the iterations, $D^{(n)}$ contains the shortest path distances between all pairs of vertices.

Applications:

- This method is valuable in network analysis, graph theory research, and operations research, especially in dense graphs where direct computation of paths is computationally intensive.
- It is also applied in computational geometry, for example, in finding the shortest path within geometric figures represented as graphs.

The last section that is being covered from this chapter this week is **Section 25.2: The Floyd-Warshall Algorithm**.

Section 25.2: The Floyd-Warshall Algorithm

The Floyd-Warshall Algorithm

The Floyd-Warshall algorithm is a dynamic programming technique for finding the shortest paths between all pairs of vertices in a weighted graph. This algorithm can handle graphs with positive or negative edge weights but cannot handle graphs with negative cycles. It is particularly well-suited for dense graphs or graphs where we need to find shortest paths between all pairs of vertices.

Algorithm Overview:

- The algorithm iteratively improves the estimate of the shortest path between all pairs of vertices through a series of intermediate vertices.
- It initializes the shortest paths as direct paths between all pairs or infinity if no direct path exists.
- For each vertex k, the algorithm updates the shortest path d[i][j] between each pair of vertices (i, j) considering k as an intermediate vertex. This update checks if d[i][k] + d[k][j] < d[i][j] and updates d[i][j] accordingly.
- After considering all vertices as intermediate points, the matrix d contains the lengths of the shortest paths between all pairs of vertices.

Steps:

The steps for the Floyd-Warshall algorithm are as follows:

- 1. Initialize the shortest path estimates using the adjacency matrix of the graph, where the weight of the edge (i, j) is used as the initial distance from i to j, and it's set to ∞ if there is no direct edge between i and j.
- 2. For each vertex k, attempt to update the shortest path d[i][j] for all pairs of vertices (i, j) by checking if a path through k is shorter.
- 3. Repeat step 2 for every vertex k in the graph.
- 4. After completing the iterations, the matrix contains the shortest distances between all pairs.

Applications:

- The Floyd-Warshall algorithm is widely used in network routing to find the shortest path in a network.
- It's also utilized in algorithms that require knowing the shortest distances between all pairs of vertices upfront, such as in some clustering and network analysis algorithms.
- This approach can serve in geographic information systems (GIS) to find the shortest paths across a network of roads or pathways.

Python Example: The Floyd-Warshall Algorithm

Below is a Python implementation of the Floyd-Warshall algorithm.

This function calculates the shortest paths between all pairs of vertices in a weighted graph using the Floyd-Warshall algorithm, represented as a matrix of edge weights.

Linear/Integer Programming And NP-Completeness

Linear/Integer Programming And NP-Completeness

13.0.1 Assigned Reading

The reading assignment for this week is from, Introduction to Algorithms:

- Chapter 29.1 Standard And Slack Forms
- Chapter 29.2 Formulating Problems As Linear Programs
- Chapter 29.3 The Simplex Algorithm
- Chapter 34.1 Polynomial Time
- Chapter 34.2 Polynomial-Time Verification
- Chapter 34.3 NP-Completeness And Reducibility
- Chapter 34.4 NP-Completeness Proofs
- Chapter 34.5 NP-Completeness Problems

13.0.2 Lectures

The lecture videos for this week are:

- Linear Programming: Introduction ≈ 13 min.
- Linear Programming: Visualizing Solutions ≈ 12 min.
- Linear Programming: Overview Of Algorithms ≈ 13 min.
- Linear Vs. Integer Programming ≈ 12 min.
- Decision Problems, Languages And Decidability ≈ 36 min.
- Reductions And NP Completeness ≈ 37 min.
- Nondeterministic Polynomial Time (NP) ≈ 29 min.
- Some NP Complete Problems ≈ 25 min.
- Polynomial Time Algorithms ≈ 29 min.

