1. Asymptotic Complexity and Basic Data Structures:

1.1. Upper bound, lower bound and tight bound definitions. Given an analytical function tha determine its asymptotic complexity. Compare two functions and determine which one wil dominate for larger and small instance sizes.

In algorithm analysis, we use asymptotic notation to describe the performance of an algorithm. Asymptotic notation refers to how an algorithm behaves as the size of the input grows without limit. The three most common types of asymptotic notation are:

- Big O notation: This represents the upper bound or worst-case scenario for the running time of an algorithm. It gives an
  upper limit on the number of operations required by the algorithm to solve the problem for any input of size n.
- <u>Bio Omega notation</u>: This represents the lower bound or best-case scenario for the running time of an algorithm. It gives
  a lower limit on the number of operations required by the algorithm to solve the problem for any input of size n.
- Big Theta notation: This represents the tight bound or average-case scenario for the running time of an algorithm. It gives an estimate of the number of operations required by the algorithm to solve the problem for any input of size n.

en an analytical function, we can determine its asymptotic complexity by looking at the growth rate of the function as the e increases. We typically ignore lower-order terms and constants and focus on the dominant term or terms that grow the

o compare two functions and determine which one will dominate for larger and small instance sizes, we can look at their rates. If one function has a higher growth rate than the other, it will dominate for larger input sizes. For smaller input sizes simining function may depend on the specific values of the input. We can also use the limit definition of asymptotic notation we two functions.

For example, consider the following two functions:

 $f(n) = 2n^2 + 5n + 1$ ;  $g(n) = n^3 + 10n$ 

As n approaches infinity, the dominant term in f(n) is  $2n^2$ , while the dominant term in g(n) is  $n^3$ . Therefore, g(n) will inste f(n) for large input sizes. For small input sizes, however, f(n) may dominate g(n) depending on the specific values of n.

1.2. Size of an instance given its implementation data structure for a selected operation. For example, using adjacency matrix, determine the asymptotic complexity of a DFS traversal.

In computer science, the size of an instance refers to the amount of input data required to perform a given operat example, the size of an instance for a sorting algorithm may be the number of elements to be sorted, or for a graph tra-algorithm like DFS, it may be the number of vertices in the graph.

The adjacency matrix is a way of representing a graph as a matrix where the rows and columns or the entries correspond to the edges. Specifically, if the graph has n vertices, then the adjacency ma-re the entry in row i and column j is 1 if there is an edge from vertex i to vertex j, and 0 otherwise.

To analyze the time complexity of a DFS traversal using an adjacency matrix, we need to count the number of basic ations performed by the algorithm as a function of the size of the instance. The basic operations in a DFS traversal are visitin tex, marking it as visited, and recursively visiting its unwisted neighbor.

If we use a standard depth-first search algorithm, the time complexity of the algorithm can be expressed in terms of the unber of vertices and edges in the graph. In particular, the time complexity of DFS on a graph represented by an adjacency trivia is QV2V2, where V is the number of vertices in the graph. This is because each vertex needs to be visited at most once, each vertex, we need to examine all V adjacent vertices in the matrix.

It's worth noting that if the graph is sparse, i.e., it has relatively few edges compared to the number of vertices, then it may be more efficient to use an adjacency list representation instead of an adjacency matrix. In that case, the time complexity of DFS would be OV +E, where E is the number of edges in the graph.

### 2. Graph Algorithms:

2.1.BFS and DFS labelling of nodes and asymptotic complexity. Given a graph understand the order in which the nodes are visited and the edges are explored.

BFS and DFS are two algorithms used for traversing graphs. Both algorithms visit each node in the graph, but they visit them different orders. BFS stands for Breadth-First Search, while DFS stands for Depth-First Search.

