

ADVANCED ALGORITHMS THEORY SWISS KNIFE



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1 TABLE OF CONTENTS

2	Gr	Graph – General Definitions		
3	De	epth First Search - DFS7		
	3.1	Description7		
	3.2	Algorithm7		
	3.3	Complexity7		
	3.4	Applications		
4	Br	eadth First Search - BFS		
	4.1	Description9		
	4.2	Algorithm9		
	4.3	Complexity9		
	4.4	Applications		
5	Mi	inimum Spanning Tree – MST		
	5.1	Generic Greedy Algorithm		
	5.2	Definitions		
6	Pr	im's Algorithm		
	6.1	Description		
	6.2	Algorithm		
	6.3	Complexity		
	6.4	Example of Execution for Exam12		
7	Ef	ficient Prim – Heap Implementation13		
	7.1	Description		
	7.2	Algorithm		
	7.3	Complexity14		
8	Kr	uskal's Algorithm		
	8.1	Description15		
	8.2	Algorithm		
	8.3	Complexity		
	8.4	Example of Execution for Exam15		
9	Ef	ficient Kruskal – Union-Find		
	9.1	Description		
	9.2	Algorithm		
	9.3	Complexity16		
1	0	Shortest Path		

11	Single-Source Shortest Path (SSSP)	
12	Non-negative weights – Dijkstra	18
12.1	Description	18
12.2	Algorithm	18
12.3	Complexity	18
13	Efficient Dijkstra – Heap	19
13.1	Description	19
13.2	Algorithm	19
13.3	Complexity	19
14	General Case: SSSP Problem	20
15	Bellman-Ford's Algorithm	21
15.1	Description	21
15.2	Algorithm	21
15.3	Complexity	21
16	All-Pairs Shortest Paths (APSP)	22
17	Floyd-Warshall's Algorithm	23
17.1	Description	23
17.2	Algorithm	23
17.3	Complexity	23
18	Maximum Flows	24
19	Ford-Fulkerson's Algorithm	25
19.1	Description	25
19.2	Algorithm	26
19.3	Complexity	26
20	NP-Hardness	27
20.1	NP-Hard Problems	28
20.2	NP-Hard Proofs	28
21	Approximation Algorithms	36
21.1	Examples of Approximations	37
22	TSP & Metric TSP	39
22.1	Travelling Salesperson Problem (TSP)	39
22.2	Metric TSP	39
22	2.2.1 Metric TSP is NP-Hard	39
22.3	2-Approximation Algorithm for Metric TSP	41
22.4	3/2 (or 1.5) Approximation Algorithm for Metric TSP	42

23	Set Cover	46
23.1	Set Cover is NP-Hard	46
23.2	Greedy Approximation Algorithm	47
24	Randomized Algorithms	48
24.1	Classification of Randomized Algorithms	48
24.2	Karger's Algorithm for Minimum Cut	49
25	Chernoff Bounds	51
25.1	Chernoff Bound Variants	51
25.2	Analysis in High Probability of Randomized Quicksort	51

Disclaimer

This file contains basically a refined version of full notes (and of my full theory file) to summarize the theory content and make it clearly visible if possible in a proper way, concise but understandable. This respects and follows, both chronologically and logically, the topics seen in 2023/2024.

2 **GRAPH – GENERAL DEFINITIONS**

- G = (V, E) as the graph itself
 - V = set of <u>vertices</u> (aka nodes)
 - $\circ \quad E \subseteq V \ x \ V$ (cartesian product = all) is a collection of <u>edges</u>
 - an edge is a pair of vertices (u, v)
 - it indicates the connection between two nodes
 - a connection of vertices allows for repetition
- <u>directed</u> graphs, which happens if $(u, v) \neq (v, u)$
- <u>undirected</u> graphs, which happens if (u, v) = (v, u)
- <u>arc</u> = edge inside directed graphs (also called *directed edges*)
- given an edge e = (u, v)
 - \circ e is <u>incident</u> on u and v (happens if vertex if one of endpoints in that edge)
 - \circ u and v are <u>adjacent</u> (there is an edge between the two vertices)
- <u>neighbors</u> of a vertex: all vertices v s.t. $(u, v) \in E$
 - o all vertices directly connected to a given vertex by an edge
- <u>degree</u> of a vertex v, denoted as d(v) or degree(v)
 - \circ the number of edges incident on v
- <u>path</u>: $u_1, u_2 \dots u_k$ and $(u_i, u_{i+1}) \in E, \forall \ 1 \le i \le k$
 - \circ finite/infinite sequence of nodes which joins a sequence of vertices via edges
- simple path: u_i (all vertices) are all distinct
 - \circ ~ same definition as above and vertices/nodes are all distinct/so are the edges
 - o e.g., 5,1,8,7,6,1,4 has 1 repeated twice so it's not simple
- <u>cycle</u>: simple path s.t. $u_1 = u_k$ (starts from a given vertex/ends at same node)
- <u>subgraph</u>: G' = (V', E') s. t.
 - $\circ \quad V' \subseteq V$
 - $\circ \quad E' \subseteq E$
 - \circ ~ the edges of E' are incident only on vertices of V'
 - $\circ \quad$ in words: it is a subset of the larger original graph
- <u>spanning subgraph</u>: a subgraph with V' = V
 - $\circ~$ a subgraph which "spans" the original graph (so there are all the vertices)
 - o following other definitions
 - subgraph obtained by edge deletions only but retaining all vertices
 - so it's a subgraph of *G* with same vertex set as *G*
- <u>connected graph</u>: if $\forall u, v \in V$, \exists a path from u to v
- <u>connected components</u>: a partition of *G* in subgraphs $G_i = (V_i, E_i), \forall 1 \le i \le k \ s. t.$
 - \circ G_i is connected $\forall i$
 - $\circ \quad V = V_1 \cup V_2 \cup \ldots \cup V_k$
 - $\circ \quad E = E_1 \cup E_2 \cup \dots \cup E_k$
 - $\circ \quad \forall i \neq j$ there is no edge between V_i and V_j
- <u>tree</u>: connected graph without cycles
 - o any two vertices are connected by *exactly* one path
- forest: set of trees (disjoint)
 - o also = undirected graph in which any two vertices are connected by at most one path

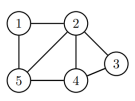
- spanning tree: a spanning subgraph connected and without cycles
- <u>spanning forest</u>: a spanning subgraph without cycles

Generally, remember:

- n = |V| (number of nodes)
- m = |E| (number of edges)
- the size of a graph is n + m

There are also multiple ways of representing:

- an adjacency list
 - an array A of n lists, one \forall vertex $v \in V$ (consider the example below)
 - \circ each containing all the vertices adjacent to v (represented by table below)



1	2,5
2	1,3,4,5
3	2,4
4	2,5,3
5	4,1,2

What if directed? Only vertices pointed for that vertex.

- Pro: space usage $\theta(n+m)$ i.e. <u>linear</u>
- Con: no quick way to determine if a given edge is in the graph
- an <u>adjacency matrix</u>

• a
$$n \times n$$
 matrix A s.t. $A[i,j] = 1$ if $edge(i,j) \in E$, 0 otherwise

- If graph is directed \rightarrow the matrix is asymmetric
- If graph is undirected \rightarrow the matrix is symmetric

In case of a weighted graph, each cell of the matrix has either the value of the edge weight (as number) w or -/null to represent null costs. This kind of graph represents costs, capacities, etc.

- Pro: Quick to determine if a given edge is present
- Con: Space required is $\theta(n^2) \rightarrow$ can be superlinear in the input size
 - o if number of vertices increases, the space required by matrix grows quadratically

3 DEPTH FIRST SEARCH - DFS

3.1 DESCRIPTION

The algorithm starts at the root node (selecting some arbitrary node as the root node in the case of a graph) and explores as far as possible along each branch before backtracking. It may find:

- new edges (discovery edge)
- non-tree edges, linked to ancestors (back edges)

Visit a vertex, then a neighbor of the vertex, then a neighbor of the neighbor – these are all neighbours, classified with adjacency lists.

3.2 ALGORITHM

procedure DFS(G,V)

visit v

 $L_V[v].ID = 1$

for all $e \in G.incidentEdges(v)$: do

if $L_E(e)$. label = null then

w = G.opposite(v, e)

if $L_V[w]$. ID = 0 then

 $L_E[e]$. label = DISCOVERY EDGE

else

$$L_E[e]$$
. label = BACK EDGE

3.3 COMPLEXITY

Given:

- n_s : number of vertices of C_s (one invocation $\forall v \in C_s$)
- m_s : number of edges of C_s (costs related to node, excluding recursive invocations inside)

The complexity overall is:

$$\theta\left(\sum_{v\in C_s}d(v)\right)=\theta(m_s)$$

More in general:
$$O(n + m) - n$$
 vertices and m edges

Written by Gabriel R.