13.0.3 Assignments

The assignment for this week is:

• Problem Set 10 - Linear/Integer Programming And NP-Completeness

13.0.4 Quiz

The quizzes for this week are:

• Quiz 11

13.0.5 Chapter Summary

The first chapter that is being covered this week is Chapter 29: Linear Programming.

Chapter 29: Linear Programming

Linear Programming (LP)

Linear Programming (LP) is a mathematical method for determining a way to achieve the best outcome (such as maximum profit or lowest cost) in a given mathematical model and is represented by linear relationships. It is one of the simplest ways to perform optimization. LP is used in various fields such as business, economics, and engineering to maximize or minimize resource allocation.

Definition and Basics:

Linear programming involves:

- A linear function to be maximized or minimized, commonly known as the objective function.
- System constraints represented by linear inequalities or equations.
- Non-negativity restrictions where the decision variables must be greater than or equal to zero.

Mathematical Formulation:

The general form of a linear programming problem is given by:

- Objective Function: Maximize (or Minimize) $z = c_1x_1 + c_2x_2 + \ldots + c_nx_n$
- Subject to the constraints:

$$a_{11}x_1 + a_{12}x_2 + \dots + a_{1n}x_n \le b_1$$

$$a_{21}x_1 + a_{22}x_2 + \dots + a_{2n}x_n \le b_2$$

$$\vdots$$

$$a_{m1}x_1 + a_{m2}x_2 + \dots + a_{mn}x_n \le b_m$$

And non-negativity constraints:

$$x_1, x_2, \dots, x_n \ge 0$$

Common Algorithms:

Several algorithms can solve LP problems, including:

- 1. The Simplex Method: This is the most widely used method for solving linear programming problems efficiently. It iterates through feasible solutions at the vertices of the feasible region until the best value is found.
- 2. Interior-Point Methods: These are more modern algorithms that approach the solution from within the feasible region and can be faster than the Simplex method for large problems.

Applications:

- LP is used in manufacturing to determine the maximum production levels for multiple products with different profit contributions and resource requirements.
- In transportation, it helps in finding the most efficient dispatching of trucks, planes, or ships that minimizes
 costs or maximizes efficiency.
- It is also used in finance and budgeting to optimize investment portfolios or advertising media mixes.

The first section that is covered from this chapter this week is Section 29.1: Standard And Slack Forms.

Section 29.1: Standard And Slack Forms

Standard and Slack Forms

Standard and Slack Forms are two ways of representing linear programming problems that facilitate the application of algorithms like the Simplex Method for finding optimal solutions. These forms convert all constraints into equalities, allowing the use of linear algebra techniques in the solution process.

Standard Form:

The standard form of a linear programming problem is where all the constraints are expressed as equalities, and all variables are non-negative. Here is how a linear programming problem is represented in standard form:

- Objective Function: Maximize (or Minimize) $z = c_1x_1 + c_2x_2 + \ldots + c_nx_n$
- Subject to the constraints:

$$a_{11}x_1 + a_{12}x_2 + \dots + a_{1n}x_n = b_1$$

$$a_{21}x_1 + a_{22}x_2 + \dots + a_{2n}x_n = b_2$$

$$\vdots$$

$$a_{m1}x_1 + a_{m2}x_2 + \dots + a_{mn}x_n = b_m$$

• And non-negativity constraints:

$$x_1, x_2, \dots, x_n \ge 0$$

Slack Form:

Slack form is a variation of the standard form used primarily within the Simplex Method. In slack form, slack variables are added to convert inequalities into equalities. This allows the algorithm to work within a consistent framework of variable dimensions.

• From the general inequality constraints $Ax \leq b$, we add a slack variable s_i to each inequality to convert it into an equality:

$$a_{11}x_1 + a_{12}x_2 + \ldots + a_{1n}x_n + s_1 = b_1, \quad s_1 \ge 0$$

$$a_{21}x_1 + a_{22}x_2 + \ldots + a_{2n}x_n + s_2 = b_2, \quad s_2 \ge 0$$

$$\vdots$$

$$a_{m1}x_1 + a_{m2}x_2 + \ldots + a_{mn}x_n + s_m = b_m, \quad s_m \ge 0$$

• These slack variables s_1, s_2, \ldots, s_m are added to ensure that the constraints form a feasible solution space.