In BFS, we start at a node, visit all its neighbors, then visit all the neighbors of those neighbors, and so on. We keep track of nodes we have visited so that we don't visit them again. We use a queue data structure to keep track of the order in which we trodes. The order in which we visit nodes using BFS is such that nodes at the same level in the graph are visited before nodes to exect level.

ors, then visit one of that neighbor's neighbors, and so on. We keep track of again. We use a stack data structure to keep track of the order in which we's is such that we go as deep as possible before backtracking.

Here's the pseudo code for DFS traversal

The symptotic complexity of both algorithms depends on the size of the graph and the way it is represented. For examt the graph is represented using an adjacency list, BFS and DFS both have a time complexity of O(V+E), where V is the number vertices and E is the number depicts in the graph. If the graph is represented using an adjacency matrix, BFS and DFS both attent complexity of O(V-E).

In terms of the order in which nodes are visited and edges are explored, BFS visits nodes in order of their distance from the starting node, while DFS explores the edges of the graph in a depth-first manner, meaning it visits all the edges of one branch before movine on to another branch.

2.2. Topological sorting and its complexity. Given a directed graph, without cycles, determine the order of a possible topological sorting and its asymptotic complexity.

The function takes a directed acyclic graph represented as an adjacency list graph as input and returns a vector sorted senting a possible topological sorting of the vertices in the graph.

The algorithm works by first initializing the indegree of all vertices to 0 and then calculating the indegree of each vertex by iterating through the edges in the graph. The algorithm then creates a queue to store vertices with indegree 0, i.e., vertices that have no incoming edges.

The algorithm then performs a BFS-like traversal of the graph by repeatedly dequeuing a vertex u with indegree 0, adding it to the sorted list, and then reducing the indegree of its neighbors by 1. If any of the neighbors of u now have indegree 0, they are added to the equeue.

# 3. Brute-Force Approach:

3.1. You will need to reason about the use of the Brute-Force approach in the development of an algorithm derive a simple pseudo-code implementation and determine its asymptotic complexity for a selected choice of a data structure. (see final note on pseudo-code).

The Brute-Force approach is a simple and straightforward way of solving problems that involves trying all possible solutions selecting the best one. Here's an example of how it can be applied to find the maximum element in an array:

In this code, we define a function findMax that takes an array arr and its size n as input and returns the maximum element in the array. We start by initializing a variable maxilement with the first element of the array. We then loop through the rest of the array and compare each element with the maxilement variable. If we find an element that is greater than the maxilement, we update maxilement with the new element. Finally, we return the maxilement.

The time complexity of this Brute-Force approach is O(n), where n is the size of the array since we need to compare each ent in the array with the maxElement variable.

Note that this is just a simple example of how the Brute-Force approach can be applied to solve a problem. In general, this approach may not be the most efficient or practical way of solving complex problems, but it can be a useful starting point for destending more continued. The problems is a simple continued to the continued of the continued

# **Graph Representation**

```
· Adjacency List vs. Adjacency Matrix
```

```
undirected
                            1 | 2 4 /
2 | 1 3 4 /
3 | 2 /
4 | 1 2 /
      1
                3
      2
                 4
directed
                            1 2 4 / 2 3 /
      1
                3
```

1.3. The min-heap as a key data structure. Complexity of its key operations siftDown and siftUp (you do not need by heart the pseudo-code of these basic operations, but need to understand how they operate). Given a sequence of value insertion determine the configuration of a minheap or max-heap and reason about the complexity of the sequence of operations.

A min-houp is a binary tree in which the parent node is always less than or equal to its children. It is a key data structure used orting and priority queue implementations. The sift(I) and siftDown operations are used to maintain the heap property after a is inserted or defeating.

The siftUp operation is used after an element is inserted into the heap. It compares the element with its parent and swaps if the parent is larger. This operation is repeated until the parent is smaller or until the root is reached. The complexity of the poperation is (Olgo n), where is the number of nodes in the heap.