There are several:

- s t path (between two generic vertices)
 - \circ done adding a *parent* field
- finding cycles
 - o use *parent* field on vertices and *ancestor* on edges
- find connected components
 - \circ run the algorithm *n* times
 - \circ $\,$ consider all untouched vertices
 - o see which have back edges, meaning they "close" the cycle
 - o otherwise, return
- find a spanning tree

4 BREADTH FIRST SEARCH - BFS

4.1 DESCRIPTION

The algorithm is iterative, starts from a source vertex and visits all vertices connected to a specific component, partitioning them in levels according to their distance. It still has discovery edges:

- but adds cross edges - which connect vertices at different levels

4.2	ALGORITHM
-----	-----------

```
procedure BFS(G, s)
       visit(s)
       L_{V}[s].ID = 1
       Create a set L_0 containing s
       i = 0
       while (!L_i.isEmpty) do:
               Create a set of vertices L_{i+1}
               for each v \in L_i do:
                       for each e \in G. incidentEdges(v) do:
                       if L_E[e]. label = null then
                               w = G.opposite(v, e)
                               if L_v[w]. ID = 0 then
                                       L_E[e]. label = DISCOVERY EDGE
                                       visit w
                                       L_V[w].ID = 1
                                       add w in L_{i+1}
                               else
```

 $L_E[e]$. label = CROSS EDGE

i = i + 1

4.3 COMPLEXITY

O(n+m)

- Same as for DFS in $\theta(n+m)$ time

So, again:

- s-t path (between two generic vertices)
 - \circ done adding a *parent* field
- finding cycles
 - use *parent* field on vertices and *ancestor* on edges
- find connected components
 - \circ run the algorithm *n* times
 - $\circ \quad \text{consider all untouched vertices}$
 - \circ ~ see which have back edges, meaning they "close" the cycle
 - o otherwise, return
- find a spanning tree

5 MINIMUM SPANNING TREE – MST

- Input: a graph G = (V, E) undirected, connected and weighted
 - A weight $w: E \to \mathbb{R}$
 - defines w(u, v) = cost of edge(u, v)
- Output: a spanning tree $T \subseteq E$ of G s.t. $w(t) = \sum_{u,v \in T} w(u,v)$ is minimized
 - \circ $\,$ Goal is minimizing the sum for all weights of every edge of the tree

5.1 GENERIC GREEDY ALGORITHM

procedure *Generic* – *MST*(*G*)

 $A= \emptyset$

while *A* does not form a spanning tree do:

find an edge (u, v) that is safe for A // crucial step

 $A = A \cup \{(u, v)\}$ // add vertices to A

return A // A is an MST

5.2 **DEFINITIONS**

- A <u>cut</u> of graph G = (V, E) is a partition of $V \to (S, V \setminus S)$
 - \circ $\;$ in words, a partition of vertices into two disjoint subsets
 - o it can be done on one or more edges
- An edge $(u, v) \in E$ <u>crosses</u> a cut $(S, V \setminus S)$ if $v \in S$ and $v \in V \setminus S$ (or viceversa)
 - \circ so, if its endpoints lie in different subsets of the partition defined by the cut
- A cut <u>respects</u> a set of edges A if no edge of A crosses the cut
- Given a cut, an edge that crosses the cut and is of minimum weight is called <u>light edge</u> (for that cut) → they are useful, because when included in MSTs, they have minimum weight

There is also the *minimum cut*, for which we have $d(v) \ge t \ \forall v \in V$, where t is a generic size of graph. Summing up all n vertices, we obtain $\sum_{v \in V} d(v) \ge tn$, concluding it's $\sum_{v \in V} d(v) = 2m$.

6 PRIM'S ALGORITHM

6.1 **DESCRIPTION**

The algorithm is iterative and selects light edges at every step, growing a spanning tree from there. Consider this gif to see the running. We have to preserve "safe edge" property – take only minimal edges not already inside of the tree.

6.2 ALGORITHM

```
procedure Prim(G,S)
```

 $X = \{S\}$

 $A = \emptyset$

while there is an edge (u, v) with $u \in X$ and $v \notin X$ do:

 $(u^*, v^*) = a$ minimum weight such edge (aka light edge)

add vertex v^* to X

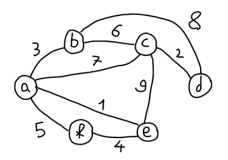
add edge (u^*, v^*) to A

return A

6.3 COMPLEXITY

O(m * n)

6.4 EXAMPLE OF EXECUTION FOR EXAM



Traversal: (a, e), (a, b), (e, f), (b, c), (c, d)

7 EFFICIENT PRIM – HEAP IMPLEMENTATION

7.1 DESCRIPTION

The previous is not so efficient in large structures. The right kind of data structure to improve the algorithm is a *priority queue*, implemented with a <u>heap</u>.

- Recap about this data structure
 - \circ insert \rightarrow add an object to the heap (possibly fast)
 - \circ extractMin \rightarrow remove an object with the smallest key (highest priority)
 - \circ delete \rightarrow given a pointer to an object, remove it
- In a heap with n objects, the complexity of these operations is O(log(n))

We can redefine the algorithm exploiting this efficient data structure basically with the same principle:

- consider a min heap starting from whatever vertex, which is the root
- from there, always extract the minimum value (means checking if it is min heap),
 - \circ then update the path

7.2 ALGORITHM

procedure Prim (G, s)

for each $v \in V$: do $key[u] = +\infty$ $\pi(v) = NULL // \pi = parent of v in the tree being built$ Key[s] = 0 H = Vwhile $H \neq 0$ do: $v^* = extractMin(H)$ for each v adjacent to v*: do if $v \in H$ and $w(v^*, v) < Key(v)$ then $\pi(v) = v^*$ delete v from H $Key(v) = w(v^*, v)$

- init $\rightarrow O(n)$
- while $\rightarrow n$ iterations
- $extractMin \rightarrow O(\log(n))$

Total cost of only *extractMin* operations: $O(n \log(n))$

- for loop: executed O(m) times in total (every vertex is explored)
 - $\circ \quad v \in H \to O(1)$
 - this here is a simple check
 - $Key(v) = w(v^*, v) \rightarrow delete + insert: O(2 \log(n)) = O(\log(n))$
 - two operations

Total cost of for loop: $O(m \log(n))$ (iterating for all adjacent nodes, quantity equal to node degree)

This way, the total complexity of the algorithm is $O(n \log(n) + m \log(n)) = O(m \log(n))$ (since G is connected, we recall) \rightarrow near-linear time.

8 KRUSKAL'S ALGORITHM

8.1 DESCRIPTION

It picks the minimum weighted edge at first and the maximum weighted edge at last. It sorts edges by weight and then adds them continuously, preserving the "safe edge" property – take only the unexplored. It does so preventing the adding of cycles.

8.2 ALGORITHM

procedure Kruskal(G)

 $A = \emptyset$

Sort safe edges of G by weight

for each edge e in non – decreasing order of weight: do

if $A \cup \{e\}$ is acyclic then:

 $A = A \cup \{e\}$

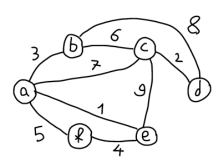
return A

8.3 COMPLEXITY

- sorting: $O(m \log(n))$
- for loop: check whether e = (u, v) closes a cycle is equivalent to check whether A contains an u v path \rightarrow DFS on $G = (V, A) \rightarrow$ complexity: O(n)

Total: $O(m * n) \rightarrow O(m \log(n)) + O(m * n) = O(m * n)$

8.4 EXAMPLE OF EXECUTION FOR EXAM



Traversal: (a, e), (c, d), (a, b), (e, f), (b, c)

9 EFFICIENT KRUSKAL – UNION-FIND

9.1 DESCRIPTION

It can be implemented as fast as Prim's, considering the most frequent operation here is cycle check (equivalently, path check), which happens when an edge is added to *A*.

We create a new data structure supporting this operation fast and to do that, we use a structure called <u>Union-Find</u> (also called *disjoint set*). This is a structure to merge *disjoint sets* (also non-overlapping in their elements) of objects and supports at least three operations:

- Init: given an array X of objects
 - it creates a Union-Find data structure with each object x ∈ X in its own set
- *Find*: given an object *x*, return the name of the set that contains *x*
 - o depth: number of edges traversed by *Find*
- Union: given two objects x, y merge the sets that contain x and y into a single set
 - o done whenever the sets are distinct
 - \circ if x, y are already in the same set, this operation does nothing

9.2 ALGORITHM

procedure Kruskal(G)

 $A = \emptyset$

U = init(V)

sort edges of E by weight

for each edge e = (v, w) in non – decreasing order of weight: do

if $Find(v) \neq Find(w)$ then:

 $A = A \cup \{(v,w)\}$

Union(v,w)

return A

9.3 COMPLEXITY

- Init: *O*(*n*)
- Sorting: $O(m \log(n))$
- 2m Find: $O(m \log(n))$
- n-1 Union: $O(n \log(n)) \rightarrow$ only when I go inside an "if" and when the edge is added
- A updating: O(n)

In total: $O(m \log(n))$

10 SHORTEST PATH

- Given a weighted graph, the <u>length</u> of a path $p = v_1, v_2, \dots v_k$ is defined as $len(P) = \sum_{i=1}^{k-1} w(v_i, v_{i+1})$
- A <u>shortest path</u> from a vertex u to a vertex v is a path with minimum length among all u v paths
- The <u>distance</u> between 2 vertices *s* and *t*, denoted as dist(s, t) is the length of a shortest path from *s* to *t*; if there is no path at all from *s* to *t* then $dist(s, t) = +\infty$

The problem itself is the following:

- Given a directed, weighted graph and a source vertex $s \in V$ and a destination $t \in V$, compute the shortest path from s to v

11 SINGLE-SOURCE SHORTEST PATH (SSSP)

- input: a directed, weighted graph G with edge weights $w: E \to \mathbb{R}$ and a source vertex $s \in V$
- output: $dist(s, v), \forall v \in V$
 - \circ shortest path to all destinations

There are two major cases to solve: a special one and a more general one.

12 NON-NEGATIVE WEIGHTS – DIJKSTRA

12.1 DESCRIPTION

Dijkstra's algorithm finds the shortest path from one vertex to all other vertices. It does so by repeatedly selecting the nearest unvisited vertex and calculating the distance to all the unvisited neighboring vertices.

- input: directed $G, s \in V, w: E \to \mathbb{R}_{\geq 0}$
- output: $dist(s, v) = len(v), \forall v \in V$
 - with len(v) coming as shorthand form of the previous one

12.2 ALGORITHM

```
procedure Dijkstra(G,s)
```

 $X = \{s\}$

len(s) = 0

 $len(v) = \infty$

while there is an edge (v, w) with $v \in X$ and $w \notin X$: do

 $(v^*, w^*) = such an edge minimizing len(v) + w(v, w)$

add w* to X

 $len(w^{*}) = len(v^{*}) + w(v^{*}, w^{*})$

12.3 COMPLEXITY

O(m * n)

13 EFFICIENT DIJKSTRA – HEAP

13.1 DESCRIPTION

Normal implementation uses adjacency list. This implementation improves efficiency by using a priority queue (usually implemented as a binary heap) to select the vertex with the smallest tentative distance efficiently. The implementation is almost identical to Prim with heaps.

