

Design and Analysis of Algorithms

José Dorronsoro
Escuela Politécnica Superior
Universidad Autónoma de Madrid

Contents

1	Elementary Graph Algorithms	3
1.1	Basic Concepts on Graphs	3
1.2	Minimum Distances on Graphs	5
1.3	All Pairs Shortest Paths	10
2	Minimum Spanning Trees	13
2.1	The Algorithms of Prim and Kruskal	13
2.2	The Disjoint Set Abstract Data Type	16
2.3	Correctness of Prim and Kruskal	23
3	Eulerian and Hamiltonian Circuits	27
3.1	Eulerian Circuits	27
3.2	Hamiltonian Circuits and an Excursion on Complexity Theory	33
3.3	The Traveling Salesman Problem	36
4	An Excursion on DNA Sequencing	40
4.1	Hamilton, Euler and DNA Sequencing	40
4.2	From Reads to the Genome	45
5	Depth First Search and Connectivity	46
5.1	Depth First Search	46
5.2	Biconnected Graphs	49
5.3	DAGs and Topological Sort	54

Before We Start

On reading and studying these notes:

From Brad DeLong's, UC Berkeley, [A note on reading big, difficult books](#):

- It is certainly true that there are many who can parrot verbal formulas yet lack knowledge of facts, terms, and concepts.
- It is certainly true that there are many who have knowledge of facts, terms, and concepts and yet lack deep understanding.
- But **I am not aware of anyone who has deep understanding of a discipline and yet lacks knowledge of facts, terms, and concepts.**
- And **those who know the facts, terms, and concepts cold are the absolute best at parrotting verbal formulas.**

1 Elementary Graph Algorithms

1.1 Basic Concepts on Graphs

Definitions

- Graph: Pair $G = (V, E)$ of a set V of vertices (nodes) and a set E of edges (u, v) with $u, v \in V$
- Edges imply direction: in (u, v) we go from u to v
- In general, graphs are **directed**
- **Undirected** graphs: $(u, v) \in E$ iff $(v, u) \in E$
- **Unweighted** graphs: we only consider edge structure
- **Weighted** graphs: edges (u, v) have weights w_{uv}
- **Multigraphs**: there might several edges between two vertices and also between a vertex and itself

Storing an Unweighted Graph

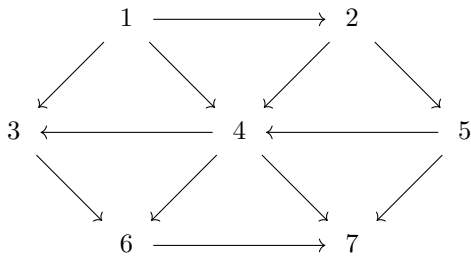
- **Adjacency matrix**: Assume $V = \{1, \dots, N\}$. Then if $(i, j) \in E$, $m_{ij} = 1$; else, $m_{ij} = 0$
 - Not for multigraphs
 - By convention $m_{ii} = 1$ (although sometimes we may consider $m_{ii} = 0$)
 - Cost: $\Theta(|V|^2) = \Theta(N^2)$
- **Adjacency list**: We can consider a pointer table $T[\]$ where $T[i]$ points to a linked list
 - If $(i, j) \in E$, then j is in one of nodes pointed by $T[i]$
 - Cost: $\Theta(|V|) + \Theta(|E|)$
 - No problem for multigraphs

- For standard graphs the cost is always $O(|V|^2)$ for both methods, since we then have

$$|E| \leq |V|(|V| - 1) = O(|V|^2)$$

An Example

- A directed graph:



The Adjacency Matrix

- The first rows of the adjacency matrix are

$$M = \begin{pmatrix} 1 & 1 & 1 & 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 1 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 1 & 0 \\ 0 & 0 & 1 & 1 & 0 & 1 & 1 \\ & & & \dots & & & \end{pmatrix}$$

The Adjacency List

- Partial adjacency list: we use a lexicographic order

$$\begin{array}{l} 1 \rightarrow 2 \rightarrow 3 \rightarrow 4 \\ 2 \rightarrow 4 \rightarrow 5 \\ 3 \rightarrow 6 \\ 4 \rightarrow 3 \rightarrow 6 \rightarrow 7 \\ \dots \end{array}$$

The Size of a Graph

- While $|V|$ and $|E|$ are in general independent, we may expect $|V| = O(|E|)$ for interesting graphs
 - $|E|$ will usually give G 's size
- G is **dense** if $|E| = \Theta(|V|^2)$
- G is **sparse** if $|E| \ll |V|^2$
- If G is dense, the adjacency matrix storage is more efficient; if G is sparse, adjacency lists are better
- We will usually work with adjacency lists, using adjacency matrices for special algorithms

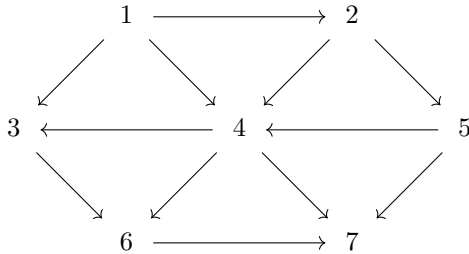
1.2 Minimum Distances on Graphs

Minimum Distance Problems

- **Path** from u to v : a subset $\pi = \{u = u_0, \dots, u_K = v\} \subset V$ with $(u_i, u_{i+1}) \in E$
- **Length** of π : $|\pi| = K = \#(\text{number of})$ edges
- First problem: given u , find a **shortest path** (i.e., a path with the smallest number of edges) π from u to any other v
- First question: how to obtain such paths?
- First idea: get a tree like “descending representation” of G starting from u and avoiding lower duplicate vertices

Minimum Distance Example

- Think of each vertex as a ball and of edges as equal length strings, and make G “hang” from u discarding “repeated” edges
- On the previous graph,

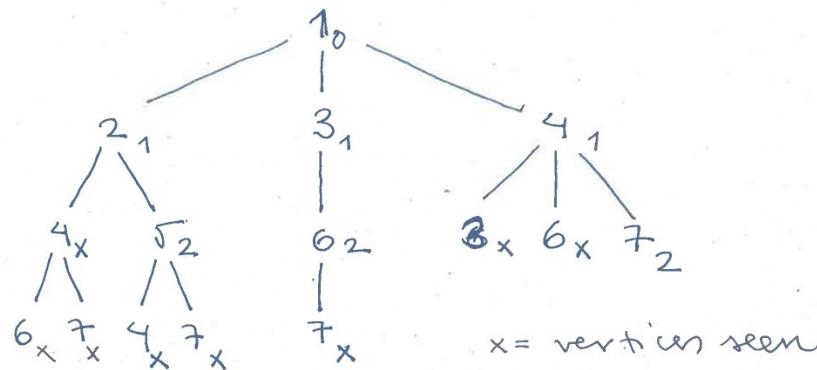


Breadth First Traversal

- We find the minimum distances by breadth first traversal (BFS) on this hanging representations

Some Observations on Minimum Distance Problems

- If $d[v]$ is the depth of v in T , it is reasonable to expect $d[v]$ to be the minimum distance from u to v
 - But we have to prove it
- If $p[v]$ is the father of v in T , we can obtain a minimum length path from u to v with edges $(w = p[v], v), (p[w], w), \dots$, and so on
- Notice that this way we have found the minimum distances from u to **all** $v \in V$
 - They are unique, but the minimum paths are not
- Q: how can we derive an algorithm for this?
- We can use a standard FIFO queue Q to process the different vertices and the tables $p[v]$ and $d[v]$



- In fact, this fits in the general framework of **Breadth First Search**

First Algorithm for Minimum Distances

- We need tables $p[v]$ for the vertex “previous” to v , $d[v]$ for the minimum distance from u to v and $v[v]$ to mark v as seen
- First, queue-based, pseudocode:

```

def dist_min(u, G):
    s[ ] = F; p[ ] = None; d[ ] = inf
    Q = q()
    d[u] = 0; Q.put(u); s[u] = T
    while not Q.empty():
        v = Q.get()
        for all z adjacent to v:
            if not s[z]: #first time z is seen
                d[z] = d[v] + c(v, z)
                p[z] = v; s[z] = T
                Q.put(z)
    return d, p
  
```

Some Observations on `dist_min`

- The table $s[]$ is redundant: $s[v] == T$ if and only if $d[v] < \infty$ (exercise: update the psc)
- We can use $p[]$ to reconstruct the minimum paths from u to all v (exercise)
- We can use $p[]$ to reconstruct the minimum distance table $d[]$ (exercise)
 - So $p[]$ would be the table to return in, say, a C function
- A vertex enters Q only once \Rightarrow the linked lists are traversed only once \Rightarrow the cost of `distMin` is $O(|E|)$, i.e., linear on G 's size

- `dist_min` is a particular instance of the general **Breadth First Search** algorithm

Breadth First Search (BFS) v 1.0

- The pseudocode of the first, queue-based version of BFS is

```
def BFS(u, G):
    s[ ] = F; p[ ] = None
    Q = q()
    doSomething(u)
    s[u] = T; Q.put(u)
    while not Q.empty():
        v = Q.get()
        for all z adjacent to v:
            if not s[z]:
                doSomething(z)
                s[z] = T; p[z] = v
                Q.put(z)
    return p
```

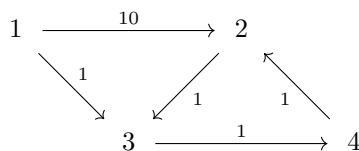
- Since we enter each list only once, if the cost of `doSomething` is $O(1)$, the cost of BFS is $O(|E|)$, i.e., linear,
- If needed, we add a driver to restart `BFS` at unseen nodes

Minimum Distances on Weighted Graphs

- $G = (V, E)$ is a **weighted** graph if there is a function $c : E \rightarrow \mathbb{R}$
 - We think of $c(i, j)$ as the cost of going from i to j
 - Although sometimes $c(i, j)$ can be negative
- **Cost** of path π : $c(\pi) = c(\{u_0, \dots, u_K\}) = \sum_1^K c(u_{j-1}, u_j)$
- Working with adjacency matrices we can store c as $m_{ij} = c_{ij}$ if $(i, j) \in E$ and $m_{ij} = \infty$ if not.
 - Now the convention is $m_{ii} = 0$
- Working with adjacency lists we can store c_{ij} in a second field of the same node of $T[i]$ that stores j

Problems ...