The siftDown operation is used after an element is deleted from the heap. It swaps the deleted node with its smallest child until the heap property is restored. This operation is repeated until the ismallest child is greater than the node or until the leaf is reached. The complexity of the siftDown operation is also O(log n).

To determine the configuration of a min-heap or max-heap after a sequence of value insertions, we start with an empty heap and add each element one at a time using the siftUp operation. The final configuration of the heap depends on the order in which the elements were inserted.

For example, if we insert the elements [3, 2, 4, 1, 5] into an empty min-heap, the resulting heap would be:



The algorithm continues until all vertices have been added to the sorted list or until the queue is empty. If a topological sorting is found, the function returns the sorted list. If the sorted list has fewer than n vertices, then the graph contains at least cycle, and the function returns a nepty vector to indicate failure.

The asymptotic complexity of the algorithm is O(V + E), where V is the number of vertices and E is the number of edges in the graph, as the algorithm needs to visit each vertex and each edge exactly once.

Here's the pseudo code in C++ for topological sorting with comments explaining what's hap

2.3. Understand and apply the Strongly-Connected components algorithm that replies on double pass over the graph (and its Reverse variant) and determine the overall asymptot complexity. Application of this algorithm to an example graph.

The Strongly-Connected components algorithm is used to identify the strongly-connected components of a directed graph strongly-connected component is a adopted where every vertex is reachable from every other vertex in the subgraph. Algorithm works by proferming a depth efficiency (OFS) on the graph, and then performing a case of DFS on the reput and the operations accord DFS on the result of the professional particular and all object reversed. The vertices are then sorted in order of their finishing times from accord DFS, and the order is used by group the vertices are through connected components.

# 4. Greedy Algorithms:

4.1. The two properties of an optimal greedy choice and optimal solution to a sub-problem. Understanding the two properties what how to prove them for a selected given greedy algorithm (Notez you will not be asked detailed proofs of these greedy properties, but rather you will be asked to reason about them for selected covered algorithms or a sample algorithm provided in the test).

- 1. Greedy choice property: A globally optimal solution can be obtained by making a locally optimal (greedy) choice
- Optimal substructure property: A problem can be solved optimally by breaking it down into subproblems and solving each subproblem optimally.

To prove these properties for a given greedy algorithm, we need to demonstrate that the algorithm satisfies these

Let's take the example of the Activity Selection problem to illustrate this. In this problem, we are given a set of activities, each with a start time and an end time, and we need to select the maximum number of non-overlapping activities that can be performed.

- 1. Sort the activities by their end times
- For each subsequent activity in the sorted list, if its start time is greater than or equal to the end time of the previously selected activity, select it and update the end time.

To prove the two properties of optimal greedy choice and optimal substructure for this algorithm, we can reason as follows:

- prove the two properties of optimal growly obsect and optimal anothershare for this algorithm, we can reason is allower.

  I. Greek challer property: A close sky to ke algorithm selects the activity that has the entiries and time among the treatments activities. This is a locally optimal choice, as it ensures that we can select the maximum number of non-overlapping activities that the state of the contractivity. By selecting the activity with the entirest end time, we there room for selecting other activities that start later. We can above that this locally optimal choice leads to a globally optimal solution that does not include the activity with the earliest of time. We can always replace this activity with the one with the earliest end time. We can always replace this activity with the one with the earliest end time and obtain a new solution that is deem as good as the previous one.
- 2. Optimal substructure property: We can break the problem of selecting the maximum number of non-over activities into subproblems by considering the extivities starting from the second one and selecting the mamber of non-overlapting selectivities than shorts. It is a subspection of the original problem, and solve is optimally using the same algorithm recovery. The optimal solution to the original problem can obtained by adding the fert activity to the solution of the subspection.



```
] = true;
i = 0; i < adj[v].size(); ++i) {
visited[adj[v][i]]) {
FS(adi add[v][i] < visited);
```

- DFS: performs a DFS on the graph to find the vertices in the order of their finishing times
- se: performs a DFS on the reverse of the graph to find the strongly-c

findSCC: performs the entire algorithm by first performing a DFS on the graph to find the finishing times, and then
performing a DFS on the reverse of the graph to find the strongly-connected components.