13.2 ALGORITHM

```
procedure Dijkstra(G, s)
                               (almost identical to Prim's implementation with heaps)
      X = \{s\}
      H = \emptyset
      key(s) = 0
      for each v \neq s: do
                key(v) = \infty
      for each v \in V: do
                insert v into H
      while H is non – empty: do
                w^* = extractMin(H)
                add w^* to X
                len(w^*) = key(w^*)
                for each edge (w^*, y) s.t. y \notin X: do
                        delete y from H
                        key(y) = \min \{key(y), len(w^*) + w(v^*, w^*)\}
                        insert y into H
```

13.3 COMPLEXITY

- considering graph as adjacency list, n vertices and m edges
- log(n) iterations because of heap usage

Total number of operations: $O((n+m)\log(n))$

(there are O(n + m) operations on heaps)

14 GENERAL CASE: SSSP PROBLEM

We reformulate the previous problem a bit:

- Input: a directed weighted graph G = (V, E) and a source vertex $s \in V$
- Output: one of the following
 - dist(s, v) \forall vertex $v \in V$
 - o a declaration that G contains a negative cycle

Need to forbid negative cycles in shortest paths, they lead to infinitely small paths, which is an NP-Hard problem.

The main Dijkstra problems are two:

- It never revisits/updates its decisions, but it should for all vertices!
 - o Once a vertex is marked as "closed", we will never develop this node again
 - If we have a vertex in open such that its cost is minimal by adding any positive number to any vertex the minimality will never change
 - Without the constraint on positive numbers the above assumption is not true
 - It assumes them to be positive to make the algorithm run faster and does this to avoid considering paths which can't be shorter
- len(v) should be an estimated distance, which needs to be updated for every vertex
 - how many times? $\leq n 1$ edges $\Rightarrow n 1$ times should be enough
 - \circ $\$ maximum number of edges in a simple path between any two vertices

15 BELLMAN-FORD'S ALGORITHM

15.1 DESCRIPTION

- Input: A directed graph G with edge weights $w: E \to \mathbb{R}$ and a source vertex $s \in V$
- Output: Either $dist(s, v) \forall v \in V$ or a declaration that G contains a negative cycle

The algorithms is used when the graph might possess negative weights and can even detect negative cycles. If the graph contains one, there is no cheapest path, instead one can make it cheaper by one more walk around said negative cycle (in n - 1 iterations it reaches a fix-point, if it doesn't it means a negative cycle exists). Still, it's slower compared to Dijkstra.

15.2 ALGORITHM

procedure Bellman - Ford(G, s) len(s) = 0 $len(v) = \infty \forall v \neq s$ for n - 1 iterations do for each $edge(u, v) \in E$: do $len(v) = min\{len(v), len(u) + w(u, v)\}$ for each $edge(u, v) \in E$: do if len(v) > len(u) + w(u, v) then return "G contains a negative cycle"

15.3 COMPLEXITY

O(m * n)

16 ALL-PAIRS SHORTEST PATHS (APSP)

- Input: A directed, weighted graph G = (V, E)
- Output: One of the following:
 - \circ dist(u, v) \forall ordered vertex pair
 - \circ a declaration that G contains a negative cycle
 - this can be problematic in finding a shortest path
 - now we would have to output n^2 shortest paths

Consider:

- If we use Bellman-Ford <u>very</u> high complexity $\rightarrow O(m * n^2)$
 - Using dynamic programming, the complexity can be reduced to $O(n^3 \log(n))$
 - \circ $\;$ This holds rewriting B-F recurrence controlling the allowable size of the input $\;$

17 FLOYD-WARSHALL'S ALGORITHM

17.1 DESCRIPTION

It's used to find the shortest paths between all pairs of nodes in a weighted graph, with positive or negative edges.

- instead of restricting the number of edges allowed in a solution, restrict the <u>identities of the</u> <u>vertices</u> that are allowed in a solution
 - \circ $\;$ in other words, now paths can pass through only certain vertices
- Basically, it compares many possible paths through the graph between each pair of vertices

It iterates on 3 vertices: *u*, *v*, *k* in 3 nested loops, testing whether using *k* in the path is better.

17.2 ALGORITHM

procedure *Floyd* – *Warshall(G)*

label the vertices $V = \{1, 2, ..., n\}$ arbitrarily

A = n x n x (n+1) array

for u = 1 to n: do

for v = 1 to n: do

if u = v then A[u, v, 0] = 0

else if $(u, v) \in E$ then A[u, v, 0] = w(u, v)

else
$$A[u, v, 0] = +\infty$$

for k = 1 to n: do

for u = 1 to n: do

for v = 1 to n: do

$$A[u, v, k] = \min \{A[u, v, k-1], A[u, k, k-1] + A[k, v, k-1]\}$$

for u = 1 to n: do

if A[u, u, n] < 0 then return "*G* contains a negative cycle"

17.3 COMPLEXITY

 $0(n^{3})$

18 MAXIMUM FLOWS

- a <u>flow network</u> is a directed graph G = (V, E) where each edge has a <u>capacity</u> $c(e) \in \mathbb{R}^+$, along with a designated <u>source</u> $s \in V$ and <u>sink</u> $t \in V$
 - for convenience, write c(e) = 0 if $e \notin E$, no edges enter s and no edges leave t
- a <u>flow</u> is a function $f: E \to \mathbb{R}^+$ satisfying the following constraints (how much stuff I send through the edges in general)
 - (capacity) $\forall e \in E, f(e) \leq c(e)$ value of the flow at most capacity of that edge
 - (conservation) $\forall u \in V \setminus \{s, t\}$ we have

$$\sum_{u \in V \text{ s.t.}(v,u) \in E} f(v,u) = \sum_{v \in V \text{ s.t.}(u,v) \in E} f(u,v)$$

- the amount of flow going in nodes must be equal to the flow going out from those (conservation of flows)
 - initially, such flow is 0, which is "how much we can pass on the edge"
- the <u>value</u> of a flow is

$$|f| = \sum_{v \in V \text{ s.t.}(s,v) \in E} f(s,v)$$

- \circ $\;$ basically, the sum of all flows going in and out vertices thanks to edges $\;$
- o as a matter of fact, the amount of stuff traveling from source to sink
- o such flow has to be less than or equal to the capacity

As for the problem itself:

- given a flow network, find a flow *f* of maximum value. Such flow is measured on *the maximum* value received in a sink node

19 FORD-FULKERSON'S ALGORITHM

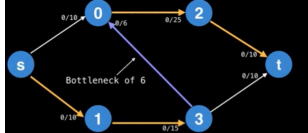
19.1 DESCRIPTION

Given a flow network G a flow f, the <u>residual network</u> of G w.r.t (with respect to) flow f, G_f , is a network with vertex set V and with edge set E_r as follows:

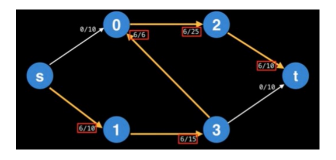
- for every edge e = (u, v) in G
 - if f(e) < c(e), add e to G_f with capacity $C_f(e) = c(e) f(e)$
 - o if f(e) > 0, add another edge (v, u) to G_f with capacity $C_f(e) = f(e)$

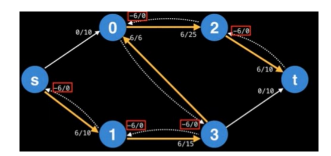
The Ford-Fulkerson (F-F) algorithm repeatedly finds an s - t path P in G_f (e.g., using BFS) and uses P to increase the current flow.

- *P* is called <u>augmenting path</u>
 - This is a path of edges in the residual graph with unused capacity greater than 0 from the source s to the sink t
 - This can only flow on edges not fully saturated yet
- In an augmenting path, the *bottleneck* is the smallest edge on the path. We can use this one to _ augment the flow along the path



- - Augmenting the flow means updating the flow values along the augmenting path (left)
 - For forward edges, this means increasing the flow by the bottleneck value
 - When augmenting the flow along the augmenting path
 - o you also need to decrease the flow along each residual edge (backward edges) by the bottleneck value (right)
 - o residual edges exist to "undo" bad augmenting paths which do not lead to a maximum flow





In figure below, in orange the augmenting path, in light-blue as written the bottleneck:

The residual graph, so, contains also residual edges. This algorithm continues to find augmenting paths and augments the flow until no more augmenting paths exist.

- The algorithm simply takes in every iteration the bottleneck
- Then considers the bottleneck and keeps incrementing selecting every possible s t path until max flow is reached
- Each iteration is then reported into the residual graph, accounting for the bottleneck
 - \circ e.g. if we chose 7/11 in an iteration, in the next iteration
 - 4 forward (remaining)
 - 7 backward (spent)

19.2 ALGORITHM

```
procedure Floyd - Warshall(G)
```

```
label the vertices V = \{1, 2, ..., n\} arbitrarily
```

A = n x n x (n+1) array

for u = 1 to n: do

for v = 1 to n: do

if u = v then A[u, v, 0] = 0

else if $(u, v) \in E$ then A[u, v, 0] = w(u, v)

else
$$A[u, v, 0] = +\infty$$

for k = 1 to n: do

```
for u = 1 to n: do
```

for v = 1 to n: do

 $A[u, v, k] = \min \{A[u, v, k-1], A[u, k, k-1] + A[k, v, k-1]\}$

for u = 1 to n: do

if A[u, u, n] < 0 then return "*G* contains a negative cycle"

19.3 COMPLEXITY

- Assume capacities are integers; then
 - \circ the flow value increases by ≥ 1 is each iteration
 - the complexity of each iteration is O(m)

Total complexity is $O(m * |f^*|)$, where f^* is a max flow

20 NP-HARDNESS

There are problems which can be solved in linear time:

- e.g. Eulerian circuit – a graph where edges are traversed all at once

There are problems which can be solved in polynomial time:

- e.g. Minimum Spanning Tree (MST), minimizing the weights inside all tree

There are also problems where no polynomial algorithms are present to solve the problem:

- e.g. Traveling Salesperson Problem (TSP), Hamiltonian Circuit

We define the following *complexity classes*:

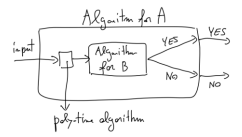
- 1) *P* is the set of decision problems that can be solved in polynomial time
- 2) *NP* is the set of decision problems with the following property:
 - a. if the answer is YES, then there is a proof of this fact (called "certificate") that can be checked in polynomial time
- 3) co NP, which is essentially the opposite of NP:
 - b. property: if the answer is NO, then there is a proof of this fact that can be checked in polynomial time

Other features of problems:

- a problem is said to be <u>NP-Hard</u> if a polynomial time algorithm for this one would imply the existence of a polynomial time algorithm for every problem in NP
- More formally, a problem is <u>NP-Hard</u> if every problem in NP reduces in polynomial time to it
 - a. unless P = NP, which is not yet solved
 - b. if a problem is NP-Hard, it provides evidence the problem may not be in P
- a problem is <u>NP-Complete</u> if it's both in NP and NP-Hard
 - a. e.g. the Cook-Levin Theorem for Boolean Satisfiability problem (SAT)
 - i. made up of clauses with conjunction/disjunction, usually 3 (3-SAT)

We use a <u>reduction</u> given it's a very powerful tool:

- a reduction is an algorithm for transforming one problem into one another
- a problem *A* reduces to *B* if there is an algorithm able to solve *B* can be translated into one which solves *A*
- remember reductions works from Y (problem I know to be hard) to X (new problem)
- the reduction is FROM Y
 - a. I already know it's NP-hard
- to X
 - a. the "new" problem



- Independent Set
 - a. given a graph G = (V, E) an <u>independent set</u> in G is a subset $I \subseteq V$ with no edges between them
- <u>(Maximum) Independent Set</u> (this one will be referred to as simply "Independent Set" meaning the latter)
 - a. compute an independent set of maximum size
- <u>SAT/3-SAT</u> (thanks to Cook-Levin Theorem)
 - a. SAT Boolean satisfiability of a formula (has to be equal to TRUE)
 - b. 3-SAT Boolean satisfiability of a formula made by 3-clauses
- Hamiltonian Circuit
 - a. a cycle that traverses all the vertices only once
- (Maximum) Clique
 - a. largest complete subgraph
- (<u>Minimum) Vertex cover</u>
 - a. minimum number of vertices that "touches" all edges

Examples of reductions:

- Using Hamiltonian circuit to solve TSP \rightarrow Ham $\leq_p TSP$
 - a. If we had a fast algorithm for TSP, we would also solve Hamiltonian problem
- Using 3SAT to solve Independent Set \rightarrow 3SAT \leq_p IndependentSet
 - a. If we had a fast algorithm for Independent Set, we would also solve 3SAT
- Using Independent Set to solve Clique \rightarrow Clique \leq_p IndependentSet
- Using Independent Set to solve Vertex Cover \rightarrow Vertex Cover \leq_p IndependentSet

20.2 NP-HARD PROOFS

- Theorem: TSP (Traveling Salesperson Problem) is NP-Hard
- *Proof*: Reduction from Hamiltonian circuit to TSP ($Ham \leq_p TSP$)

Wait a minute: TSP is not a decision problem!

No worries. Define *TSP* as:

- input: G = (V, E) complete, undirected, weighted graph $k \in \mathbb{R}$
- output: \exists in *G* a Hamiltonian circuit of cost $\leq k$?

We could try to use all possible k values, but k is not guaranteed to be polynomial; using only the cycles will not work either.

What we actually do: Pick an arbitrary input instance for Ham. and create the following input for TSP:

- G' = (V, E') complete, undirected, weighted graph with:

$$w(e \in E') = \begin{cases} 1, & \text{if } e \in E \\ +\infty, & \text{otherwise} \end{cases}$$

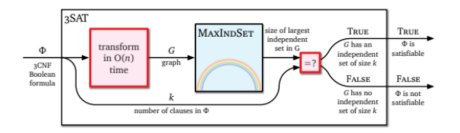
If we use k = m this reduction takes poly-time (($O(n^2)$). Then:

- if G has an Hamiltonian circuit, then the TSP algorithm run on G' returns an Hamiltonian circuit with cost n
- if G doesn't have a Hamiltonian circuit, then any Hamiltonian circuit in G' must have ≥ 1 edge not in G, hence of weight ∞. Hence, in this case, a TSP algorithm run on G' returns a Hamiltonian circuit of cost > n

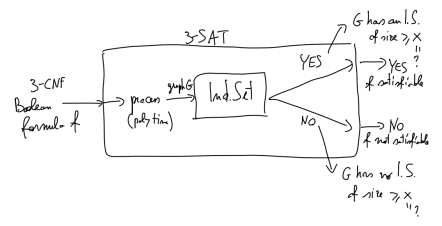
If we had a fast algorithm for TSP we would also solve the Hamiltonian circuit problem.

Reduction from 3SAT (problem in logic) to *Independent Set* (problem in graphs) \rightarrow 3SAT \leq_p IndependentSet

They seem totally unrelated problems, but let's see what we have to do (figure here is from "Algorithms" book of Jeff Erickson, suggested in particular for the whole NP-Hardness chapter):



What we are conjecturing is the following:



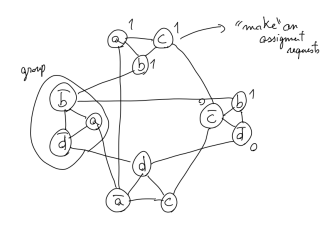
Basically, the presence of an independent set in the constructed graph corresponds to a satisfying truth assignment for the 3SAT instance.

Let's see the main ideas (figure representing the scenario):

- pick an arbitrary 3CNF Boolean formula f with k clauses

$$(a \lor b \lor c) \land (b \lor \neg c \lor \neg d) \land (\neg a \lor c \lor d) \land (a \lor \neg b \lor \neg d)$$

- vertices: each vertex represents one literal in f
 - a. a group of 3 vertices represents a clause (one of the k clauses)
 - i. assignment request = choose vertices and make a request



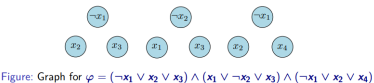
- edges:
 - a. We add an edge between a literal and its inverse, for all the literals
 - b. We add an edge between every pair of vertices that are in the same group

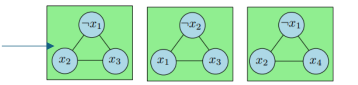
There are two ways to think about 3SAT: (this reasoning coming from here)

- 1. Find a way to assign 0/1 (false/true) to the variables such that the formula evaluates to true, that is each clause evaluates to true
- 2. Pick a literal from each clause and find a truth assignment to make all of them true. You will fail if two of the literals you pick are in conflict, i.e., you pick x_i and $\neg x_i$

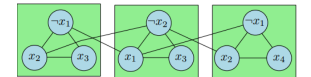
The reduction works this way:

- the graph will have one vertex for each literal in a clause
- connect the 3 literals in a clause to form a triangle; the independent set will pick at most one vertex from each clause, which will correspond to the literal to be set to true





- connect 2 vertices if they label complementary literals; this ensures that the literals corresponding to the independent set do not have a conflict



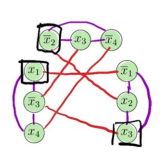
- Take k to be the number of clauses, ensuring they are all "covered"

Remember what *satisfiable* means:

- it asks whether the variables of a given Boolean formula can be consistently replaced by the values TRUE or FALSE in such a way that the formula evaluates to TRUE

 $\varphi = (x_1 \vee \overline{x}_3 \vee x_4) \land (\overline{x}_2 \vee x_3 \vee \overline{x}_4) \land (\overline{x}_1 \vee x_2 \vee x_3)$

- 1. Create a vertex for each literal.
- 2. Connect each literal to the other two literals in the same clause.



- 3. Connect each literal x_i to \overline{x}_i .
- 1) idea: independent set represents $conflicts \Rightarrow add$ an edge between every pair of vertices that are inconsistent (*asking* for opposite assignments to the same variable)
 - a. in words: if you choose one vertex, it means it's part of a clause
 - b. you have to choose other two vertices which are sure to be different because they are a different independent set
 - c. if you choose one vertex, you <u>have</u> to choose the complement
 - i. in order to realize the AND inside the formula

Observation: an independent set with ≥ 1 vertex in each group gives a satisfying truth assignment \rightarrow should look for indipendent sets of size $\geq k$ to say "YES, f it's satisfiable".

Issue: an independent set now is free to recruit <u>multiple</u> vertices from a group, so I might output "YES, f is satisfiable" even if this not true! \Rightarrow idea: force the recruitment of <u>one</u> vertex per group.

2) add one edge between every pair of vertices that one in the same group

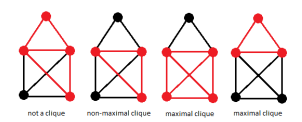
Claim: G contains an independent set of size exactly $k \Leftrightarrow$ the formula f is satisfiable

Proof:

- 1) suppose f is satisfiable. Pick any satisfying assignment. Each clause in f has $\geq 1 TRUE$ literal. Thus, we can choose a subset S of k vertices in G that contains exactly one vertex per group such that the corresponding k literals are all TRUE. The set S is an independent set because it does not contain both endpoints of any edge of a group, nor of any edge that connects inconsistent literals (as it is derived from a consistent truth assignment)
- 2) suppose *G* contains an independent set of size *k*. Each vertex in *S* must be in a different group. Assign *TRUE* to each literal of *S*. Since inconsistent literals are connected by an edge, this assignment is consistent. Since *S* contains 1 verftex per group, each clause in *f* contains (at least) one *TRUE* literal \Rightarrow *f* is satisfiable

- (Maximum) Clique: compute the longest complete subgraph
 - a. other name for a complete graph (from now on, the problem will be called Clique)
 - b. below, a useful figure to clearly see the problem

Show that Clique is NP-Hard.