- Applying our first algorithm to the graph



- Working here with the tree like representation of G is now trickier
which is obviously wrong

	d	p	v	d	p	v	d	p	v
1	0	-	T	0	-	T	0	-	T
2	∞	-	F	10	1	T	10	1	T
3	∞	-	F	1	1	T	1	1	T
4	∞	-	F	∞	-	-	2	3	T
Q	1			2, 3			3, 4		

Fixing The First Algorithm

- The node 2 gets out of Q too soon \Rightarrow we have to change the ordering in Q
- We use a **priority queue** Q that orders vertices using the current value of $d[v]$
- Now v is seen when it **leaves** Q (and not when it enters Q)
- We also need (again) a table $s[v]$ to check whether v has left Q and, hence, we do not consider it any longer
- This leads to **Dijkstra's** algorithm for positive costs

Dijkstra's Algorithm

- Dijkstra's pseudocode is:

```
def dijkstra(u, G):
    s[ ] = F; p[ ] = None; d[ ] = inf          # 1
    Q = pq()
    d[u] = 0
    Q.put( (d[u], u) )
    while not Q.empty():                      # 2
        v = Q.get()                           # 3
        if not s[v]:
            s[v] = T
            for all z adjacent to v:          # 4
                if d[z] > d[v] + c(v, z):
                    d[z] = d[v] + c(v, z)
                    p[z] = v
                    Q.put( (d[z], z) )        # 5
    return d, p
```

Dijkstra's Algorithm II

- **Example:** First steps of Dijkstra's algorithm on the previous graph

	d	p	v	d	p	v	d	p	v
1	0	-	F	0	-	T	0	-	T
2	∞	-	F	10	1	F	10	1	F
3	∞	-	F	1	1	F	1	1	T
4	∞	-	F	∞	-	-	2	3	F
PQ	1 ₀			3 ₁ , 2 ₁₀			4 ₂ , 2 ₁₀		

Dijkstra's Cost

- The five commented numbers in the psc determine its cost

- The cost of (1) is clearly $O(|V|)$
- Using a PQ over a binary heap the cost of `Q.put`, `Q.get` is $O(\log |Q|)$
 - Q will contain at most an item for every edge, so $|Q| = O(|E|)$
 - Thus, the cost of (3) over all iterations in (2) is $O(|E| \log |E|)$
- We enter (4) **once** per node; thus the total number of joint iterations in (2) and (4) is $|E|$
- Hence, the cost of (5) over all iterations is $O(|E| \log |E|)$
- Since usually $|E| = O(|V|^2)$, the overall cost is

$$\begin{aligned} O(|V|) + O(|E| \log |E|) &= O(|V|) + O(|E| \log |V|^2) \\ &= O(|V|) + O(|E| \log |V|) \end{aligned}$$

- This will be $O(|E| \log |V|)$ for most graphs, i.e., log linear in a graph's size

Observations on Dijkstra's Algorithm

- We allow that several instances of the same v be in Q
- We can stop the algorithm earlier using a counter of seen vertices (exercise)
 - But have to clear Q , so ...
- Dijkstra **works** for positive weights: at its end
 - $d[v]$ contains the minimum distance from u to any other v
 - And we can get the minimum paths using $p[v]$
- But **all this has to be proved**
- Dijkstra is an example of the general **breadth first search** graph algorithm
 - And also of a **greedy** algorithm

Breadth First Search (BFS) v 2.0

- The pseudocode for general, PQ based BFS, is

```
def BFS(u, G):
    s[ ] = F; p[ ] = None; Q = pq()
    d[u] = 0; Q.put( (d[u], u) )
    doSomething(u)
    while not Q.empty():
        _, v = Q.get()
        if not s[v]:
            s[v] = T
            for all z adjacent to v:
                doSomething(z)          #perhaps change d[z]
                p[z] = v; Q.put( (d[z], z) )
    return p
```

- If needed, we add a driver to restart BFS at unseen nodes
- If the cost of `doSomething` is $O(1)$ and we work with a PQ over min heaps, the cost of BFS is $O(|E| \log |V|)$

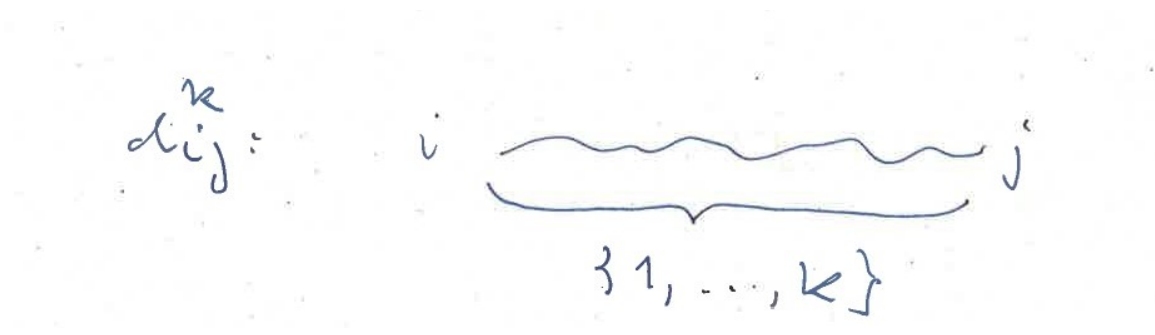
1.3 All Pairs Shortest Paths

All Pairs Shortest Paths

- If (G, c) is a weighted directed graph, we can consider in principle three minimum distance problems:
 - For u, v fixed, find **only** the minimum distance between u and v
 - For u fixed, find the minimum distance between u and **all other** $v \in V$
 - For **all** $u, v \in V$, find the minimum distance between u and v
- While the first problem seems easier, no algorithm for general graphs is better than the best one for the second problem
 - Notice that a minimal path from u to v is also minimal for all vertices in between
- We can solve the third problem iterating an algorithm for the second one over all $u \in V$
 - For instance, iterating Dijkstra over all $u \in V$ has a cost $|V| \times O(|E| \log |V|) = O(|V||E| \log |V|)$
 - If G is dense, the cost is then $O(|V|^3 \log |V|)$

Improving on Dijkstra I

- Assume $V = \{1, \dots, N\}$ and the cost c is nonnegative
- Denote by d_{ij} the minimum distance between i and j
- We define d_{ij}^k be the minimum distance between i, j but where **the intermediate nodes are taken only from $\{1, \dots, k\}$**

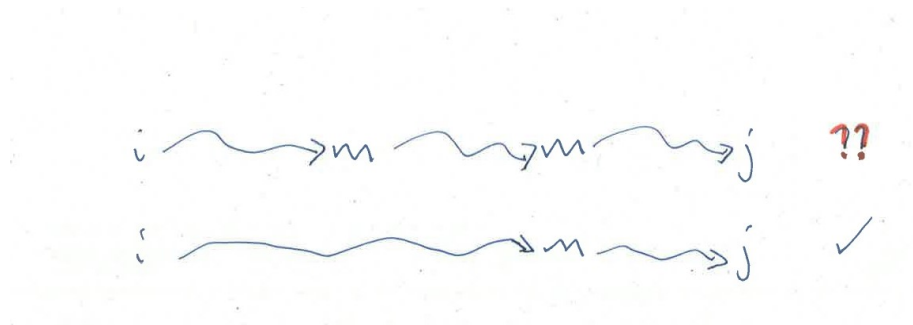


- It is clear that

$$d_{ij}^0 = c(i, j), \quad d_{ij}^N = d_{ij}$$

- It is clear that **no vertex is repeated on the optimal path** that gives d_{ij}^k

Improving on Dijkstra II



- Obviously, an optimal path between i and j with $\{1, \dots, k\}$ as intermediate nodes may or may not contain k
- If it doesn't, we have

$$d_{ij}^k = d_{ij}^{k-1}$$

- If it does, we have

$$d_{ij}^k = d_{ik}^{k-1} + d_{kj}^{k-1}$$

for we have:

- A path from i to j is **optimal iff the partial subpaths** between i and k and k and j **are optimal**, i.e.,

$$d_{ij}^k = d_{ik}^k + d_{kj}^k$$

- But a path **having another** k between i and k or between k and j **cannot be optimal**:

* We can simply remove the subpath from k to k to get a better path

- Thus, it is then obvious that

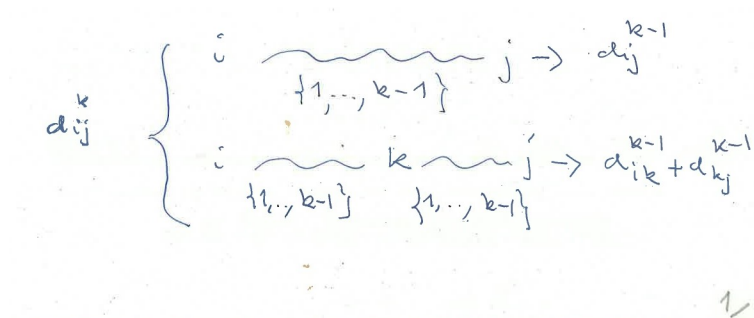
$$d_{ik}^k = d_{ik}^{k-1}, d_{kj}^k = d_{kj}^{k-1}$$

Dynamic Programming Solution

- We can conclude

$$d_{ij}^k = \min\{d_{ij}^{k-1}, d_{ik}^{k-1} + d_{kj}^{k-1}\}$$

and $d_{ij} = d_{ij}^N$



Floyd–Warshall Algorithm

- Working with adjacency matrices, this suggests the following (quite bad) pseudocode

```
def FW_0(m_c):
    n = m_c.shape[0]
    d = np.empty((n, n, n+1))
    #d[i, j, k+1]: d_min from i to j with intermediate [0, ..., k-1]
    d[:, :, 0] = m_c
    for k in range(n):
        for i in range(n):
            for j in range(n):
                t = d[i, k, k] + d[k, j, k]
                d[i, j, k+1] = min(d[i, j, k], t)
    return d[:, :, n]
```

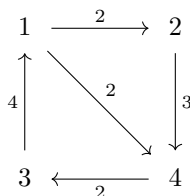
- The algorithm is \pm obviously correct
 - In fact, it also works for negative c provided there are **no negative cycles**

Floyd–Warshall Cost

- The time cost is $O(N^3)$, better than iterated Dijkstra for dense graphs
- The space cost is at first sight also $O(N^3)$ as we use N matrices $N \times N$; but in fact a single matrix D is enough, for
 - We first “retain” d_{ik}, d_{kj}
 - Then for i or $j \neq k$ we set $c = d_{ik} + d_{kj}$, and we can overwrite d_{ij} as $d_{ij} = \min\{d_{ij}, c\}$
- Exercise (easy): rewrite FW taking advantage of this
 - Is it now a good **Python** algorithm?
- Exercise (more difficult): how can we recover the optimal paths?
- Observation: FW is our first example of a problem solvable by a **Dynamic Programming (DP)** algorithm, which exploits
 - An optimization problem with an **optimal substructure** (obvious: any optimization problem has it) that we are able to make **explicit**
 - The explicit substructure formula also ensures FW to be correct