The overall asymptotic complexity of this algorithm is O(V+E), where V is the number of vertices and E is the number of edges in the graph. This is because we perform two DFS traversals over the graph and its reverse, each taking O(V+E) time. To apply this algorithm to an example graph, we can create an adjacency list representation of the graph and pass it to the dSCC function. This will return a vector of strongly-connected components, where each component is represented as a vector

ivity struct represents an activity with a start and finish time. The compareActivities functreasing order of finish time.

The article/solution function that as vector of activities and netwes the maximum number of activities that can be selected. If first costs the activities that can be selected. If first costs the activities by finish time using the cost function. It then institutes to the official first time to make elected costs to activity whose tender activities and selected costs to activity whose unit time is great the activity and selected costs. It is not activities and selected costs to activity whose unit time is great the activity of the activity's finish time and increments selected Cossat. Finally, it returns selected Cossat.

In the main function, we create a vector of sample activities and call the activitySelection function to determine the insum number of activities that can be selected.

4.2. Fractional Knapsack algorithm and why it produces an optimal solution, Given an example, determine the outcome of the algorithm in terms of the sequence of selected items. Illustrate why the same greedy algorithm discussed in class does not work optimally for the case of the integer Knapsack problem.

- 1. The items are sorted in non-increasing order of their value/weight ratio using the compare fun
- The maximum value that can be obtained is initialized to 0.
- For each item, the amount of the item that can be added to the knapsack is calculated as the m remaining capacity divided by the item's weight.
- 5. The algorithm returns the maximum value that can be obtained

The Fractional Kapsack algorithm produces an optimal solution because it always selects the item with the highest value/weight ratio, which maximizes the value per unit of weight. However, the same greedy algorithm does not work optimally for the case of the integer Kanpack problem because it may select an item with a lower value/weight ratio if it fits better in the kanpacak, which may not result in the optimal solution.

un-Cost Spanning Tree algorithms (Prim and Kruskal) and their implementation time as well as examples of application in restricted graphs. Properties of the algorithms of data structures that best suit their implementation. Proof of correctness of the no of the MST (choice of the lightest and safe edge) as well as regarding the two poritims properties. For restricted graphs, explore or develop possibly simpler and reason about their correctness and asymptotic time complexity. greedy algorithm pr algorithms and rea

Minimum-Cost Spanning Tree (MCST) algorithms aim to find the tree with the minimum total weight in each con hted graph. Two well-known algorithms for finding the MCST are Prim's algorithm and Kruskal's algorithm.

- 1. Prim's Algorithm:

  - From the set of vertices not yet included in the tree, choose the one that has the minimum distance to the
    tree and add it to the tree.
  - Repeat step 2 until all vertices are included in the tree.

In the above code, we define a Node struct to store the index of the node and its distance from the source node. We also Graph class with a contractor to initialize the number of vertices, an additing function to add edges to the graph, and a function to implement Dijkstris Algorithm to find the shortest path from a source node to all other vertices.

In the dijkstrn function, we initialize a priority queue to maintain the vertices sorted by their distance from the source vector to store the distances of all vertices from the source (initialized as INT\_MAX), and a boolean array to keep tra-certices visited so for (initialized as false). We also set the distance of the source vertex to itself as 0.

Then, we start a loop while the priority queue is not empty. In each iteration of the loop, we extract the vertex with the und distance value from the priority queue. We mark this vertex as visited and iterate over its adjacent vertices. For each at vertex, we update its distance value in the distances vector if it can be improved by going through the current vertex.

To do this, we calculate the tentative distance as the sum of the current vertex's distance and the weight of the edge connecting the current vertex to the adjacent vertex. If this tentative distance is less than the current distance of the adjacent vertex, we update in distance value and add to the priority queue.