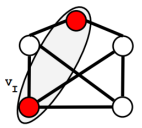


Solution (a nice graphical explanation here)

Decision version:

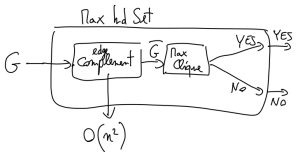
- Input: < G = (V, E), k >
- Output: \exists in *G* a clique of size *k*?

We operate a reduction from Maximum Independent Set (Ham. circuit is not really related to it; as you can see <u>here</u>, one can use 3SAT in order to show Clique is NP-complete). Figure here shows Independent Set.



- Intuition
 - a. clique: vertices with <u>all</u> edges between them
 - b. maximum independent set: vertices with no edges between them
- Definition
 - a. given a graph G = (V, E), its <u>edge-complement</u> $\overline{G} = (V, \overline{E})$ has the same vertex V and an edge set \overline{E} such that $(u, v) \in \overline{E} \Leftrightarrow (u, v) \notin E$ (so, no common edges)
- Observation
 - a. a set of vertices S is independent in $G \Leftrightarrow S$ is a clique in $\overline{G} \Rightarrow$ the largest independent set in G has the same size as the largest clique in \overline{G}

To make it super complete, let's draw the schema of what we are doing – takes $O(n^2)$ time, given the constant work needed to traverse all edges *and* vertices:



Definition: a <u>vertex cover</u> of a graph is a set of vertices that includes that at least one endpoint of every edge of the graph

b. Side figure represents such, to be clearer to you

Another problem is:

<u>(Minimum) Vertex Cover</u>: compute the smallest vertex in a given graph
 a. From now on, only called Vertex Cover

Show that Vertex Cover is NP-Hard.

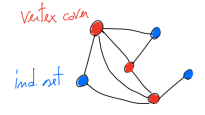
<u>Solution</u> (once again, a nice graphical explanation of this one <u>here</u>)

Decision version:

- Input: < G = (V, E), k >
- Output: \exists in *G* a vertex cover of size *k*?

We operate a reduction from Maximum Independent Set (once again, this is the most similar problem to the one we are proving)

- Observation
 - a. a set of vertices S is independent in $G \Leftrightarrow V \setminus S$ is a vertex cover of G
 - i. in blue there is an independent set (actually the biggest one)
 - ii. the other ones are the vertex cover

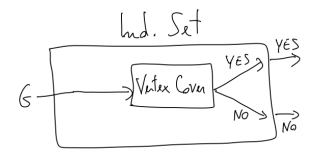


 \Rightarrow the longest independent set in G has size n-k , where k is the size of the smallest vertex cover of G

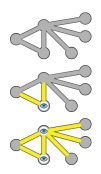
Independent set:

- Input: < G = (V, E), n k >
- Output: \exists in *G* an independent set of size n k?

Once again, let's represent this in a complete way:



Exercise



Written by Gabriel R.

- Show that:
 - a. Vertex Cover \leq_p Independent Set
 - b. Clique \leq_p Vertex Cover

 \Rightarrow these 3 problems are equivalent.

Solution (official = shorter)

- "same" as Independent Set \leq_p Vertex Cover
- we can consider the following figure for this one
 - a. consider a clique of size 4 in the middle (left)
 - b. if you take the complement of this one (right)
- *G* has a clique of size $k \Leftrightarrow \overline{G}$ has a vertex over of size n k
 - a. proof: see the book (§ p. 1106 of 4th edition theorem 34.12)



Solution (longer and better explained)

a. Suppose that we have an efficient algorithm for solving Independent Set, it can simply be used to decide whether G has a vertex cover of size at most k, by asking it to determine whether G has an independent set of size at least n - k

Given an instance of the Vertex Cover problem, consisting of a graph G = (V, E) and an integer k representing the size, we construct an instance of the Independent Set problem as follows:

- 1. Let G' = G (i.e., the graph for the Independent Set instance is the same as the original graph G).
- 2. Let k' = |V| k (i.e., the target size of the independent set is the number of vertices in *G* minus the size of the vertex cover *k*).

To show that this reduction is correct, we need to prove the following:

- 1. If G has a vertex cover of size $\leq k$, then G' has an independent set of size $\geq k'$.
- 2. If G' has an independent set of size $\geq k'$, then G has a vertex cover of size $\leq k$.

Let's prove both (1) and (2):

- Suppose *C* is a vertex cover of size $\leq k$ in *G*. Then, the set $V \setminus C$ is an independent set in *G'* (since *C* covers all the edges, no two vertices in $V \setminus C$ can be adjacent). Furthermore, $|V \setminus C| \geq |V| k = k'$
- Suppose S is an independent set of size $\geq k'$ in G'. Then, the set $V \setminus S$ is a vertex cover in G (since S is independent, every edge must have at least one endpoint in $V \setminus S$). Furthermore, $|V \setminus S| \leq |V| k' = k$.
- b. To show that $Clique \leq_p Vertex Cover$, we need to provide a polynomial-time reduction from the Clique problem to the Vertex Cover problem. Here's one way to construct the reduction:

Given an instance of the Clique problem, consisting of a graph G = (V, E) and an integer k, we construct an instance of the Vertex Cover problem as follows:

- 1. Let G' = G (i.e., the graph for the Vertex Cover instance is the same as the original graph G).
- 2. Let k' = |V| k (i.e., the target size of the vertex cover is the number of vertices in *G* minus the size of the clique *k*).

To show that this reduction is correct, we need to prove the following:

- 1. If G has a clique of size $\geq k$, then G' has a vertex cover of size $\leq k'$.
- 2. If G' has a vertex cover of size $\leq k'$, then G has a clique of size $\geq k$.

Proof of (1): Suppose *C* is a clique of size $\geq k$ in *G*. Then, the set $V \setminus C$ is a vertex cover in *G'* (since *C* is a clique, every edge must have at least one endpoint in $V \setminus C$). Furthermore, $|V \setminus C| \leq |V| - k = k'$.

Proof of (2): Suppose *S* is a vertex cover of size $\leq k'$ in *G'*. Then, the set $V \setminus S$ is a clique in *G* (since *S* is a vertex cover, every edge must have both endpoints in $V \setminus S$, which means $V \setminus S$ is a clique). Furthermore, $|V \setminus S| \geq |V| - k' = k$.

21 APPROXIMATION ALGORITHMS

These kinds of algorithms are are efficient algorithms that find approximate solutions to optimization problems (in particular NP-hard problems) with *provable* guarantees on the distance of the returned solution to the optimal one. They solve problems not solvable in polynomial time using approximation.

An optimization problem can be described as follows:

 $\Pi: I \propto S$

where Π = approximation problem, I = set of inputs and S = set of solutions.

 $c {:} S \to \mathbb{R}^+$

Above, the cost function c maps each solution to a positive real number.

$$\forall i \in I, S(i) = \{s \in S : i \Pi_s\}$$

Above, the the set of feasible solutions, and our goal follows.

$$s^* \in S(i) \text{ and } c(s^*) = \min / \max \{c(S(i))\}$$

Here, we want to find the best solution s^* for a minimization/maximization problem. Specifically, we want to find it for the specific instance of that problem ($i\Pi s$).

Definition: Let Π be an optimization problem and let A_{Π} be an algorithm for Π that returns, $\forall i \in I, A_{\Pi} \in S_i$. We say that A_{Π} has an <u>approximation factor</u> of $\rho(n)$ if $\forall i$ in I such that |i| = n we have:

- minimization problem (basically, an explicit lower-bound of the optimal solution)

$$\frac{c(A_{\Pi}(i))}{c(s^*(i))} \le \rho(n)$$

- maximization problem (basically, an explicit upper-bound of the optimal solution)

$$\frac{c(s^*(i))}{c(A_{\Pi}(i))} \le \rho(n)$$

Here, we assume that *c* maps each feasible solution to a real number ≥ 1 .

Goal: $\rho(n) = 1 + \epsilon$, with ϵ as small as possible.

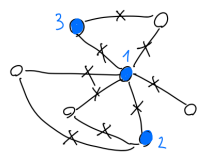
Definition: An <u>approximation scheme</u> for Π is an algorithm with 2 inputs $A_{\Pi}(i, \epsilon)$ that $\forall \epsilon$ is a $(1 + \epsilon)$ -approximation.

- In this case we just have to choose how much approximation we want by tuning the value of ϵ
- In other words: fixed an instance *i* of size *n*, the quality is ϵ (whatever ϵ is)

Very first algorithm you can think of? Use a greedy approach:

- select the vertex for the highest degree
- "remove" the touched edges
- repeat

Consider the following figure – take 3 as the highest, then 2 and 1 and remove touched edges as said:



Unfortunately, for this algorithm $\rho(n) = \Omega(\log(n))$.

How to prove a LB (Lower Bound)? It's enough to show one "bad" input instance.

Another algorithm (greedy approach):

- choose any edge
- add its endpoints to the solution
- "remove" the covered edges
- repeat

We'll show that this is a 2-approximation algorithm.

procedure Approx_Vertex_Cover(G)

 $V'=\emptyset$

E' = E

while $E' \neq \emptyset$: do

Let (u, v) be an arbitrary edge of E'

$$V' = V' \cup \{u, v\}$$

$$E' = E' \setminus \{(u, z), (v, w)\}$$

// remove edges that have u and v as endpoints

return V'

Complexity: O(n + m)

We'll show $\frac{|V'|}{|V^*|} \le 2$

Written by Gabriel R.

Given A = set of selected edges:

- A is a matching: $\forall e, e' \in A \Rightarrow e \cap e' = \emptyset$
 - a. i.e. set of edges with no vertices in common
 - b. every edge is disjoint, so there is no couple of edges sharing a common node
- *Approx_Vertex_Cover* selects a <u>maximal</u> matching: \forall edge $y, A \cup y$ is <u>not</u> a matching
 - a. this is a matching which cannot be increased
 - i. not possible to select an edge which touches other vertices

Proof:

1. lower bound to the optimal solution V^*

What can one say about $|V^*| vs |A|$?