Applying Floyd–Warshall

- Example:



- We iteratively compute the intermediate matrices

$$D^k = (d_{ij}^k), \quad k = 0, 1, \dots, N$$

- Observe that going from D^{k-1} to D^k **we just copy** $d_{ik}^k = d_{ik}^{k-1}, d_{kj}^k = d_{kj}^{k-1}$

From D^0 to D^1

- We have

$$\begin{aligned} d_{23}^1 &= \min\{d_{23}^0, d_{21}^0 + d_{13}^0\} = \min\{\infty, \infty + \dots\} = \infty \\ d_{24}^1 &= \min\{d_{24}^0, d_{21}^0 + d_{14}^0\} = \min\{3, \infty + \dots\} = 3 \end{aligned}$$

and so on, to get

$$D^0 = \begin{pmatrix} 0 & 2 & \infty & 2 \\ \infty & 0 & \infty & 3 \\ 4 & \infty & 0 & \infty \\ \infty & \infty & 2 & 0 \end{pmatrix} \rightarrow D^1 = \begin{pmatrix} \mathbf{0} & \mathbf{2} & \infty & \mathbf{2} \\ \infty & \mathbf{0} & \infty & 3 \\ \mathbf{4} & 6 & \mathbf{0} & 6 \\ \infty & \infty & 2 & \mathbf{0} \end{pmatrix}$$

- And similarly we get D^2, D^3 and D^4

2 Minimum Spanning Trees

2.1 The Algorithms of Prim and Kruskal

Trees

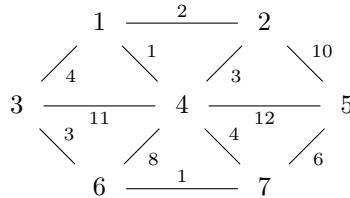
- An undirected graph $G = (V, E)$ is **connected** if for every pair $u, v \in V$ there is a path π in G from u to v
- A **cycle** π in a graph $G = (V, E)$ is a path that starts and ends at the same point
- A **tree** is an undirected connected graph that is also **acyclic**, i.e., there are no cycles in E
- A tree T is a **spanning tree** (ST) for $G = (V, E)$ if $T = (V, E_T)$ with $E_T \subset E$
- If G is weighted, the **cost** of an ST T is

$$c(T) = \sum_{(u,v) \in E_T} c(u, v)$$

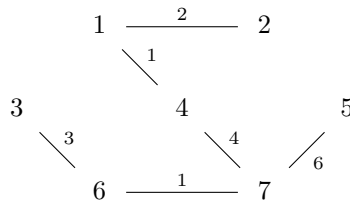
- $T = (V, E_T)$ is a **minimum spanning tree** (MST) for the undirected graph $G = (V, E)$ if for any other ST $T' = (V, E'_T)$ we have $c(T) \leq c(T')$

MST Examples

- On the graph



a first MST with cost 17 is



Prim's Algorithm

- Changing slightly Dijkstra's gives **Prim's** algorithm for finding MSTs

```
def prim(G, u):
    s[ ] = F; p[ ] = NULL; c_t[ ] = inf          # 1
    Q = pq()
    c_t[u] = 0; Q.put( (c_t[u], u) )
    while not Q.empty():                        # 2
        _, v = Q.get()                          # 3
        if not s[v]:
            s[v] = T
            for all z adjacent to v:           # 4
                if not s[z] and c_t[z] > c(v, z):
                    c_t[z] = c(v, z)
                    p[z] = v
                    Q.put( (c_t[z], z) )        # 5
    return p, c_t
```

- The second `if not s[z]` didn't appear in Dijkstra; do we need it here?

Observations on MSTs

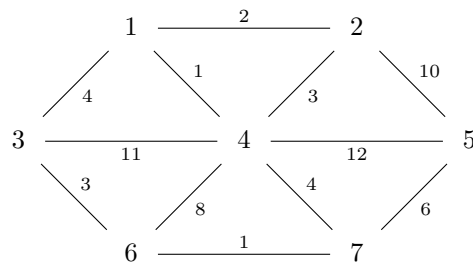
- There may be several minimum spanning trees in a graph but the **minimum cost is unique**
- We recover the MST with the table $p[]$ and have $c(T) = \sum_{v \neq u} c(p[v], v)$
- The cost of Prim is $O(|E| \log |V|)$ if the PQ is built over a min heap
- **Prim works:** at the end, the edges $(p[v], v)$ of a MST E_T are given by $p[v]$ and the $c[v]$ give their costs
 - But again this has to be proved
 - And, since it is correct, we do not need to check $s[v] == T$ for if z already seen, $c_t[z] \leq c(v, z)$ (although it saves time)
- Prim and Dijkstra are examples of a **greedy** algorithms

Greedy Algorithms

- A greedy algorithm tries to solve a **global optimization problem** by making **locally optimal choices** at each of its steps
 - Simple example: the **Nearest Neighbor** algorithm for the **Traveling Salesman Problem** (TSP)
 - In Dijkstra: we maintain a table $d[v]$ of **partially minimum distances** from u to v computed over a subset of all paths from u to v
 - In Prim we maintain a table $c_t[v]$ of **locally minimum edge costs** of a partial spanning subtree that is progressively grown from a starting node u
- Greedy strategies are often quite natural
 - But a too simple greedy approach often results on wrong algorithms, with greedy TSP an example
 - Also the greedy ideas behind Dijkstra and Prim are not that obvious
 - And less so that they result in correct algorithms
- **Kruskal's** is another, much clearer example of a greedy algorithm to obtain a MST

A First Look at Kruskal's Algorithm

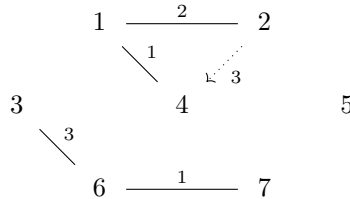
- Main idea: sort the edges of E in a PQ by increasing costs and build a graph (forest?) of partial STs
 - Starting from single node trees $T_u = (\{u\}, \emptyset)$ and
 - Adding edges from the PQ that do not produce cycles
- Example:



How to Apply Kruskal?

- Solving ties lexicographically, the sorted edges are
 $(1, 4), (6, 7), (1, 2), (2, 4), (3, 6), (1, 3), (4, 7), (5, 7), (4, 6), \dots$

- We add edges to the partial ST as



- But trying to add $(2, 4)$ we get a **cycle**, so we drop it and add next $(3, 6)$
- We keep going until we get an MST

Elements of Kruskal's Algorithm

- To implement Kruskal we need a PQ, a way of storing the selected edges and a way to maintain the forest of partial subtrees and to detect cycles
- No problem with the PQ and we can simply gradually build the final MST graph over the Kruskal forest of the partial subtrees
- At first sight maintaining trees and detecting cycles in them looks complicated and costly
- However, observe that (u, v) **gives a cycle iff u and v are in the same subset $V_{T'}$ of the vertices of a tree T' in the Kruskal forest**
 - 2 and 4 are in the set $\{1, 2, 4\}$
 - Thus we do not need to work with trees but with **subsets**
- We do this with a new abstract data type, the **Disjoint Set**

2.2 The Disjoint Set Abstract Data Type

Disjoint Set

- A **Disjoint Set (DS)** over a universal set U is a dynamic family S of disjoint subsets of U (i.e., a **partition** of U), each of which is **represented** by a certain element x and that has the following primitives:
 - `init_DS(U, S)`: receives the universal set U and returns the initial S as the family of atomic subsets $\{\{u\} : u \in U\}$
 - `find(x, S)`: receives an element $x \in U$ and returns the representative of the subset S_x of S that contains x
 - `union(x, y)`: receives two representatives x, y , computes their union $S_x \cup S_y$ and returns a representative of the subset $S_x \cup S_y$

Observations on the Disjoint Set

- The subsets of a Disjoint Set are never split

- They can only change to bigger subsets
- The Disjoint Set is never empty
- After `init_DS` we start with a partition with $|U|$ subsets;
 - Thus, the maximum number of unions is $|U| - 1$
- Even if we don't have yet a data structure for DS, its primitives allow us to write a first pseudocode for Kruskal

Kruskal's Algorithm

```
def kruskal(G):
    T = (V, E={})                                #empty graph for the MST
    init_DS(V, S)                                # 1

    Q = pq()
    for all (u, v) in E:
        Q.put((c(u, v), (u, v)))                # 2

    while not Q.empty():                          # 3
        _, (u, v) = Q.get()                      # 4
        x = find(u, S)
        y = find(v, S)
        if x != y:
            add((u, v), E)                       # 6
            union(x, y, S)                       # 7

    return T
```

Observations on Kruskal's Algorithm

- Here we build the MST T on a graph initially without edges (when writing a program this may change)
- The algorithm may not return a ST, for instance if G is not connected
 - We can control this introducing a counter c and increasing it when a new edge is added to L
 - c should have the value $|V| - 1$ when the PQ is empty
 - Exercise: add code to control this situation
- The maximum number of unions is $|V| - 1$
- Even if we achieve a efficient implementation of `union` and `find`, the cost of Kruskal will be at least $O(|E| \log |V|)$ because of building the PQ in (1)
 - So it won't improve on Prim

A First Data Structure for DS

- We assume $V = \{1, \dots, N\}$
- A simple idea is keep each subset in a list with the representative in the first node
- We also construct a pointer (dict?) table $T[\]$ where $T[i]$ points to the list that contains i
- The cost of `find` is clearly $O(1)$

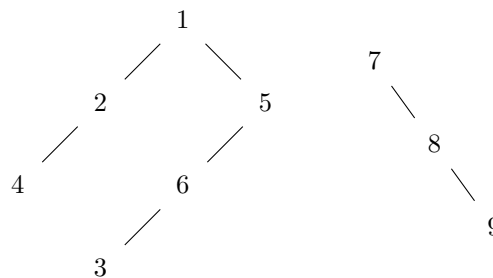
- To implement `union(x, y, S)` we can concatenate the list $T[y]$ after the list $T[x]$ and then make sure that for each j in $T[y]$ we have $T[j] = T[x]$
- However this is not satisfactory as the cost of the union is then
 - $|T[x]|$ (to find the end point) plus
 - $|T[y]|$ (to reset the pointers of $V_{T(y)}$)
- This can be improved upon but we will do something different

Our Data Structure for DS

- Our data structure stores DS as trees (**not to be confused** with those of the Kruskal forest)
- The representative x of a subset S is at the **root** of the subset tree T_S
- The cost of `union(x, y, S)` is then just $O(1)$, as we simply make, say, T_{S_y} a child subtree of the x root
- To implement `find(u, S)` first we need a fast way to locate the tree of u and, then, to go from the u node to the root
- This can be easily done if we place the subsets on a table $p[]$:
 - $p[u]$ is the index of the father of u
 - $p[x] = -1$ for a root x , i.e., a representative