Finally, when all vertices have been visited, the distances vector contains the shortest path distances from the source to all other vertices in the graph.

The time complexity of Dijkstra's algorithm is O((E+V)log(V)), where E is the number of edges and V is the number of ces in the graph. This is due to the use of the priority queue data structure in the algorithm.

nece in the gipet. Insile auto et use of the prototy queue and streams the suprime.

A.S. Max-Flow Ford-Fulkerson Algorithm — The Edmonds-Karp implementation using BFS for selection of augmentation paths. Examples and selected cases for DAG. Understanding extreme cases of time complexity for fractional and irrational edge weights. Implementation complexity, Use of Max-Flow in Maximal Bipartite Matching and reasoning about their correctness. Understanding the Min-Cut/Max-Flow Theorem and apply it to sample case to determine the impossibility of specific flow values. For the Max-Flow algorithm you will not be asked to proof the correctness of this greedy algorithm.

Here is the pseudo code implementation of the Edmonds-Karp algorithm for finding maximum flow in a net

In this algorithm, we first initialize a graph represented by an algorithm, the first initialize a graph represented by a flag case, of list, a capacity matrix to store the capacity of each edge, as flow matrix to store the flow through each edge, and a parent array to store the sugmenting path. We then define a function that these a source mode and a target not one input an element represent for there exists an anguenting path from the source to the target, and false etherwise. This functions uses a Breadth-First Search algorithm to find the shortest augmenting goals and store the target, and false etherwise. The function target and extra source node and a starget node as imput and extrems the maximum those from the source to the target. This function tark tasks as some node and a starget node as imput and extrems the maximum thow from the source to the target. This function repeatedly calls the 10st function to find suggenteding paths and updates the flow along the aggreeting gaths and not once augmenting paths and paths exist.

We can improve the time complexity of this algorithm by using the fact that the max function is associative and mutative. This means that we can compute the maximum of multiple elements in parallel, rather than recursively computing aximum of two elements at a time. Here's an improved implementation of the find, max function that takes advantage of this

In this improved implementation, we first check if there are only two elements, and return the maximum of those two elements. Otherwise, we divise the array into two balves, and recursively compute the maximum of the left half and the maximum of the right half-fraidly, we return the maximum of these two maximums.

The running into  $\theta$  is improved algorithm can be analyzed using the Master Theorem. In this case, we have  $\alpha = 2$  (since we are dividing the array into two halves), b = 2 (since each subproblem has half the size of the original problem), and d = 1 (since the time complexity of comparing the maximum of two elements in G(1)). Therefore, we have:

# $T(n)=2T(n/2)\pm O(1)$

According to the Master Theorem, this recurrence has a running time of O(nlog(n)), which is much faster than the previous

By analyzing the terms in the Master Theorem for a specific code example or recurrence, we can identify which parts of the orithm contribute most to the running time, and how we can modify be algorithm to improve its time complexity. Changing a fiftic section of the code can lead to shanges in the asymptotic complexity, and we need to carefully analyze the impact of these fiftic section of the code can lead to shanges in the asymptotic complexity, and we need to carefully analyze the impact of these

```
r (auto v : adj[u]) {
  int weight = v.second, node = v.first;
  if (ivis[node] && dist[node] > weight) {
      dist[node] = weight;
      ps.gush(make_pair(dist[node], node);
    }
}
```

- Sort all the edges in non-decr
- Pick the smallest edge. Check if it forms a cycle with the spanning tree formed so far. If the cycle is not formed, include this edge.
- Repeat step 2 until there are (V-1) edges in the spanning tree

ion of Kruskal's Algorithm using the Union-Find algo

```
t int MAXN = 1e5 + 5;
parent[MAXN], rank[MAX
```

The time complexity of the Edmonds-Karp algorithm is O(VE'2), where V is the number of vertices and E is the number of edges. This is because in the worst case, we may have to update the flow along every edge in the graph for every iteration of the algorithm. However, in practice, the algorithm usually performs much better than this worst-case bound.