A is a matching \Rightarrow in V^{*} there must be ≥ 1 vertex \forall edge of A (right figure)

In whatever vertex cover, particularly V^* , we have to cover all graph edges and, in particular, all A edges. But A is a matching (so, every edge of A is disjoint), so:

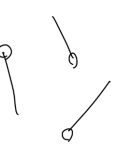
$$|V^*| \ge |A|$$

2. upper bound to the optimal solution V'

What can one say about |V'| vs |A|?

- |V'| = 2|A| by construction and so:

$$(1.+2.) \Rightarrow |V'| \le \frac{2|A|}{2} \le 2|V^*| \Rightarrow \frac{|V'|}{|V^*|} \le 2$$



22 TSP & METRIC TSP

22.1 TRAVELLING SALESPERSON PROBLEM (TSP)

Definition: Given a complete, undirected graph and a function $w: E \to \mathbb{R}^+$, output a <u>tour</u> $T \subseteq E$ (i.e. a cycle that passes through every vertex exactly once) minimizing $\sum_{e \in T} w(e)$. Collectively:

$$T \subseteq E \text{ minimizing } \sum_{e \in T} w(e)$$

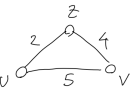
- $w: E \to \mathbb{R}^+$: we will work only on positive weights).
 - a. We can do this without loss of generality (wlog) because every TSP tour has the same number of edges ⇒ we can add a large weight to each edge, such that edges have nonnegative weights

22.2 METRIC TSP

<u>Metric TSP</u> is a special case of TSP where the weight function *w* satisfies the triangle inequality:

 $\forall u, v, z \in V$, it holds that $w(u, v) \leq w(u, z) + w(z, v)$

The following is an example of that:



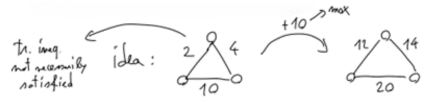
The problem can be shown to be NP-Hard, using an instance of TSP to build this one and using an Hamiltonian circuit to show we can assign a weight to each edge being "balanced" overall in the choice, always ensuring the best choice.

22.2.1 Metric TSP is NP-Hard

Theorem: Metric TSP is NP-Hard

Proof: $TSP \leq_p Metric TSP$

The idea is the following (where inequality is not strictly satisfied):



Given an instance of the TSP problem $\langle G = (V, E), w, k \rangle$, we build an instance of *Metric TSP* $\langle G' = (V, E), w', k' \rangle$ such that the triangle inequality is satisfied in G'. In order to to this, we can define the weight function w' as follows:

$$w'(u, v) = w(u, v) + W$$

giving
$$W = \max_{u, v \in V} \{w(u, v)\}$$

$$\langle G = (V_{1}E), w, k \rangle$$

$$\langle G' = (V,E), w', k' \rangle$$

$$W'(v,v) = W(v,v) + W$$

$$\lim_{v,v \in V} \{w(v,v)\}$$

Think of a value k' in such a way there if there exist an Hamiltonian circuit, there will be one in G' in such a way the cost of the tour will work for every edge, so:

$$k' = k + nW$$

To be shown yet:

- 1) w' satisfies triangle inequality
- 2) \exists an Hamiltonian circuit of cost k in $G \Leftrightarrow \exists$ Hamiltonian circuit of cost k' in G'

Let's see how to solve them formally:

1) $w'(u,v) \leq^{?} w'(u,w) + w'(w,v)$ (is it at most the weight of the others)? $w(u,v) + W \leq^{?} w(u,w) + w(w,v) + 2W$ (does this hold adding a general weight)? $w(u,v) \leq^{?} w(u,w) + w(w,v) + W$ (simply adding W both members) $w(u,w) + w(w,v) + W - w(u,v) \geq^{?} 0$ (is it true this is at most 0)? $y(u,w) + w(w,v) + W - w(u,v) \geq^{?} 0$ (is it true this is at most 0)?

We only ask if the definition of triangle inequality is satisfied correctly. Note it's important that the weights of edges are non-negative (otherwise, last part does not hold)

2)

- a. $\Rightarrow: \exists$ Ham. circuit of cost k in G. Note that an optimal solution contains exactly n edges and the same circuit in G' introduces a weight for every edge (so, $+W \forall$ edge). Thus, the cost of said tour in G' is k + nW
- b. just remove the $+W \forall$ edge to obtain a Ham. circuit of cost k in G

Let's see how to solve them formally:

3) $w'(u,v) \leq^{?} w'(u,w) + w'(w,v)$ (is it at most the weight of the others)? $w(u,v) + W \leq^{?} w(u,w) + w(w,v) + 2W$ (does this hold adding a general weight)? $w(u,v) \leq^{?} w(u,w) + w(w,v) + W$ (simply adding W both members) $w(u,w) + w(w,v) + W - w(u,v) \geq^{?} 0$ (is it true this is at most 0)? $v(u,w) + w(w,v) + W - w(u,v) \geq^{?} 0$ (is it true this is at most 0)?

We only ask if the definition of triangle inequality is satisfied correctly. Note it's important that the weights of edges are non-negative (otherwise, last part does not hold)

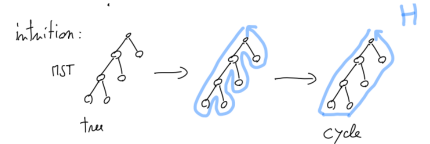
4)

- a. $\Rightarrow: \exists$ Ham. circuit of cost k in G. Note that an optimal solution contains exactly n edges and the same circuit in G' introduces a weight for every edge (so, $+W \forall$ edge). Thus, the cost of said tour in G' is k + nW
- b. just remove the $+W \forall$ edge to obtain a Ham. circuit of cost k in G

22.3 2-APPROXIMATION ALGORITHM FOR METRIC TSP

What is the most similar problem to *Metric TSP*? MST (Minimum Spanning Tree). We give the following *intuition*:

- we give an MST
- we want to build a cycle: what to do on a tree to achieve it?
- basically, there is a DFS traversing all the nodes
- the cycle forms having all nodes touched exactly once



We can simply solve this problem by adding the edge (e, a) to the *Preorder* list and make it an Hamiltonian circuit. We are free to add every edge we want because the graph is complete by definition. To do so, we define the following:

procedure Approx - Metric - TSP(G):

$$V = \{v_1, v_2, \dots v_n\}$$

$$r = v_1 //root from which Prim is run$$

$$T^* = Prim(G, r)$$

$$< v_{i_1}, v_{i_2}, \dots v_{i_n} > H' = PREORDER(T^*, r)$$

// lists all the vertices in the tree in an ordered fashion following a preorder walk

return $< H', v_{i_1} \ge H // basically, close the cycle$

This algorithm uses Prim as a subroutine to compute the MST. As such, this is super-fast and can be characterized as a near-linear algorithm.

So, to fully summarize:

- 1. Given a complete weighted graph G, pick any vertex v as the root, and find a minimum spanning tree T, using Prim's algorithm
- 2. Compile a list L of vertices encountered in a preorder traversal of T
- 3. Return *L* as a tour

22.4 3/2 (OR 1.5) APPROXIMATION ALGORITHM FOR METRIC TSP

<u>Christofides algorithm</u> (or Christofides–Serdyukov algorithm) was born in 1976.

Reason for 2-approximation factor was the fact the preorder traversal of T^* used every edge of T^* exactly <u>twice</u>. We'll try to improve on this by constructing a tour that traverses MST edges only <u>once</u>.

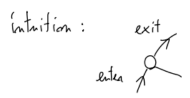
We give a couple of definitions useful for this context:

- A path (or cycle) is Eulerian if it crosses every edge of the graph exactly once
- A connected graph is Eulerian if there exists an Eulerian cycle

If the MST was Eulerian (cannot be) then we would have a 1-approx algorithm (which would be optimal, given one would cross every edge exactly once). *Approx_Metric_TSP* is finding a "cheap" Eulerian cycle in the MST, but effectively needs to double its edges.

Question: is there a cheaper Eulerian cycle?

Theorem: A connected graph is Eulerian \Leftrightarrow every vertex has even degree. The intuition is the following: enter a vertex, then exiting from it using a new edge, doing that without using edges more than once.



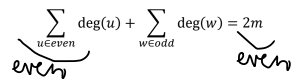
We want to focus on the *odd* degree vertices, given I have to cross again vertices (the even ones are fine, given we don't pass on them again). So, let's handle the <u>odd-degree</u> vertices of the MST explicitly.

Property: in any (finite) graph, the number of vertices of odd degree is even.

Proof: We use the following equality:

$$\sum_{v \in V} \deg(v) = 2m$$

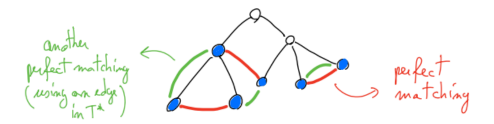
Basically, the sum of odd vertices with even ones, will get us an even result, that's the main intuition. So, we can split such summation into two parts:



Since the result must be even, the sum of degrees must be even too. But this happens only if the number of odd degree vertices is even.

Idea: augment the initial MST T^* with (the cheapest basically) a minimum-weight <u>perfect matching</u> (perfect means that it includes <u>all</u> the vertices) between the vertices that have odd degree in the MST.

For instance, let's consider the following MST, coloring in blue the odd-degree vertices. Imagine we add a perfect matching colored in red.



 \Rightarrow the resulting graph has only even-degree vertices, i.e. is an Eulerian graph.

Let's write the algorithm, which does exactly for things:

Christofides(G)

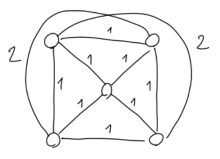
1) $T^* \leftarrow Prim(G,r)$ // $T^* = (V, E^*)$

2) Let *D* be the set of vertices of T^* with odd-degree. Compute a min-weight perfect matching M^* on the graph induced by *D* // this can be done in polynomial time (Edmonds, 1965)

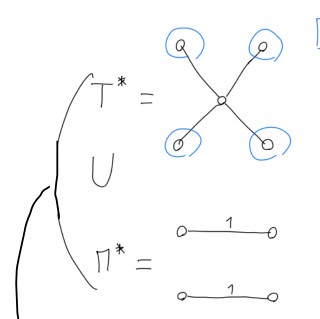
3) The graph $(V, E^* \cup M^*)$ is Eulerian // any edge in both E^* and M^* appears twice in this (multi)graph.