An Example of the DS for the DS

- For a subset partition over the universal set $[1, 2, 3, 4, 5, 6, 7, 8, 9]$



the associated table would be

$[-1, 1, 6, 2, 1, 5, -1, 7, 8]$

Union and Find over Trees

- To initialize the DS we simply need $p[i] = -1$ for all i
- The simplest pseudocode for `find` is

```

def find(u, p):
    while p[u] != -1:
        u = p[u]
    return u
  
```

- The pseudocode for a naive union is

```
def union(x, y, p):
    p[y] = x    #join second tree to first
    return x
```

Improving Union

- Since the cost of `find` is $O(\text{height}(T_x))$ it is clear that we should join the shorter tree into the taller one
- For this we need to keep a tree's height h
 - We simply can change $p[x]$ at the root x from -1 to $-h$

- We then change the pseudocode for union as

```
def union_height(x, y, p):
    if p[y] < p[x]:    #T_y is taller
        p[x] = y; return y
    elif p[y] > p[x]:  #T_x is taller
        p[y] = x; return x
    else:              #T_x, T_y have the same lenght
        p[y] = x; p[x] -= 1; return x
```

- We also change the while condition on `find` to

```
while p[u] >= 0:
```

The Cost of Find

- **Proposition.** If $\text{prof}(T)$ denotes the depth of a DS tree T , we have $\text{prof}(T) \leq \lg |T|$
- **Proof Sketch:**

- Use induction on $|T|$, with an obvious base case $|T| = 1$
- Assume $\text{prof}(T') \leq \lg |T'|$ for $|T'| < |T| = k$ and that we join T_y into T_x with $|T_x \cup T_y| = k$
- If $\text{prof}(T_y) < \text{prof}(T_x)$,

$$\text{prof}(T_x \cup T_y) = \text{prof}(T_x) \leq \lg |T_x| \leq \lg |T_x \cup T_y|$$

and the same argument works when $\text{prof}(T_x) < \text{prof}(T_y)$,

- If $\text{prof}(T_y) = \text{prof}(T_x)$ and, say, $|T_y| \leq |T_x|$,

$$\begin{aligned} \text{prof}(T_x \cup T_y) &= 1 + \text{prof}(T_y) \leq 1 + \lg |T_y| = \lg 2|T_y| \\ &\leq \lg |T_x \cup T_y| \end{aligned}$$

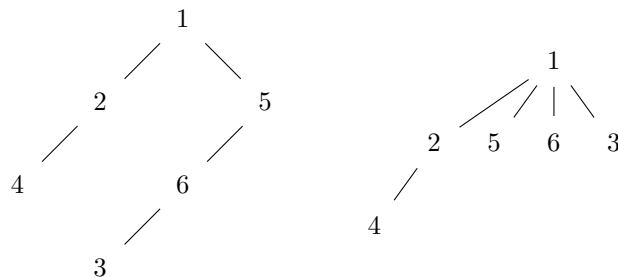
Improving Find

- Thus, the cost of `find(x, p)` is also $O(\log |S_x|) = O(\log N)$
- Moreover, we can further improve on this
- Observe that when finding the representative of u we also find the **representative of all the v between u and the root of its tree**

- We can thus change `find` to update `p[v]` for all v between u and the root
- In other words, we can **compress the path** from u to the root

The Effect of Path Compression

- Left: tree state after `find(3)`; right: state after `find_cc(3)`



Find with Path Compression

- Recall that after finding the representative of u , we also know it for all the other nodes between u and the root of the tree
- We thus improve `find` as follows:

```
def find_cc(u, p):
    # find the representative
    z = u
    while p[z] >= 0:
        z = p[z]

    # compress the path from u to the root
    while p[u] >= 0:
        y = p[u]
        p[u] = z
        u = y
    return z
```

Path Compression and Union by Rank

- The problem is now that, after `find`, we no longer have in $-p[x]$ the tree's height
- We do nothing about this other than calling $-p[x]$ the tree's **rank**
- We change nothing on `union` although it is no longer a union by height but a **union by rank**
- However the joint cost of unions and finds considerably improves
- **Proposition:** *If on a DS with N elements we do L unions by rank and $M = \Omega(N)$ path compression finds, the overall cost is*

$$O(L + M \lg^* N)$$

The \lg^* Function

- We define $\lg^* H = K$ if K is the smallest integer such that after K binary logs we have

$$\lg(\dots \lg(\lg H) \dots) \leq 1$$

- For instance $\lg^* 65536 = \lg^* 2^{16} = 4$, but then

$$\lg^* 2^{65536} = 1 + \lg^* 2^{16} = 5$$

- Now 2^{65536} is a huge number:
 - Find out how many digits its decimal expression has (easy)
 - Then try to write it using millions, billions, googols and so on! ;-)
- For practical purposes $\lg^* H = O(1)$

Back to Kruskal's Algorithm

- Assume we work with union by rank and path compression and go back to Kruskal's pseudocode

```
def kruskal(G):
    T = (V, E={})                #empty graph for the MST
    p = init_DS(V)               # 1
    Q = pq()

    for all (u, v) in E:
        Q.put( (c(u, v), (u, v))) # 2

    while not Q.empty():         # 3
        _, (u, v) = Q.get()      # 4

        x = find_cc(u, p)        # 5
        y = find_cc(v, p)

        if x != y:
            add((u, v), E)       # 6
            union(x, y, p)       # 7

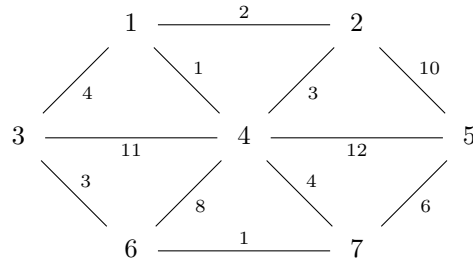
    return T
```

The Cost of Kruskal's Algorithm

- Clearly the cost of (1) is $O(|V|)$ and that of (2) is $O(|E| \log |V|)$
- The cost of (4) accumulated over (3) is again $O(|E| \log |V|)$
- Since the single cost of (6) and (7) is $O(1)$ and only happens when $x \neq y$, their accumulated costs are $O(|V|)$
- Finally, since we must do at least one `find_cc` for each node, the total number is $\Omega(N)$ and, therefore, the cost of (5) accumulated over (3) is $O(|E| \lg^* |V|)$, that is, essentially $O(|E|)$
- Summing things up, the cost of Kruskal is $O(|E| \lg |V|)$, dominated by the PQ operations
- In particular the DS operations do not penalize the algorithm

Applying Kruskal's Algorithm

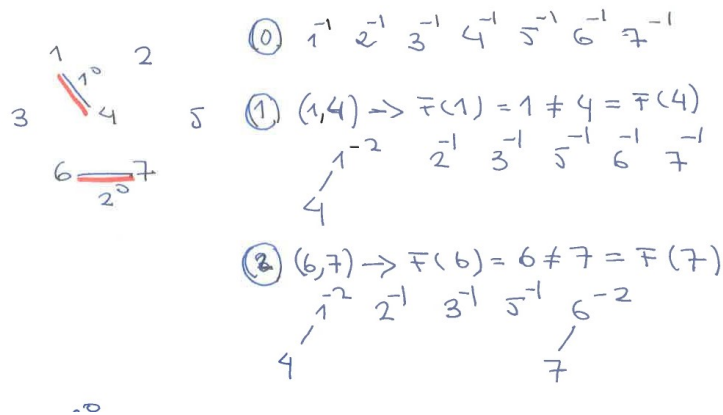
• **Example:**



- The PQ is $(1, 4), (6, 7), (1, 2), (2, 4), (3, 6), (1, 3), (4, 7), (5, 7), (4, 6), (2, 5), (3, 4), (4, 5)$

Applying Kruskal's Algorithm (II)

- We maintain separately the Kruskal forest and the DS forest



Applying Kruskal's Algorithm (III)

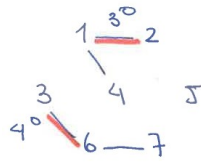
- We process the remaining edges from the PQ

Applying Kruskal's Algorithm (IV)

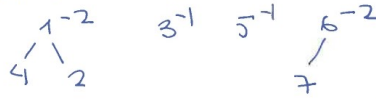
- We process the remaining edges from the PQ

Applying Kruskal's Algorithm (V)

- We process the remaining edges from the PQW until it is empty
- The MST may not change but the DS forest may

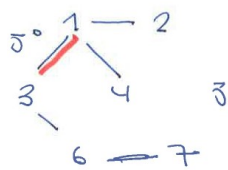
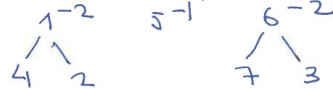


$$(3) (1,2) \rightarrow F(1) = 1 \neq 2 = F(2)$$

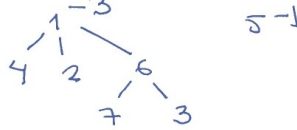


$$(4) (2,4) \rightarrow F(2) = 1 = F(4) \rightarrow \times$$

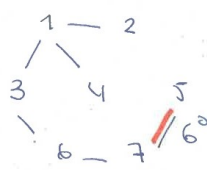
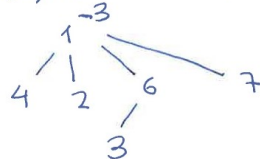
$$(5) (3,6) \rightarrow F(3) = 3 \neq 6 = F(6)$$



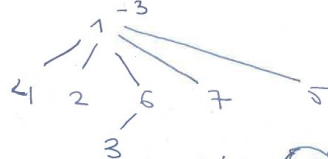
$$(6) (1,3) \rightarrow F(1) = 1 \neq 6 = F(3)$$



$$(7) (4,7) \rightarrow F(4) = 1 = F(7)$$

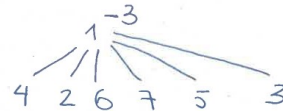


$$(8) (5,7) \rightarrow F(5) = 5 \neq 1 = F(7)$$



$$(9) (4,6): \text{in combination } (10) (2,5): \text{idem}$$

$$(11) (3,4) \rightarrow F(3) = 1 = F(4) \rightarrow$$



2.3 Correctness of Prim and Kruskal

Cuts and Minimal Crossings

- Assume we have an undirected weighted graph $G(V, E)$ with cost c
- A **cut** P of G is a partition of V into two disjoint subsets $P = (S, V - S)$
- An edge (u, v) **crosses** P if either $u \in S$ and $v \in V - S$ or viceversa
- A subset $A \subset E$ **preserves** P if no edge in A crosses P
- An edge (u, v) that crosses P is **minimal** w.r. to P if $c(u, v) \leq c(w, z)$ for any other edge (w, z) that crosses P