Conversion from Standard to Slack Form:

Conversion involves the addition of slack variables to inequalities to turn them into equalities. For instance, if a constraint is $x_1 + 2x_2 \le 10$, it is rewritten as $x_1 + 2x_2 + s = 10$ with $s \ge 0$ being the slack variable.

Applications:

- Both forms are crucial in computational solutions of linear programs. Standard form provides a unified way to write linear programs, which simplifies theoretical analysis.
- Slack form is particularly useful in the implementation of the Simplex algorithm, which requires all program constraints to be in equality form with non-negative variables.

The next section that is being covered from this chapter this week is **Section 29.2: Formulating Problems As Linear Programs**.

Section 29.2: Formulating Problems As Linear Programs

Formulating Problems As Linear Programs

Formulating problems as Linear Programs (LPs) involves translating a real-world scenario or decision-making process into a mathematical model that meets the structure required for linear programming. This process includes identifying variables, constraints, and an objective function that are all linear.

Steps to Formulate a Linear Program:

The process of formulating a linear programming problem typically involves the following steps:

- 1. **Define the variables:** Clearly define the decision variables that represent the quantities to be determined.
- 2. **Formulate the objective function:** Establish a linear objective function to maximize or minimize. This function usually represents cost, profit, or some other metric to optimize.
- 3. **Establish the constraints:** Set up constraints based on the limitations or requirements of the problem. These constraints must also be linear equations or inequalities.
- 4. Ensure non-negativity: Include non-negativity constraints for all variables, meaning that all decision variables should be greater than or equal to zero.

Company Production Problem

Consider a company that manufactures two types of products using two different resources. The goal is to maximize profit given the resource constraints.

• Variables:

$$x_1 =$$
Number of units of Product 1
 $x_2 =$ Number of units of Product 2

• Objective Function:

Maximize
$$z = 20x_1 + 30x_2$$

This represents maximizing the total profit, where Product 1 contributes \$20 per unit, and Product 2 contributes \$30 per unit.

• Constraints:

$$3x_1 + 2x_2 \le 18$$
 (Resource 1)
 $2x_1 + 4x_2 \le 16$ (Resource 2)
 $x_1, x_2 \ge 0$

These constraints ensure that the use of Resource 1 and Resource 2 does not exceed the available amounts.

Key Considerations:

- It is crucial to ensure that all relationships (objective and constraints) are linear. This is what defines an LP and allows the use of linear programming techniques.
- Variables must accurately represent all dimensions of the problem to avoid oversimplification or inaccurate
 modeling.
- The choice between maximizing or minimizing the objective function should reflect the ultimate goal of the problem-solving scenario.

Applications:

Linear programming can be applied to a wide range of problems, including:

- Resource allocation tasks, such as budgeting and logistics.
- Scheduling problems, including workforce and manufacturing schedules.
- Blending problems, where components must be combined in optimal proportions (e.g., in food production or chemical manufacturing).

The last section that is being covered from this chapter this week is Section 29.3: The Simplex Algorithm.

Section 29.3: The Simplex Algorithm

The Simplex Algorithm

The Simplex Algorithm is a popular method for solving linear programming problems that uses an iterative process to move from one vertex of the feasible region to an adjacent vertex with a non-decreasing value of the objective function, eventually reaching the optimal solution.

Algorithm Overview:

The Simplex Algorithm, developed by George Dantzig in 1947, is a cornerstone in the field of operational research for solving optimization problems in linear programming.

- The algorithm involves iterations with two main operations: pivot operations that move to a better adjacent feasible solution, and checking for optimality.
- It requires the linear programming problem to be in standard form and typically uses the slack form during the computation.