4) Return the cycle that visits all the vertices of G in the order of their first appearance in the Eulerian cycle (basically, skipping all repeated vertices – shortcutting)

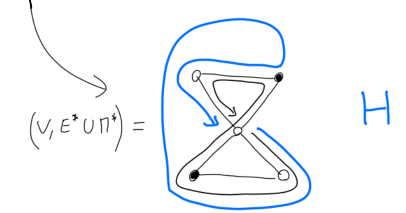
Consider the following example, connecting all vertices:



Now take the odd-degree vertices T^* and compute the minimum-weight perfect matching M^* .



Putting all of this together (merging it all) we get:



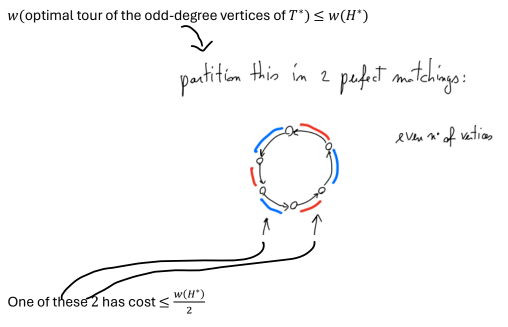
Analysing the algorithm:

- $w(H) \le w(T^*) + w(M^*)$ a. by triangle inequality
- $w(T^*) \le w(H^*)$

The goal to reach is $w(H) \leq \frac{3}{2}w(H^*)$. We would need to prove:

- $w(M^*) \leq \frac{1}{2}w(H^*)$ (by triangle inequality)

We will do the following clever step:



Putting all pieces together we get:

$$w(H) \le w(H^*) + \frac{w(H^*)}{2} = \frac{3}{2}w(H^*)$$

23 SET COVER

<u>Set cover</u> is an optimization problem that models many problems requiring resources to be allocated. It aims to find the least number of subsets that cover some universal set.

Its inputs are:

- I = (X, F) = instance of the set covering problem
- X = set of elements of any kind, called "universe"
- *F* ⊆ {*S*: *S* ⊆ *X*} = *B*(*X*)
 a. *B* stands for "Boolean": set of all subsets of *X*

There is a constraint that needs to be always respected: $\forall x \in X, \exists S \in F : x \in S$ i.e., "F covers X"

Optimization problem: (smallest subset of F having its members covering all X) \rightarrow find $F' \subseteq F$ s.t.

- 1) F' covers X
- 2) min |F'|

Example:

 $X = \{1, 2, 3, 4, 5\}$

 $F = \{\{1,2,3\},\{2,4\},\{3,4\},\{4,5\}\}$

 $\Rightarrow F^* = \{\{1,2,3\},\{4,5\}\}\$

23.1 SET COVER IS NP-HARD

Assertion: Set Cover (in its decision version $\langle (X, F), k \rangle$) is NP-hard.

Proof: Vertex Cover \leq_{p} *Set Cover*

- Given an instance of Vertex Cover Problem $\langle G = (V, E), k \rangle$
- we create an instance of Set Cover problem < (X, F), k >

Basically $\langle G = (V, E), k \rangle \rightarrow f \langle (X, F), k \rangle$

where:

- X = E
- $F = \{S_1, S_2, \dots S_n\}$ one \forall vertex $\in V$, 1,2, ... n
- $S_i = \{e = (u, v) \text{ such that } u = i \text{ or } v = i\}$, which is the set of covered edges by node e

Basically, there are |V| = n subsets S_i , and each subset is the set of edges incident to vertex *i*. Now show that finding a Set Cover of size $k \Leftrightarrow$ finding a Vertex Cover of size k.

- \Rightarrow Suppose $\{S_1, S_2, \dots, S_k\}$ is a set cover for X. Then, every edge in E must be incident to at least one vertex u_1, \dots, u_k . This happens because every element if one node of the adjacency list and so we find the minimal number of nodes touching all edges of graph, guaranteeing it will be minimal (for all sizes, given, even if less than k). Therefore, it forms a vertex cover of size k in G.
- \leftarrow Suppose $u_1, ..., u_k$ is a vertex cover in *G*. Then, S_i covers all the edges incident to vertex u_i . Therefore, $\{S_1, ..., S_k\}$ is a set cover of size *k* for *X*

23.2 GREEDY APPROXIMATION ALGORITHM

The greedy method works by picking at each stage, the set *S* that covers the greatest number of remaining elements that are uncovered:

- choose the subset that contains the largest number of uncovered elements
- remove from X those covered elements
- repeat until $X = \emptyset$

 $Approx_Set_Cover(X, F)$

U = X

 $F' = \emptyset // solution$

while $U \neq \emptyset$: do

// take the set of F covering as many elements as possible

 $let S \in F = |S \cap U| = \max_{S' \in F} \{|S' \cap U|\}$

 $U \leftarrow U \setminus S$ // update the list of available elements, removing those from S

 $F \leftarrow F \setminus \{S\}$ // remove the sets already considered inside of F

 $F' \leftarrow F' \cup \{S\}$

return F'

Correctness: (it does the job – it covers all the elements) At every iteration |U| decreases by at least one.

Complexity:

- n. of iterations $\leq |X|$ (every $S_i \in F$ contains at least an element)
- n. of iterations $\leq |F|$ (every $S_i \in F$ contains at least two elements)
- \Rightarrow n. of iterations \leq min {|X|, |F|}
- \forall iterations the complexity is $\leq |X| * |F|$ (scanning all elements and decreasing elements in both sets)
- $\Rightarrow O(|X| * |F| * min\{|X|, |F|\})$

24 RANDOMIZED ALGORITHMS

<u>Randomized algorithms</u> are algorithms that may do *random* choices, basically using a source of randomness in its logic. We give some basic examples:

- Example 1: Randomized quicksort (RQS)

Quicksort but chooses the pivot at random so to break the unlucky element choice and get on average a good probability on result.

- Example 2: Verifying polynomial identity

Checks if polynomials are equivalent and there are different approaches:

- check all the terms (slow)
- choose a random integer, compute the polynomials and check if they are equal
 - a. it may be wrong, outputting YES even If they are different

24.1 CLASSIFICATION OF RANDOMIZED ALGORITHMS

We divide these into two main categories:

- 1) randomized algorithms that <u>never fail</u>, which are called "<u>LAS VEGAS</u>" algorithms
 - a. (e.g., randomized quicksort)

$$\forall i \in I, A_R(i) = s \quad s.t.(i,s) \in \Pi$$

where $\Pi \subseteq I \times S$ is the decisional problem, *i* is an input instance, A_R is the random algorithm which applied to the input instance produces a solution *s* s.t. the couple (*i*, *s*) belongs to Π

Randomness comes into play in the analysis of the complexity – because it depends from the randomness of the choices. $\forall n, T(n)$ is a <u>random variable</u> of which we usually study its expectation E[T(n)] or $\Pr(T(n) > c * f(n)) \rightarrow \leq \frac{1}{n^k}$ (so, for some constants c and k, we say that T(n) = O(f(n)) with high probability (here, T(n) is called complexity function) – this second one is more powerful than the first, so Pr more powerful than E.

- 2) randomized algorithms that may fail are called "MONTE CARLO" algorithms
 - a. e.g., verifying polynomial identities

$$\forall i \in I, A_R(i) = s \quad s.t.(i,s) \notin \Pi$$

We study $Pr((i, s) \notin \Pi)$ as a function of $n = |i| \rightarrow family of random variables (binary)$

Moreover, even T(n) may be a random variable. For decision problems, these algorithms can be divided into:

- <u>one-sided</u>: they may fail only on one answer
 - a. e.g., can make right all YES instances but may be wrong on all NO instances
 - two-sided: they may fail in both answers
 - a. e.g., it can make wrong all YES instances but can make wrong all NO instances

24.2 KARGER'S ALGORITHM FOR MINIMUM CUT

A quite simple MONTE CARLO and elegant algorithm created in 1993. Let's start from the problem itself it wants to solve: the <u>minimum cut</u> revolves finding a cut of minimum size, that is, the minimum number of edges whose removal disconnects the graph. A couple of useful definitions to see the problem:

Definition: A <u>multiset</u> is a collection of objects with repetitions allowed. It's usually denoted between a couple of brackets, as you can see here.

 $S = \{\{objects\}\}\$ $\forall object \ o \in S, m(o) \in \mathbb{N} \setminus \{0\}$

where m = multiplicity, so how many copies of "o" are in S.

Definition:

- a multigraph $G = (V, \mathcal{E})$ s.t. $\forall V \subseteq \mathbb{N}, V$ finite and \mathcal{E} is a multiset of elements (u, v) s.t. $u \neq v$
- Note: A simple graph G = (V, E) is also a multigraph.

Definition:

- given $G = (V, \mathcal{E})$ connected, a <u>cut</u> $C \subseteq \mathcal{E}$ is a multiset of edges s.t. $G' = (V, \mathcal{E} \setminus C)$ is not connected.

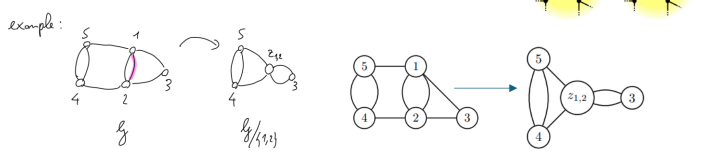
Let's give Karger's idea here:

- choose an edge at random
- "contract" the two vertices of that edge, removing all the edges incident both vertices

This works with very low probability, but let's use the trick we saw already: repeating this a good enough number of times, can actually refine the analysis and obtain a good level of probability.

We see below two examples of the same contraction in Karger and on the right a generic contraction.

- Basically, it makes the two vertices to collapse in just one vertex connected with all the previous adjacent vertices
- If as a result there are several edges between some pairs of (newly formed) vertices, retain them all.