A Meta MST Algorithm

- Consider the following meta–algorithm to find MSTs

```
def metaMST(G, c):
    T = (V, E={})           #empty graph for the MST

    while |E| < |V|:
        find a cut P preserved by E
        select (u, v) minimal w.r. to P
        add((u, v), E)

    return T
```

- Notice that `metaMST` is also a kind of greedy meta–algorithm
 - At each step a minimal edge is added to the partial MST

Prim as an Example of metaMST

- Recall that Prim works with a table $v[]$ of seen nodes and that the nodes still in Q are ordered by their cost at insertion
- Assume that a node v has been extracted from Q just before is marked as seen, and take
 - $P = (\{seen\ nodes\}, \{others\})$
 - $E = \{(p[w], w) : w \in \{seen\ nodes\}\}$
- Then we have
 1. E preserves P for if $(p[w], w) \in E$, both w and $p[w]$ are seen
 2. $(p[v], v)$ crosses P , for v is still unseen but $p[v]$ was processed when v entered Q , i.e., it is seen by now
 3. If other (w, z) crosses P we have $v[w] = T$, $v[z] = F$ and, hence, $z \in Q$ since it is adjacent to the already seen node w
 4. Since we extract v but not z , $c(p[v], v) \leq c(w, z)$ and, thus, $(p[v], v)$ is minimal
- Hence, Prim is a particular case of metaMST

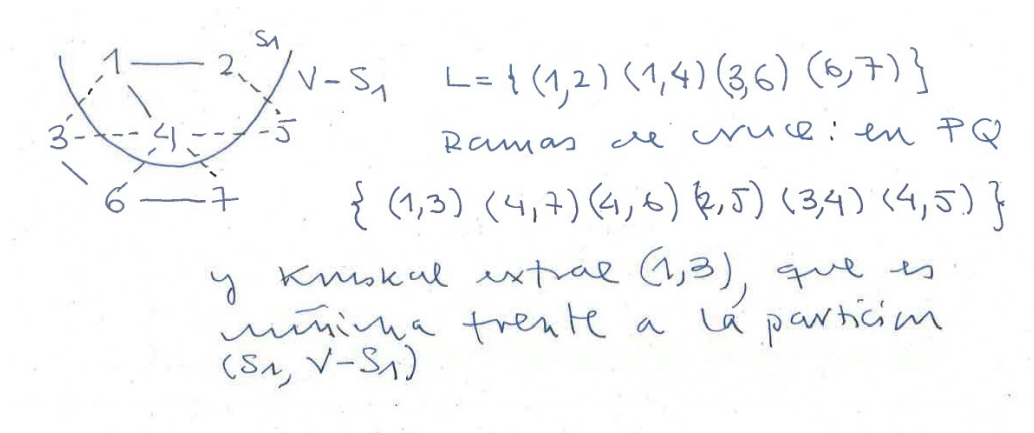
Kruskal as an Example of metaMST

- Assume that we are about to add the edge (u, v) and let

- E be the edges already selected
- $P = (S_u, V - S_u)$ where S_u is the subset of the tree T_u that contains u
- Then we have
 1. E preserves P , for if $(w, z) \in E$, w and z are in the same subtree T , which cannot happen if $w \in S_u$ and $z \in V - S_u$
 2. (u, v) crosses P by our choice of P
 3. Any other (w, z) crossing P must connect different subtrees and cannot make a cycle
 4. But then (w, z) must still be in Q : if it has left Q but is not in E , it would have made a cycle, which it cannot
 5. Thus, $c(u, v) \leq c(w, z)$ and (u, v) is minimal w.r. P
- Hence, Kruskal is a particular case of metaMST

A Kruskal metaMST step

- In the previous example, assume we are going to add $(1, 3)$
- The partition, the preserving edges and the crossing ones are



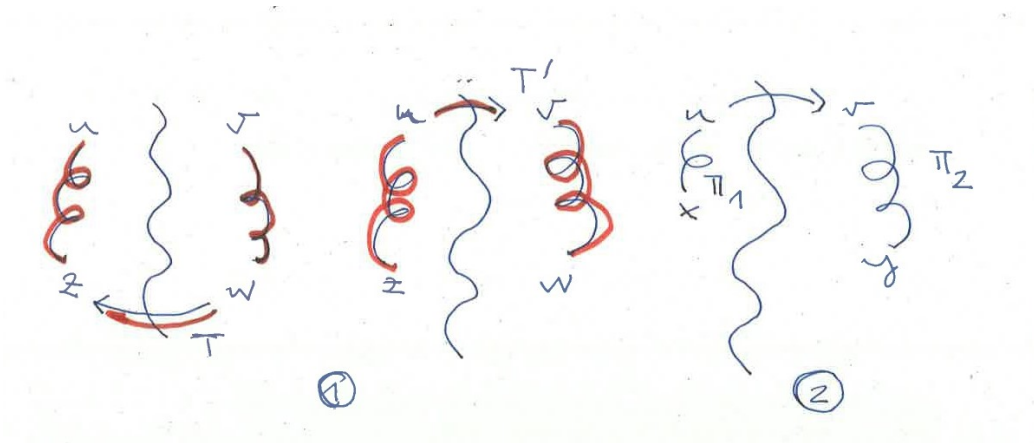
Correctness of metaMST I

- Thus, if metaMST is correct, Prim and Kruskal will also be correct
- **Proposition.** Let $G = (V, E)$ be a undirected, connected, weighted graph and assume $A \subset E$ verifies $A \subset E_T$ for some MST T . Then, if A preserves some P and (u, v) is minimal w.r. to P , we have $A \cup \{(u, v)\} \subset E_{T'}$ for some MST T'
- **Proof sketch I:**
 - Assume $T = (V, E_T)$ is a MST
 - Then $\pi = E_T \cup \{(u, v)\}$ is a cycle with an edge (w, z) that crosses P

- Define $T' = (V, E_{T'})$ with $E_{T'} = (E_T - \{(w, z)\}) \cup \{(u, v)\}$
- Clearly $c(T') \leq c(T)$ and have to prove that T' is a spanning tree
- Since $V_{T'} = V$, we just have to check T' is connected

Correctness of metaMST II

- **Proof sketch II:** Building T' and going from x to y by T'



Correctness of metaMST III

- **Proof sketch III:** let x, y be two nodes; we show they can be connected by T'
 - If x, y are in the same subset of P they can clearly be joined by T and, hence, by T' , as they coincide there
 - Assume x, y at different subsets of $P = (S_1, S_2)$ with x, u and y, v in the same sides
 - There are paths π_1 from x to u in S_1 and π_2 from v to y in S_2 ; hence they are in T and also in T'
 - Then $\pi = \pi_1 \cup \{(u, v)\} \cup \pi_2$ is a path in T' from x to y
 - Thus T' is connected, $c(T') \leq c(T)$ and $V_{T'} = V$
 - Thus T' is an MST

Loop Invariants

- The proposition says that **after each iteration the selected edges are part of a MST**
- This is an example of a **loop invariant**:
A condition that remains true after each loop and that “leads” the algorithm towards a correct solution
- The standard way to prove the correctness of an iterative algorithm is to find an adequate loop invariant for its iterations

- Example: loop invariants for InsertSort or BubbleSort
 - InsertSort: after iteration i , $i = p + 1, \dots, u$, the subtable $T[p], \dots, T[i]$ is sorted
 - BubbleSort: after iteration i , $i = u, \dots, p + 1$, the subtable $T[i], \dots, T[u]$ is sorted

Correctness of metaMST II

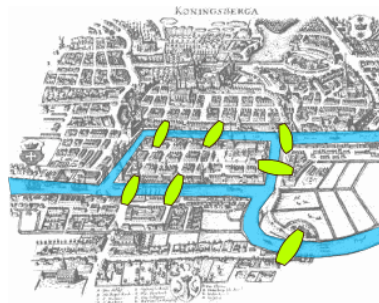
- **Corollary.** *metaMST returns a MST*
- **Proof sketch:**
 - We just exploit the loop invariant provided by the previous proposition
 - Let $L_0 = \emptyset \subset L_1 \subset \dots \subset L_{N-1}$ be the successive subsets metaMST produces
 - If L_j is a subset of some MST, the proposition shows that so is L_{j+1}
 - But obviously L_0 is a subset of some MST and, thus, so is L_{N-1} and since it has $N - 1$ edges, (V, L_{N-1}) is a MST
- **Corollary** *Prim and Kruskal return MSTs*

3 Eulerian and Hamiltonian Circuits

3.1 Eulerian Circuits

The Bridges of Königsberg

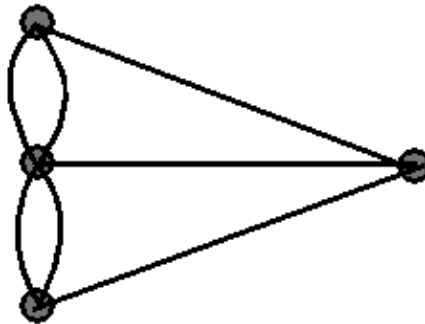
- The bridges of Königsberg (East Prussia) over the Pregel river circa 1700:



- The problem: find a promenade that crosses all bridges but only once
- Exercise: google `pregel graph`

The Bridges of Königsberg as a Graph Problem

- We can depict the bridges of Königsberg as a multigraph (i.e., we allow for multiple edges between two nodes)



- The problem: find a circuit that passes **through all edges but only once**
- Such a circuit in a multigraph is called an **Eulerian circuit (EC)**

Euler's Insight

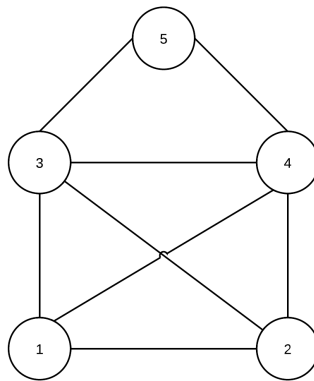
- Leonhard Euler showed in 1736 (*Solutio problematis ad geometriam situs pertinentis*) that such a circuit is not possible
- If G is an undirected graph, we define the **degree** $\deg(w)$ of a node w as the number of edges that leave w (or that enter w)
- Assume that $\pi = \{(u = u_0, u_1), \dots, (u_{K-1}, u_K = u)\}$ is an EC for G
- If $w \neq u$ is a node in π , we subtract 1 from $\deg(w)$ each time we enter w or leave it
 - Since at the end we have passed by all the edges of w , **we must have at the beginning $\deg(w)$ even**
- Similarly each time we enter u inside π we subtract 1 from $\deg(u)$ and also when we leave u ; moreover, when we start and end π we also subtract 1 from $\deg(u)$
 - Thus, **we must also have $\deg(u)$ even**