Steps of the Simplex Algorithm:

The key steps in the Simplex Algorithm are as follows:

- 1. **Initialization:** Convert the LP problem into slack form and identify a basic feasible solution (BFS).
- 2. Checking for Optimality: Determine if the current BFS is optimal by checking if there are any negative coefficients in the objective function row. If not, the current solution is optimal; otherwise, proceed to the next step.
- 3. **Pivot Operation:** Select a non-basic variable to enter the basis and a basic variable to leave the basis, based on the objective to increase (or decrease) the value of the objective function. This step is often done using the pivot rule, such as the smallest coefficient rule or the largest increase rule.
- 4. **Update Basis:** Perform row operations to update the tableau to reflect the new basis. This involves Gaussian elimination to make the entering variable a basic variable.
- 5. **Iteration:** Repeat the checking and pivot operations until the optimal solution is found.

Computational Considerations:

- The efficiency of the Simplex Algorithm can significantly vary based on the choice of pivoting rule; however, in practice, it performs very efficiently on most problems.
- While the worst-case computational time can be exponential, this is rare, and the Simplex Algorithm is polynomial on average for random problems.

Applications:

- The Simplex Algorithm is extensively used in industries for optimizing resources, maximizing profits, minimizing costs, and improving operational efficiency.
- It is applicable in various sectors including manufacturing, transportation, finance, and any area where resource allocation is involved under constraints.

Python Example: Simplex Algorithm

Below is a conceptual outline of implementing the Simplex Algorithm in Python.

```
def simplex_algorithm(A, b, c):

# A is the matrix of coefficients, b is the right-hand side vector,

# c is the coefficients of the objective function

tableau = initialize_tableau(A, b, c)

while not is_optimal(tableau):

pivot_column = select_pivot_column(tableau)

pivot_row = select_pivot_row(tableau, pivot_column)

pivot(tableau, pivot_row, pivot_column)

update_tableau(tableau, pivot_row, pivot_column)

return get_solution(tableau)
```

This code skeleton outlines the main steps of the Simplex Algorithm, focusing on setup, iteration, and solution extraction phases, and it needs detailed functions to handle each specific step.

The next chapter that is being covered this week is **Chapter 34: NP-Completeness**.

Chapter 34: NP-Completeness

NP-Completeness

NP-Completeness is a fundamental concept in theoretical computer science that helps classify computational problems according to their inherent difficulty. This concept is crucial in understanding the limits of what problems can be efficiently solved using algorithms.

Introduction to Complexity Classes:

NP-Completeness deals with the complexity classes NP, P, and NP-complete. Here are the basic definitions:

- P (Polynomial Time): This class includes those problems that can be solved in polynomial time, i.e., the amount of time required to find a solution can be expressed as a polynomial function of the size of the input data.
- NP (Nondeterministic Polynomial Time): This class consists of those problems for which a given solution can be verified in polynomial time. Note that while all problems in P are also in NP, whether all NP problems are in P is an unresolved question known as the P vs NP problem.
- NP-Complete: This is a subset of NP that are amongst the hardest problems in NP. A problem is NP-Complete if every problem in NP can be reduced to it in polynomial time.

Understanding Reductions:

Reductions are a way to show that a problem X is at least as hard as a problem Y. If Y can be "reduced" to X (written as $Y \leq_p X$), and X can be solved in polynomial time, then Y can also be solved in polynomial time.

- A common type of reduction used in proving NP-Completeness is the polynomial-time reduction.
- To prove a problem is NP-Complete, one must show it is in NP, and that every problem in NP can be reduced to this problem in polynomial time.

Significant Theorems and Concepts:

- Cook-Levin Theorem: This theorem states that the Boolean satisfiability problem (SAT) is NP-Complete, making it the first known NP-Complete problem.
- NP-Hard: Problems that are at least as hard as the hardest problems in NP but are not necessarily in NP themselves. These problems do not need to have a polynomial-time solution verification.