12

Definition: given $G = (V, \mathcal{E})$ and $e = (u, v) \in \mathcal{E}$, the contraction of G with respect to e, $\frac{G}{e} = (V', \mathcal{E}')$ is the multigraph with $V' = V \setminus \{u, v\} \cup \{z_{u,v}\}$ with $z_{u,v} \notin V$ coming from the fusion of u and v:

$$\mathcal{E}' = \mathcal{E} \setminus \{\{(x, y) \text{ s. } t. (x = u) \text{ or } (x = v)\}\}$$
$$\cup \{\{(z_{u,v}, y) \text{ s. } t. (u, y) \in \mathcal{E} \text{ or } (u, y) \in \mathcal{E}, y \neq u \text{ and } y \neq u \text{ or } (y \neq u)\}$$

We describe the algorithm here:

 $\begin{array}{ll} \textit{FULL_CONTRACTION} \ (G = (V, \mathcal{E})) \\ & \text{for } i = 1 \text{ to } n - 2 \text{: do} \\ & e \leftarrow \textit{RANDOM}(\mathcal{E}) & // \textit{choose an edge at random in multiset} \\ & G' = (V', \mathcal{E}') \leftarrow \frac{G}{e} & // \textit{contraction on graph with random edge selection} \\ & V \leftarrow V' & // \textit{construct the new graph with vertices and edges} \\ & \mathcal{E} \leftarrow \mathcal{E}' \\ & \text{return} \ |\mathcal{E}| & // \textit{return the graph and its cardinality} \end{array}$

Consider $k \rightarrow$ how many times to repeat *FULL_CONTRACTION*, which depends on the probability of making a mistake – hence, it depends on the analysis of the algorithm)

 $KARGER(G = (V, \mathcal{E}), k)$ $min - cut = \mathcal{E}$ for i = 1 to k: do $t = FULL_CONTRACTION(G)$ if |t| < |min - cut| then min - cut = t repeats FULL - CONTRACTION(k times to reduce the probability of errorto be determined by the analysis

return *min – cut*

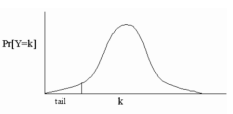
The analysis suggests us the following:

• in order to obtain $\Pr(\text{Karger succeds}) > 1 - \frac{1}{n^d}$ we need to repeat "Full contraction" $k = \frac{dn^2 \ln n}{2}$ times

25 CHERNOFF BOUNDS

<u>Chernoff bounds</u> are tools from modern probability theory that are frequently used in the analysis of randomized algorithms. They're a more powerful version of the Markov's lemma. This mainly uses *indicator* random variables (that is, variables which can have either value 0 or 1).

Generally, Chernoff bounds are a tool which allow to study the concentration of an event around its mean (specifically, in the "tail" – see figure – so, far from the mean) and to overcome the previous fact we use them.



- The markup is a little loose, not very significant
- Better augmentation allows me to move from analysis to the average case to the more desirable high probability analysis

The idea between Chernoff bounds is to transform the original random variable into a new one, such that the distance between the mean and the bound we will get is significantly stretched. It answers the question about how tight the bound we can get when having more information about the distribution of the random variables.

We give Chernoff's lemma here: let $X_1, X_2, ..., X_n$ independent indicator random variables where $E[X_i] = p_i, 0 < p_i < 1$. Let $X = \sum_{i=1}^n X_i$ and $\mu = E[X]$. Then $\forall \delta > 0$:

$$\Pr(X > (1+\delta)\mu) < \left(\frac{e^{\delta}}{(1+\delta)^{(1+\delta)}}\right)^{\mu}$$

In words: the outcome concentrates around the min is very high – to the contrary, the probability of deviating from the min should be very low.

25.1 CHERNOFF BOUND VARIANTS

Consider the following variants of Chernoff bounds (weaker but easier to state and to use):

1)
$$\Pr(X < (1 - \delta)\mu) < e^{-\frac{\mu\delta^2}{2}}, 0 < \delta \le 1$$

2) $\Pr(X > (1 + \delta)\mu) < e^{-\frac{\mu\delta^2}{2}}, 0 < \delta \le 2e - 1$

25.2 ANALYSIS IN HIGH PROBABILITY OF RANDOMIZED QUICKSORT

As an example of application of Chernoff bounds, we do the analysis of Randomized Quicksort, in which we remember the pivot is chosen at random (possibly, not very far from the median). This is a LAS VEGAS algorithm, since it always sorts.

RandQuickSort(S)

|S| = n, all distinct

if $|S| \leq 1$ then return $\langle S \rangle$

p = RANDOM(S) // pick a "pivot" element uniformly at random from S

 $S_{1} = \{x \in S \text{ s.t. } x < p\} // O(n) \text{ time}$ $S_{2} = \{x \in S \text{ s.t. } x > p\} // O(n) \text{ time}$ $Z_{1} = RandQuickSort(S_{1})$ $Z_{2} = RandQuickSort(S_{2})$ return $\langle Z_{1}, p, Z_{2} \rangle$

We want to approximate as close as possible to the actual median. The generic event E can be characterized as the "good choice" of the pivot between all the statistically possible choices.

So, calculating the cost of the algorithm:

- total work at each level is mostly linear, so $\leq c * n$
- depth of the recursion tree = min $\left\{ integer \ i \ s. \ t. \left(\frac{3}{4}\right)^i n \le 1 \right\} = \left[\log_{\frac{4}{3}}(n) \right] = O(\log(n))$

So, continuing:
$$\left(\frac{3}{4}\right)^i n \le 1 \Leftrightarrow \left(\frac{3}{4}\right)^i \le \frac{1}{n} \Leftrightarrow \left(\frac{4}{3}\right)^i \ge n \Leftrightarrow \log_{\frac{4}{3}}\left(\frac{4}{3}\right)^i \ge i \ge \log_{\frac{4}{3}}(n)$$

$$\Rightarrow T_{RQS}(n) = O(nlog(n))$$

Fix <u>one</u> root-leaf path *P* and the following lemma says, "with h.p. the path chosen in short"; specifically shorter than $\log (n)$ ".

Lemma: $\Pr(|P| > a * \log_{\frac{4}{3}}(n)) < \frac{1}{n^3}$

We are trying to find the probability there is at least one big path around the mid value.

If this is true, we're done, applying the very frequent/very famous following lemma.

Lemma (Union bound): for any random events $E_1, ..., E_K$:

$$\Pr(E_1 \cup E_2 \cup \dots \cup E_k) \le \Pr(E_1) + \Pr(E_2) + \dots + \Pr(E_k)$$

If the lemma is true, it follows that:

- Given the event E_i = the path p_i has length > $a * \log_{\frac{1}{2}}(n)$:

$$\Pr\left(\exists \ path > a * \log_{\frac{4}{3}}(n)\right) = \Pr\left(\bigcup_{i=1}^{n} E_{i}\right) \leq^{union \ bound}$$
$$\leq \sum_{i=1}^{n} \Pr(E_{i}) <_{lemma} n * \frac{1}{n^{3}} = \frac{1}{n^{2}}$$
$$\dots \geq 1 - \frac{1}{n^{2}}$$

 $\Rightarrow T_{RQS}(n) = O(nlog(n))$ w.h.p.

Suppose *p* is always the <u>median</u> of *S*; then

$$T_{RQS}(n) = \begin{cases} 2T_{RQS}\left(\frac{n}{2}\right) + O(n), & n > 1\\ 0, & n \le 1 \end{cases}$$

So, basically:

- there are *n* nodes (excluding leaves associated to \emptyset) $\Rightarrow \le n$ paths root-leaf
- we will show these paths are not so long; using any method (Master theorem or whatever), we get the same height of the tree

$$T_{ROS}(n) = ^{Master Theorem} O(n \log(n))$$

However, p is the median with probability $\frac{1}{n}$, which is very low.

(The following part is useful for the exam)

The event *E* can be characterized as "in the first $l = a * \log_{\frac{4}{3}}(n)$ nodes of *P* there have been $< \log_{\frac{4}{3}}(n)$ lucky choices". I'm studying this last event:

- $X_i, 1 \le i \le l = a * \log_{\frac{4}{2}}(n)$
- $X_i = 1$ if at the i^{th} vertex of P there is a lucky choice of the pivot

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$$\Pr(X_i = 1) = \frac{1}{2} \forall i$$

- X_i are independent

We want the probability of $P(\sum_{i=1}^{l} X_i) < \log_{\frac{4}{2}}(n)$ to be bound (and to be very low).

Given $X = \sum_{i=1}^{l} X_i$, its expected value is as follows:

$$\mu = E[X] = E[\sum_{i=1}^{l} X_i] = \sum_{i=1}^{l} E[X_i] = \sum_{i=1}^{l} \frac{1}{2} = \frac{l}{2} = \frac{a}{2} \log_{\frac{4}{3}}(n)$$

Now, let's apply the following Chernoff bound:

$$\Pr(X < (1 - \delta)\mu) < e^{\frac{-\mu\delta^2}{2}}, 0 < \delta \le 1$$

$$\downarrow$$

$$(1 - \delta)\mu = \log_{\frac{4}{3}}(n)$$

$$(1 - \delta)\frac{a}{2}\log_{\frac{4}{3}}(n) = \log_{\frac{4}{3}}(n)$$

One possible choice is $a = 8, \delta = \frac{3}{4}$.

$$\Pr(X < \log_{\frac{4}{3}}(n)) < e^{-\frac{8}{4} * \log_{\frac{4}{3}}(n) * \frac{9}{16}}$$
$$= e^{-\frac{8}{4} * \log_{\frac{4}{3}}(n) * \frac{9}{8}}$$
$$< e^{-\log_{\frac{4}{3}}(n)}$$

Written by Gabriel R.

$$= e^{\frac{-\ln(n)}{\ln\left(\frac{4}{3}\right)}}$$
$$= \left(e^{-\ln(n)}\right)^{\frac{1}{\ln\left(\frac{4}{3}\right)}}$$
$$= \left(\frac{1}{n}\right)^{\frac{1}{\ln\left(\frac{4}{3}\right)} \approx 3,47}}$$
$$< \frac{1}{n^3}$$