There Are No ECs in Königsberg

- It follows from the previous analysis that a necessary condition to have an EC is that $\deg(v)$ is even for all $v \in V$
- Since all the nodes in the previous multigraph have odd degrees, Euler concluded that no Eulerian circuit is possible in Königsberg
- As we shall see later, Euler also proved that the condition is sufficient:
If $\deg(v)$ is even for all nodes v of an undirected graph G , then there is an Eulerian circuit in G

Drawing Houses Without Lifting the Pen

- A child's game is to try to draw the house below without lifting the pen from the sheet



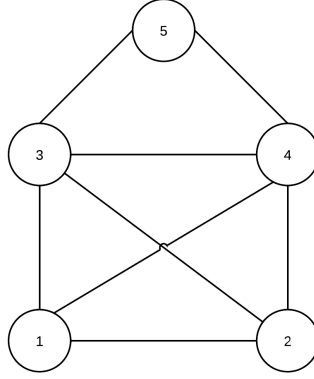
- It is very easy if we start at nodes 1 or 2 but impossible if we start from 3, 4 or 5

Euler's Insight Again

- Assume that $\pi = \{(u = u_0, u_1), \dots, (u_{K-1}, u_K = v \neq u)\}$, $u \neq v$, is such an **Eulerian path** (EP)
- If $w \neq u, v$ is a node in π , each time we enter w we subtract 1 from $\deg(w)$ and also when we leave w ;
 - Since at the end we have passed through all the edges of w , we must have at the beginning $\deg(w)$ even
- Similarly each time we enter u inside π we subtract 1 from $\deg(u)$ and also when we leave u ; moreover, since we start π at u , we also subtract 1 from $\deg(u)$
 - Thus, we must also have $\deg(u)$ odd
- Similarly each time we enter v inside π we subtract 1 from $\deg(v)$ and also when we leave v ; moreover, since we end π at v , we also subtract 1 from $\deg(v)$
 - Thus, we must also have $\deg(v)$ odd
- Thus, **a necessary condition to have an EP is that $\deg(w)$ is even for all w except the first node u and the final one v of π**

Back to Drawing Houses

- Since $\deg(1) = \deg(2) = 3$ we can find an EP for the house drawing if we start at either 1 or 2



- But since $\deg(3) = \deg(4) = \deg(5)$ even, it is impossible to draw an EP for the house starting at them
- And there is no EP in Königsberg either.

Euler's Theorem for Circuits

- **Theorem.** If $G = (V, E)$ is a connected undirected **multigraph**, there is an EC in G iff $\deg(u)$ is even for all $u \in V$

Proof sketch: We argue by induction on $|V|$

- The theorem is obviously true if $|V| = 2$ and assume it also to be true for any $G' = (V', E')$ such that $|V'| < |V|$
- Start walking from a node u and subtract from $\deg[w]$ when passing by a node w until we arrive at v such that $\deg(v) = 0$ after we enter v and, thus, cannot leave it
- It is easy to see that $v = u$ for if not, $\deg[v]$ is odd
- Thus, we have found a cycle π
- Set $G' = (V', E')$ with $V' = V - \{w : \deg_G[w] = \deg_\pi[w]\}$ and $E' = E - E_\pi$

Euler's Theorem for Circuits II

Proof sketch (cont.):

- Since $|V'| \leq |V| - 1$ and $\deg_{G'}(w) = \deg_G(w) - \deg_\pi(w)$ is even, we can apply induction on the connected components G_1, \dots, G_K of G'
- Consider the nodes in π such that $\deg_\pi[w] < \deg_G[w]$ and let's enumerate these components by their first appearances w_1, \dots, w_K in π
- Each w_i is in the connected component $G_i = (V_i, E_i)$, which has an EC π_i
- Let $\tilde{\pi}$ the circuit we get “collating” these π_i with π
- Then we have

$$\# \text{ edges in } \tilde{\pi} = \# \text{ edges in } \pi + \sum_i |E_i| = |E|$$

and, hence, $\tilde{\pi}$ is an EC

Euler's Theorem for Paths

- **Corollary.** *If G is a connected undirected graph, there is an EP π in G iff $\deg(w)$ is even for all $w \in V$ except for two vertices u and v . Moreover, then π starts at u and ends at v or viceversa*

Proof sketch: We just show the condition to be sufficient:

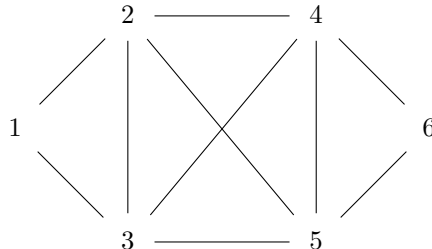
- Consider $G' = (V, E' = E \cup \{(u, v)\})$, i.e., we add an extra edge (u, v) to E
- Since $\deg_{G'}(u) = \deg_G(u) + 1$, $\deg_{G'}(v) = \deg_G(v) + 1$ and $\deg_{G'}(w) = \deg_G(w)$ for all other w , all the G' degrees are even and there is an EC π' in G'
- Let's write π' as $\pi' = \{(v, z), \dots, (w, u), (u, v)\}$, with the last edge the one we added to get G' .
- Then removing this edge we get the EP $\pi = \{(v, z), \dots, (w, u)\}$.

How to Find an EC

- We simply to follow the proof's argument
- We start at any u_1 and build $\pi_1 = \{(u_1, v_2), \dots, (v_{K-1}, v_K)\}$ subtracting 1 from $\deg(w)$ each time we enter or leave w and where we stop because after entering v_K we have $\deg(v_K) = 0$
 - It is then clear that $u_1 = v_K$
- Let $G_1 = (V_1, E_1)$ the graph obtained after removing π_1 from E and all the $w \in V$ for which $\deg(w) = 0$ after π , i.e., for which $\deg_\pi(w) = \deg_G(w)$
 - Clearly u_1 at least will be removed, i.e., $|V_1| < |V|$
 - If $|V_1| = 0$, clearly π_1 is an EC in G
 - If however $|V_1| > 0$, there is a first u_2 in π_1 such that $\deg_{G_1}(u_2) > 0$
 - We can thus **restart the above process on G_1** obtaining a new circuit π_2 and a “remaining” graph G_2
- If we repeat the preceeding and find circuits π_1, \dots, π_M until $V_M = \emptyset$, then we can “collate” the π_j circuits to get an EC π for G

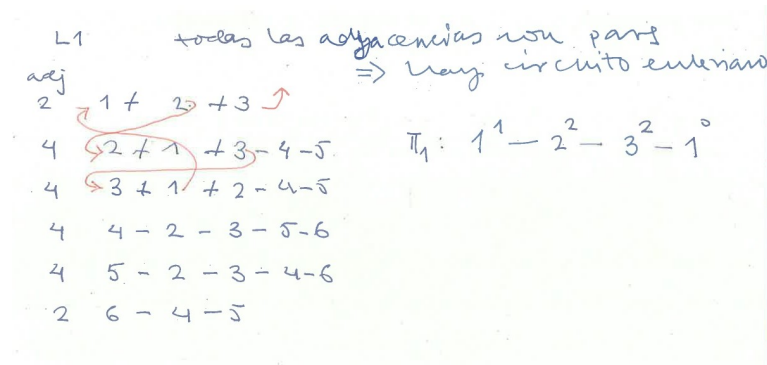
How to Find an EC II

- This is essentially Hierholzer's algorithm
- We do not write a pseudocode (good exercise!) but it is clear that its cost will be $O(|E|)$
- **Example:**

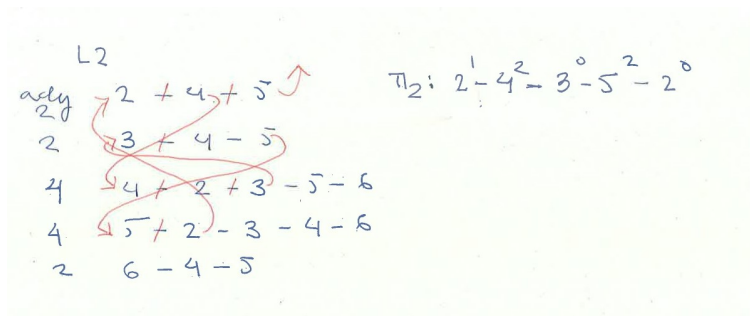


EC Steps I

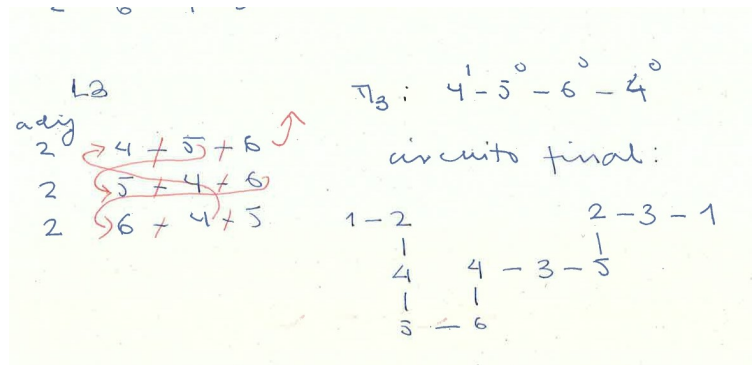
- The first steps to find an EC are

**EC Steps II**

- The next steps to find an EC are

**EC Steps III**

- The final steps to find an EC are



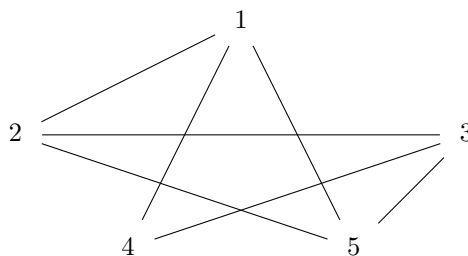
3.2 Hamiltonian Circuits and an Excursion on Complexity Theory

Hamiltonian Circuits

- If G is an undirected connected graph, a **Hamiltonian circuit** (HC) is a circuit on G that visits **only once each node** other than the initial
- Finding HCs may be trivial in some cases, such as complete graphs
- There are also sufficient conditions for special graphs
- But for general graphs, while finding ECs has an $O(|E|)$ cost, finding HCs is much costlier
- In fact, essentially the only general algorithm is an exhaustive search with backtracking