The Edmonds-Karp algorithm can be used to solve many types of network flow problems, including the Maximal Bipartite ching problem. In this problem, we are given a bipartite graph and we want to find a matching that covers as many vertices as sible. A matching is a set of edges such that no two edges share a common vertex.

To use the Edmonds-Karp algorithm for this problem, we first convert the bipartite graph into a flow network by adding a source node and a sink node and connecting the source node to one partition of the bipartite graph and the other partition to the sink node. We thus not the capacity of each edge to 1, indicating that each vertex can be matched with at most one other vertex. The maximum flow in this network corresponds to the maximum matching in the bipartite graph.

The correctness of the Edmonds-Karp algorithm can be proved using the Max-Flow Min-Cut theorem. This theorem states that the maximum flow in a network is equal to the minimum cut, where a cut is a partition of the vertices into two disjoint sets such that the source is in one set and the sides is the other set. The minimum cut corresponds to the minimum number of edges that need to be removed to disconnect the source from the sink.

In the Edmonds-Karp algorithm, we always choose the shortest augmenting path from the source to the sink, which corresponds to a cut in the residual graph. Therefore, the flow along this path is equal to the capacity of the minimum cut. By repeating this process, we eventually find the maximum flow, which is equal to the capacity of the minimum cut.

The time complexity of the Edmonds-Karp algorithm can be improved to O((VE)log(U)) using the Dinic's algorithm, where U is the maximum capacity of any edge in the network. However, this improvement is not necessary for most practical

5.2. Examples of the recurrences of the Merge-Sort and Hanoi Towers algorithms and their

# 5.2.1.Merge-Sort Algorithm

The Merge-Sort algorithm is a popular sorting algorithm that uses the divide and conquer approach to sort an array. The algorithm consists of two main steps - dividing the array into two halves and recursively sorting them, and merging the sorted halves to produce the final sorted array. Here's the peach endocode for the Merge-Sort algorithm.

```
merge(int arr[], int lef
int n1 = mid - left + 1;
int n2 = right - mid;
```

In the merge\_sort function, we first check if the left index is less than the right index. If it is, we calculate the middle index of recursively call the merge\_sort function on the left and right halves of the array. Once both halves are sorted, we merge them ing the merge function.

# T(n) = 2T(n/2) + O(n)

The time complexity of Prim's algorithm for finding the minimum spanning tree of a graph is  $O(V^*2)$  with an adjacency matrix representation and  $O(E \log V)$  with an adjacency list representation, where V is the number of vertices and E is the number of edges in the graph.

On the other hand, Kruskal's algorithm has a time complexity of  $O(E \log E)$  or  $O(E \log V)$ , whichever is smaller, where E is the number of edges in the graph and V is the number of vertices.

In general, Kruskal's algorithm is faster than Prim's algorithm when the graph is sparse (i.e., has relatively few edges compared to the number of vertices), while Prim's algorithm is faster when the graph is dense (i.e., has relatively many edges compared to the number of vertices). However, the actual time complexity depends on the implementation and the specific characteristics of the input graph.

4.4. Dijkstra Shortest path algorithm and why it produces an optimal shortest path from a single 74. Typista shortes pain agoithm and will produce a dynamic another pain tool a single source. Examples of application in special cases for DAGs or graphs with restricted edge weights where simpler algorithm could be developed. Correctness and proof of greedy properties in general or in specific (restricted graphs).

Dijkstra's Algorithm is a greedy algorithm that solves the shortest path problem for a weighted directed graph with non-negative edge weights, producing a shortest path tree. It works by maintaining a set of vertices whose shortest distance from the source node is know, and a set of vertices whose shortest distances in unknown. At each step, it selects the vertex with the smallest known distance from the source node and updates the distances of its adjacent vertices if a shorter path is found.