Practical Implications and Applications:

Understanding NP-Completeness has practical implications in various fields:

- It helps in determining the boundaries of feasible computational solutions, guiding researchers and practitioners in whether to look for exact, approximate, or heuristic solutions.
- In areas like cryptography, scheduling, and network design, knowing whether a problem is NP-Complete can influence the approach to problem-solving (e.g., using approximate algorithms instead of exact ones).

Example Problems:

Some classical NP-Complete problems include:

- Traveling Salesman Problem (TSP)
- Graph Coloring
- Knapsack Problem
- Hamiltonian Cycle Problem

These problems are widely studied not only for their theoretical importance but also for their real-world applications in optimizing tasks, resource allocation, and planning under constraints.

The first section that is being covered from this chapter this week is **Section 34.1: Polynomial Time**.

Section 34.1: Polynomial Time

Polynomial Time

Polynomial time is a term used in computational complexity theory to describe the class of problems known as "P" or "Polynomial Time Problems." These are problems that can be solved by an algorithm whose running time grows polynomially with the size of the input.

Definition and Characteristics:

- A problem is said to be in polynomial time if there exists an algorithm that solves the problem and the execution time of the algorithm is a polynomial function of the size of the input for the algorithm.
- Formally, a problem is in class P if it can be solved in time $\mathcal{O}(n^k)$ for some non-negative integer k, where n is the size of the input.

Examples of Polynomial Time Algorithms:

Examples of problems that can be solved in polynomial time include:

- Determining if a number is prime (can be done in $\mathcal{O}(n^6)$ time using the AKS primality test).
- Finding the greatest common divisor (GCD) of two numbers using the Euclidean algorithm.
- Sorting a list of numbers using Merge Sort or Quick Sort.

Importance of Polynomial Time:

- Polynomial time algorithms are considered efficient and feasible for practical use because their running times grow at a manageable rate as the size of the input increases.
- The class P is central in the field of complexity theory, particularly in the study of NP-completeness. The famous "P vs NP" problem asks whether every problem whose solution can be verified in polynomial time can also be solved in polynomial time.

Conceptual Understanding:

- The notion of polynomial time relates to worst-case running time—it considers the maximum time needed to solve the problem across all possible inputs of a particular size.
- This contrasts with "exponential time," where the running time grows exponentially with the input size, which is typically not feasible for large inputs.

Theoretical Implications:

- Polynomial time serves as a benchmark for algorithm efficiency. Algorithms not bounded by polynomial time are usually impractical for large inputs due to their high computational demands.
- Understanding which problems can be solved in polynomial time helps in classifying computational problems and guides researchers in developing more efficient algorithms.

The next section that is being covered from this chapter this week is **Section 34.2: Polynomial-Time Verification**.

Section 34.2: Polynomial-Time Verification

Polynomial-Time Verification

Polynomial-Time Verification is a crucial concept in computational complexity theory that pertains to the class of problems known as NP, or "Nondeterministic Polynomial Time." This class includes problems for which a given solution can be verified quickly (in polynomial time), even if finding the solution might be slow.

Definition and Fundamentals:

- A problem is in NP (Nondeterministic Polynomial Time) if given a "certificate" or a proposed solution, the correctness of the certificate can be verified in polynomial time with respect to the size of the input.
- This does not necessarily mean that a solution can be found in polynomial time, only that if a potential solution is provided, its validity can be confirmed or denied quickly.

Examples of Polynomial-Time Verification

Common scenarios where solutions can be verified in polynomial time include:

- Boolean Satisfiability Problem (SAT): Given a Boolean formula and a truth assignment for its variables, verifying whether this assignment satisfies the formula can be done in linear time, which is polynomial.
- **Graph Coloring:** Checking if a given coloring of a graph (where colors are assigned to vertices) is valid (no adjacent vertices share the same color) can also be completed in polynomial time.
- Hamiltonian Path Problem: Given a path in a graph, verifying whether it is a Hamiltonian path (visits every vertex exactly once) can be achieved in polynomial time.