Hamiltonian Circuits II

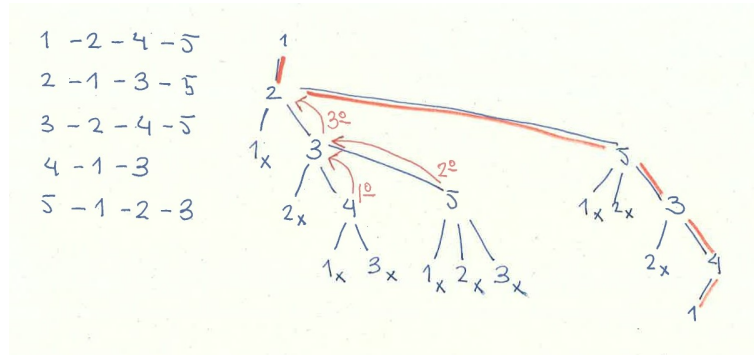
- **Example:**



- Since the number of node orderings is $N!$, the search's cost can be very high
- Actually, finding HCs in general graphs is an example of an **NP-complete** problem

Backtracking Search

- An example of a HC search



P and NP I

- We will make a brief (and light) excursion on Complexity Theory
- We consider **decision problems** \mathcal{P} : there is a set of **solution inputs** $S_{\mathcal{P}}$, for which **the decision on an input I is 1 iff $I \in S_{\mathcal{P}}$**
 - To decide whether a graph has an EC or HC is a decision problem but notice that an algorithm does not have to actually find an EC or HC to solve them
 - Optimization problems can be partially reduced to decision problems using a bound C : change *find the optimum* by *find a solution with cost $\leq C$*
- For an input I we can consider its size $|I|$ to be the number of bits needed to store it
- We say that \mathcal{P} is in the class P if there is an algorithm A with cost polynomial on $|I|$ that solves \mathcal{P} , i.e., $A(I) = 1$ iff $I \in S_{\mathcal{P}}$
 - Note that to be in class P does not mean that A is efficient: if its cost is $O(|I|^{1000})$, \mathcal{P} is in P

P and NP II

- Decision-EC is in P : we check in linear time whether or not there are ECs in G by counting degrees and checking that they are even
- An algorithm $C(I, S)$ is a **certifier** for \mathcal{P} if
 - For every input $I \in S_{\mathcal{P}}$ there is at least another input S to C such that $C(I, S) = 1$
 - If $I \notin S_{\mathcal{P}}$, then $C(I, S) = 0$ no matter which S is used
- We can see S as a kind of certificate (solution?) for I that the C validates
 - For the EC or HC problems, S can just be a possible EC or HC
- We say that \mathcal{P} is in the class NP if there is a certifier C that runs in polynomial time on the sizes $|I|$ and $|S|$

- For instance, if $I = G$ and S is a possible CH, we can check it in polynomial time
- Thus HC belongs to NP

P and NP III

- Clearly $P \subset NP$: if $\mathcal{P} \in P$ and A solves it, set $C(I, S) = A(I)$; then
 - If $I \in S_{\mathcal{P}}$, then $C(I, S) = A(I) = 1$ for any S
 - If $I \notin S_{\mathcal{P}}$, we will have $C(I, S) = A(I) = 0$ no matter the S presented
- Big question: $P = NP$?
- If yes, there would be a polynomial time algorithm for HC
- It is one of the Millenium Problems of the Clay Mathematics Institute with a 1M \$ prize
 - For more details see [Clay Institute's P vs NP page](#)
- General opinion: $P \neq NP$
- Reason: **NP -complete problems**

NP-complete Problems

- We say that \mathcal{P}_1 is **reducible** to \mathcal{P}_2 if there is a map

$$T : \{\text{inputs of } \mathcal{P}_1\} \rightarrow \{\text{inputs of } \mathcal{P}_2\}$$
 such that I_1 **has a solution for** \mathcal{P}_1 **iff** $T(I_1)$ **has a solution for** \mathcal{P}_2
 - Or: $I \in S_{\mathcal{P}_1}$ iff $T(I) \in S_{\mathcal{P}_2}$
- Thus, **if A is an algorithm that solves \mathcal{P}_2 , then $A \circ T$ solves \mathcal{P}_1 :**

$$I \in S_{\mathcal{P}_1} \text{ iff } T(I) \in S_{\mathcal{P}_2} \text{ iff } A(T(I)) \equiv A \circ T(I) \equiv 1$$
- If T has polynomial cost, we say that \mathcal{P}_1 is **polynomially reducible** to \mathcal{P}_2
- We say that problem \mathcal{P} is **NP -complete** if **any other** $\mathcal{P}' \in NP$ is polynomially reducible to \mathcal{P}
- Notice that we can **prove that** $P = NP$ if we just find one NP -complete problem \mathcal{P} such that $\mathcal{P} \in P$

Is There Any NP -complete Problem?

- At first sight the NP -complete definition seems very strict so a natural question is whether there any such problem
- Answer: yes, and in fact many!!
 - HC is such a problem
 - TSP will be another

- The first (basically) *NP*-complete problem found is 3-SAT
- Given a Boolean expression B written using only AND, OR, NOT operators, and parentheses, the **satisfiability problem (SAT)** is to decide whether there is some assignment of T and F to the variables that will make B true
- The k -SAT problem deals with expressions in **conjunctive normal form** (i.e., as a sequence of OR clauses joined by AND) with k variables or their negation per clause

Cook's Theorem

- Example: 3-SAT deals with expressions like

$$(x_{11} \text{ OR } !x_{12} \text{ OR } x_{13}) \text{ AND } (!x_{21} \text{ OR } x_{22} \text{ OR } !x_{23}) \text{ AND } (x_{31} \text{ OR } !x_{32} \text{ OR } x_{33}) \text{ AND } \dots$$
- **Cook's Theorem (1971)**: 3-SAT is NP-complete
 - However, 2-SAT $\in P$
- More to read: Chapter 5 of H. Wilff's book [Algorithms and Complexity](#)
- Much more to read: M.R. Garey and D.S. Johnson. **Computers and Intractability: A Guide to the Theory of NP-Completeness**. W.H. Freeman, 1979.
- But are P , NP and *NP*-complete problems just academic curiosities?

3.3 The Traveling Salesman Problem

The Traveling Salesman Problem

- TSP: Given a weighted complete graph G , **find a HC** (trivial) **with minimum cost**
- It is an optimization problem with obvious practical interest: many persons have to solve it every morning
 - Decision version: given a weighted complete graph G and a bound C , is there a HC π such that $c(\pi) \leq C$?
- TSP is **NP-hard**: every problem in NP can be polynomially reduced to TSP
 - Or P is NP-hard problem if 3-SAT or HC reduce polynomially to P
 - A NP-hard problem may not have to be NP-complete (e.g., the halting problem) or to be a decision problem (e.g., TSP)
 - Also, TSP-decision for general graphs is NP-complete
 - But TSP-decision is also NP-complete for “real world” problem versions, such as for cities in the plane with Euclidean distances
- Many related problems of great practical interest in planning, logistics or DNA sequencing are also NP-complete

From TSP to HC

- Fact: **HC is polynomially reducible to TSP**
- Assume `tsp(V, c)` is a routine that returns the TSP solution for G with cost c and consider the following routine for HC:

```
def tsp_2_hc(V, E):
    for any u, v in V:
        if (u, v) in E:
            c(u, v) = 1
        else:
            c(u, v) = 2

    p = tsp(V, c)
    if cost(p) == |V|:
        return p
```

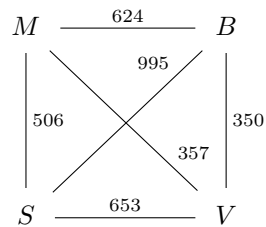
- `tsp_2_hc` solves HC for π is a HC on G iff $c(u, v) = 1$ for any $(u, v) \in \pi$ iff $c(\pi) = |V|$
- Thus TSP has not only practical but also theoretical interest

A TSP Example

- Simple example:

```
["madrid", "barcelona", "sevilla", "valencia"]
```

- The (complete) graph is



- More examples in [Traveling Salesman Algorithms](#)

A Greedy TSP Solution

- Simple greedy approach: Nearest-Neighborhood (NN) TSP, that simply visits the nearest unseen city

```
def nn_tsp_circuit(distance_matrix, node_ini=0):
    num_cities = distance_matrix.shape[0]
    circuit = [node_ini]

    while len(circuit) < num_cities:
        current_city = circuit[-1]

        # sort cities in ascending distance from current
        options = list(np.argsort(distance_matrix[ current_city ]))

        # add first city in sorted list not visited yet
        for city in options:
            if city not in circuit:
                circuit.append(city)
                break

    return circuit + [node_ini]
```

What Can We Do About TSP?

- The greedy solution of the previous problem is M, V, B, S, M
- On average, NN gives a path that is about 25% longer than the optimum
 - But one can set up special instances of TSP where NN gives the worst route
- If c satisfies the triangle inequality $c(u, v) \leq c(u, z) + c(z, v)$ for any z , we have

$$c(\pi_{NN}) = O(\log |V|) \times c^*,$$

with π_{NN} the NN solution and c^* the optimal cost

- TSP has great practical importance, but there is no cost effective **exact** algorithm for general graphs
 - So, it may be very hard to find the best route to, say, deliver mail (at least in big cities)
- Q: What can we do?