Here's the pseu

```
* (outo is = adj(u).legic(); it = adj(u).enf(); ++it) {
int v = ito-first;
int velget = ito-second;
};
{
// (first)
```

### 5. Divide and Conquer Algorithms:

5.1. Formulation of recurrence and the Master Theorem. Deriving a recurrence from a code and "solving" it. How to improve time complexity by analysis of the terms in the Master Theorem for a specific code example, or recurrence example. For instance, you will need to know if changing a specific section of the code, will lead to a change in its asymptotic complexity, or by how much you need to make a change in which section of the recursive function implementation that will lead to changes in the time complexity of the overall algorithm.

Divide and conquer algorithms are a class of algorithms that solve a problem by breaking it down into smaller subproblems, solving those subproblems independently, and then combining the solutions to the subproblems to obtain the solution to the original problem. The key idea is to divide the problem into smaller subproblems that are easier to solve, and then combine the solutions to these subproblems to obtain the solution to the original problem.

One of the most important aspects of drivide and conseque algorithms is the formulation of a recurrence relation that describes the running time of the algorithm. The recurrence relation expresses the running time of the algorithm as a function of the size of the input.

One of the most famous tools for analyzing the time complexity of divide and conquer algorithms is the Master Theorem which provides a way to classify the running time of a class of divide and conquer algorithms based on their recurrence relation.

```
octor arr(sco.),
} olso {
int mid = (start + end) / 2;
int left max = find_max(arr, s
int right max = find_max(arr, return max(left_max, right_max)
```

In this code, the find, max function takes an array arr and two indices start and end that specify the range of the array to seaffor for the maximum element. If the range consists of a single element, the function returns that element. Otherwise, the function recursive/calls itself on the left and right hardse of the array and returns the maximum of the two values.

To analyze the time complexity of this algorithm, we can use a recurrence relation. Let T(n) be the running time of the rithm on an array of size n. Then, we have:

# T(n) = 2T(n/2) + O(1)

rsively calls itself twice on subproblems of size n/2, and the time rehis is because the algorithm num of the two values is O(1).

To solve this recurrence relation, we can use the Master Theorem. The Master Theorem states that if a recurrence rela-

# T(n) = aT(n/b) + f(n)

 $T(n) = O(n \land log\_b(a))$ 

if  $f(n) = O(n \log_b(a))$ , and

T(n) = O(f(n))

 $if \ f(n) = Omega(n \land log\_b(a)).$ 

In our example, we have a = 2, b = 2, and f(n) = O(1), so we can apply case 2 of the Master Theorem, which gives us

 $T(n) = O(n \land log\_2(2)) = O(n)$ 

The Towers of Hanoi problem is a classic problem that involves moving a tower of disks of different sizes from one peg to another, with the constraint that only one disk can be moved at a time and a larger disk cannot be placed on top of a smaller disk. The problem can be showed using the recurring spapeous, where we recurrively more the smiller tower to an intermident peg, move the largest disk to the destination peg, and then move the smaller tower from the intermediate peg to the destination peg Hers's the penduchoods for the Towers of Hanoi algorithm:

The input arguments to the function are n, the number of disks in the tower, source, the peg the tower is currently on, tination, the peg we want to move the tower to, and intermediate, the third peg that we can use to move the tower.

The base case of the recursion is when we have only one disk left, in which case we simply move it from the source peg to the destination peg. In the recursive case, we first move the top =1 disks from the source peg to the intermediate peg using the destination peg, as the temporary peg. Then, we move the largest disks from the ource peg to the destination peg. Finally, we move the top =1 disks from the ource peg as the temporary peg. Then,

The time complexity of the Towers of Hanoi algorithm is  $O(2^n n)$ , since in the west case we need to make  $2^n n - 1$  moves to move a tower of a disks.