Importance of Polynomial-Time Verification:

- This property is what defines the complexity class NP. It is significant because it includes many important problems for which no polynomial-time solving algorithm is known, but whose solutions can be verified quickly.
- It helps in understanding computational feasibility when direct solving is impractical or unknown.

Role in NP-Completeness:

- To prove that a problem is NP-Complete, it must first be shown that it belongs to NP. This is done by demonstrating that any proposed solution can be verified in polynomial time.
- After establishing that a problem is in NP, it must be shown that it is as hard as any problem in NP, usually by a polynomial-time reduction from another NP-Complete problem.

Theoretical and Practical Implications:

- Polynomial-time verification allows researchers to use heuristic and approximation algorithms to find potential solutions to complex problems and then verify their correctness efficiently.
- In practical applications, especially in fields like cryptography and network security, being able to verify a solution quickly is often more critical than being able to find one efficiently.

The last section that is being covered from this chapter this week is **Section 34.3: NP-Completeness And Reducibility**.

Section 34.3: NP-Completeness And Reducibility

NP-Completeness and Reducibility

NP-Completeness and Reducibility are central concepts in computational complexity theory that provide a framework for classifying and understanding the difficulty of various computational problems. These concepts are crucial for proving whether problems in NP can be considered as hard as the hardest problems in NP, known as NP-complete problems.

Defining NP-Completeness:

- A problem is defined as NP-complete if it is both in NP and as hard as any problem in NP. This means that any problem in NP can be reduced to this problem in polynomial time.
- The first problem proven to be NP-complete was the Boolean satisfiability problem (SAT), via the Cook-Levin theorem.

Understanding Reducibility:

Reducibility is a method used to compare the difficulty of computational problems. A problem P is reducible to a problem Q (denoted as $P \leq_p Q$) if an instance of P can be transformed into an instance of Q in polynomial time.

- **Polynomial-Time Reduction:** If problem A can be reduced to problem B in polynomial time and B can be solved in polynomial time, then A can also be solved in polynomial time.
- This type of reduction is used to prove NP-completeness by showing that a known NP-complete problem can be reduced to the problem in question in polynomial time.

Steps to Prove NP-Completeness:

To prove that a problem is NP-complete, follow these steps:

- 1. Show that the problem is in NP. This involves demonstrating that a given solution can be verified in polynomial time.
- 2. Choose a known NP-complete problem and reduce it to the problem in question, ensuring that the reduction process itself runs in polynomial time.
- 3. Demonstrate that this reduction is correct and preserves the problem structure, implying that solving the reduced problem would solve the original NP-complete problem.

Significance of Reducibility:

- Reducibility is important because it helps classify problems based on their computational complexity. By using reductions, complexity theorists can create a hierarchy of problems based on their difficulty.
- It also assists in identifying whether new problems are NP-complete, helping computer scientists understand whether these problems are likely to have polynomial-time solutions or not.

Applications and Implications:

- Understanding NP-completeness and reducibility is vital for fields such as cryptography, algorithm design, and artificial intelligence, where decision-making processes depend heavily on problem complexity.
- In practice, knowing that a problem is NP-complete may lead to the focus on heuristic or approximation algorithms rather than exact solutions, especially in real-world applications where solutions need to be found within reasonable time constraints.



Exam 4

Exam 4

14.0.1 Exam

The exam for this week is:

- Spot Exam 4 Notes
- Spot Exam 4



Quantum Algorithms

Quantum Algorithms

15.0.1 Lectures

The lecture videos for this week are:

- Computation And Physics ≈ 30 min.
- Qubits And Quantum Operators ≈ 37 min.
- Entanglements And Bell's Inequality ≈ 22 min.
- Grover's Search Algorithm ≈ 42 min.



Final Exam

Final Exam

16.0.1 Exam

The exam for this week is:

- Final Exam Notes
- Final Exam