Approximation Algorithms

- Alternative: **approximate** algorithms
- **Definition:** Given an optimization problem \mathcal{P} , an **approximate algorithm** for \mathcal{P} with bound $\lambda \geq 1$ is an algorithm A that for every input I returns a solution $s_A(I)$ such that

$$c^*(I) \leq c(s_A(I)) \leq \lambda c^*(I)$$

with $c^*(I)$ the optimal cost for \mathcal{P} on I

- NN is not exactly an approximate algorithm for TSP, since its bound is $O(\log |V|)$ and depends on $|V|$

Approximation Algorithms for TSP

- **Proposition:** *If the cost function is Euclidean, i.e., it verifies*

$$c(u, v) \leq c(u, w) + c(w, v) \text{ for all } u, v, w \in V,$$

then there is an approximate algorithm for TSP with $\lambda = 2$

- Algorithm:

```
def euclideanTSP(g, c):
    find a MST t on g
    duplicate its edges to obtain a graph g_1

    #now each node in g_1 has degree 2 and there is an EC
    find a EC p_1 in g_1

    shortcut seen edges in p_1 to get HC p
    return p
```

Approximation Algorithms for TSP• **Proof sketch:**

- Let T_1, π_1 and π be the MST, the EC and the HC returned by the algorithm
- Let π^* be an optimal HC and remove an edge on π^* to get a spanning tree T^*
- Since T_1 is an MST, we have $c(T_1) \leq c(T^*) \leq c(\pi^*)$
- By the Euclidean distance property, if we shortcut the segment $u \rightarrow w \dots \rightarrow z \rightarrow v$ to $u \rightarrow v$, we have

$$\underbrace{c(u, v)}_{\pi} \leq \underbrace{c(u, w) + c(w, x) + \dots + c(z, v)}_{\pi_1}$$

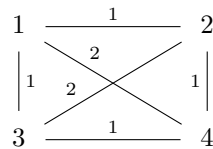
- We then we conclude that

$$c(\pi) \leq c(\pi_1) = 2c(T_1) \leq 2c(\pi^*)$$

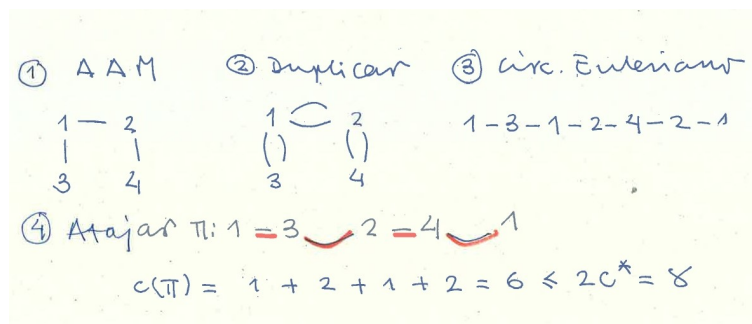
- The **Christofides** algorithm improves this to $\lambda = 1.5$ (see [this article](#) in Wired for more about the algorithm)

Approximation Algorithms for TSP II

- To learn more: Johnson, McGeoch, [The Traveling Salesman Problem: A Case Study in Local Optimization](#)
 - Or the movie [The Travelling Salesman](#)
- **Example**

**Applying The Algorithm**

- The steps to find an approximate TSP solution



4 An Excursion on DNA Sequencing

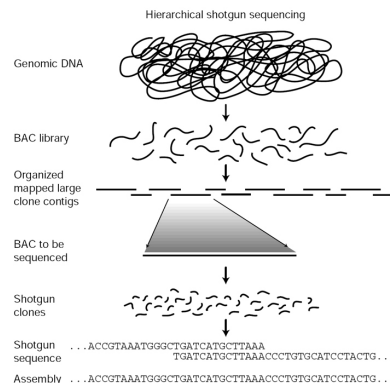
4.1 Hamilton, Euler and DNA Sequencing

DNA Sequencing

- **Note:** this is a very, very light description of DNA Sequencing
- Goal: decompose a gene into a sequence of four letters $\{A, C, G, T\}$ that correspond to DNA bases
- **Shotgun sequencing** follows a four step process:
 - Blast the gene into random short fragments (“reads”) of 100–500 bases
 - Identify read subsequences by hybridizing them on a DNA microarray
 - Reconstruct each read from these subsequences
 - Reconstruct the entire gene from the reads
- First two steps: biochemistry
- Third step: Hamiltonian or (better) Eulerian circuits
- Fourth step: compute the Shortest Superstring Problem solving TSP (plus more algorithms and a lot of biochemistry)

Shotgun Sequencing

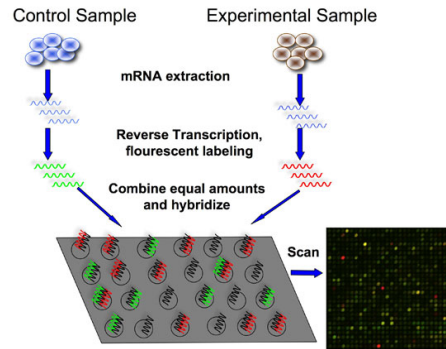
- Idealized hierarchical shotgun sequencing strategy



From [Nature](#)

Microarray Hybridization I

- Scheme of the process:



From bitesizebio.com/7206/introduction-to-dna-microarrays

Microarray Hybridization II

- The process steps are:
 - Put all the possible length ℓ probes, i.e., DNA subsequences of a fixed length ℓ , into the spots of a microarray
 - Put a drop of fluorescently labeled DNA into each microspot of the array
- The DNA fragment hybridizes with those microspots that are complementary to a certain substring of length ℓ of the fragment
 - Thus, the DNA subsequences in those microspots are also part of the DNA fragment to identify
- This way we get all possible length ℓ subsequences that make the fragment but they are **unordered**

ℓ -mers and the Spectrum

- We call the sequence on each one of the probes an **ℓ -mer**
- The **ℓ -spectrum** $sp(S, \ell)$ of a sequence S is the set of all the ℓ -mers from S
- For instance, $S = \text{[TATGGTGC]}$ we have

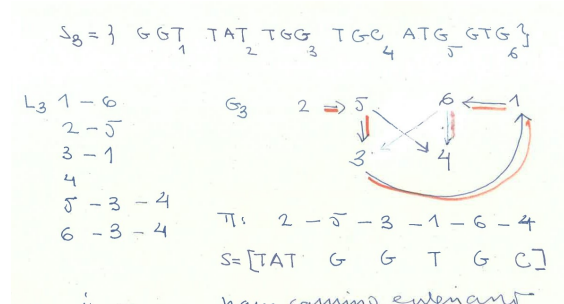
$$sp(S, 3) = \{\text{TAT}, \text{ATG}, \text{TGG}, \text{GGT}, \text{GTG}, \text{TGC}\}$$
- We have $|sp(S, \ell)| \leq |S| - \ell + 1$
- After hybridization, the hybridized probes in the microarray give us **an unordered version of** $sp(S, \ell)$ that we have to correct to recover S
- The **overlap** $\omega(s_1, s_2)$ between two ℓ -mers s_1, s_2 is the longest length of a suffix of s_1 that is also a prefix of s_2
- We clearly have $\omega(s_1, s_2) \leq \ell - 1$ and if s_2 follows s_1 in S , we must have $\omega(s_1, s_2) = \ell - 1$

Sequencing by Hamiltonian Paths

- We can reconstruct the sequence S by finding an ordering s_{i_1}, \dots, s_{i_K} of $sp(S, \ell)$ such that $\omega(s_{i_j}, s_{i_{j+1}}) = \ell - 1$
- This suggests to define the graph $G_\ell(S) = (V_\ell, E_\ell)$ where
 - $V_\ell = sp(S, \ell)$ and
 - $(s, s') \in E_\ell$ iff $\omega(s, s') = \ell - 1$
- Notice that reconstructing S is equivalent to **pass once through all the nodes of $G_\ell(S)$**
- In other words, **we can reconstruct S by finding a Hamiltonian path in $G_\ell(S)$**

Sequencing by Hamiltonian Paths II

- Example: consider $s = [\text{TATGGTGC}]$ and the unordered 3-spectrum
 $sp(S, 3) = \{\text{GGT}, \text{TAT}, \text{TGG}, \text{TGC}, \text{ATG}, \text{GTG}\}$
- By inspection, the adjacency list and graph, the HC and the recovered sequence are



Sequencing by Eulerian Paths

- The obvious problem of HP sequencing is the lack of efficient algorithms to solve the HP problem
- Alternative: **try to have ℓ -mers on the edges instead of on nodes**:
 - If $s \in sp(S, \ell)$ and s_1 is its $\ell - 1$ prefix and s_2 its $\ell - 1$ suffix, we can consider s as the edge connecting nodes s_1 and s_2
 - Now we have $\omega(s_1, s_2) = \ell - 2$
- We define now the graph $G_{\ell-1} = (V_{\ell-1}, E_{\ell-1})$ where
 - $V_{\ell-1} = sp(S, \ell - 1)$ and
 - $(s, s') \in E_{\ell-1}$ iff they are respectively prefix and suffix of an $s \in sp(S, \ell)$
 - Equivalently, $(s, s') \in E_{\ell-1}$ iff $\omega(s, s') = \ell - 2$

- Notice that now reconstructing S is equivalent to **pass once over all the edges of $G_{\ell-1}$**
- In other words, **we can reconstruct S by finding a EP in $G_{\ell-1}$**

Eulerian Circuits on Directed Graphs

- However, $G_{\ell-1}$ is a **directed** graph: we have to adapt the Eulerian circuit/path theory to these graphs
- In an directed graph $G(V, E)$ we have to distinguish between **incident and adjacent edges**
- For any $u \in V$, we say that (u, v) is an **adjacent** (outgoing) edge and (w, u) an **incident** (incoming) edge
- The **indegree** $in(u)$ of u is the number of incoming edges to u
- The **outdegree** $out(u)$ is the number of outgoing edges from u

Eulerian Circuits on Directed Graphs II

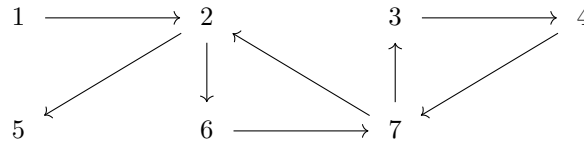
- Assume that $\pi = \{(u = u_0, u_1), \dots, (u_{K-1}, u_K = u)\}$ is an Eulerian circuit on G
- If $w \neq u$ is a node in π ,
 - Each time we enter w we subtract 1 from $in(w)$ and also from $out(w)$ when we leave w
 - Since at the end we have passed through all the edges of w , $in(w) = out(w) = 0$
 - Thus, **we must have at the beginning** $in(w) = out(w)$
- Similarly, for u
 - Each time we enter u inside π we subtract 1 from $in(u)$ and also from $out(u)$ when we leave it
 - Moreover, when we start we subtract 1 from $out(u)$ and also subtract 1 we subtract 1 from $in(u)$ when we finish
 - Thus, **we must also have** $in(u) = out(u)$

Euler's Theorem for Directed Graphs

- **Euler's Theorem.** Assume G is a weakly connected directed graph. A necessary and sufficient condition to have an EC in a directed G is that $in(v) = out(v)$ for all $v \in V$
- **Corollary.** A necessary and sufficient condition to have an Eulerian path $\pi = \{(u = u_0, u_1), \dots, (u_{K-1}, u_K = v)\}$ in a directed graph G is that we have $in(w) = out(w)$ for all $w \in V$ different from u and v and also $in(v) = out(v) + 1$, $in(u) = out(u) - 1$
- Essentially the same $O(|E|)$ algorithm we saw for undirected graphs can be applied to directed ones
- Thus we can efficiently sequence genomic reads

Applying Euler on Directed Graphs

- Consider the graph



- The adjacency list and the first exploration give

a	i		way circuits eulerian
1	0	① + 2	all 1 a 5
2	2	2 + 5 - 6	$\pi_1: 1-2-5$
1	1	3 - 4	
1	1	4 - 7	
0	1	⑤ + 7	
1	1	6 - 7	
2	2	7 - 2 - 3	

Applying Euler on Directed Graphs II

- The second and third steps and the final EC are

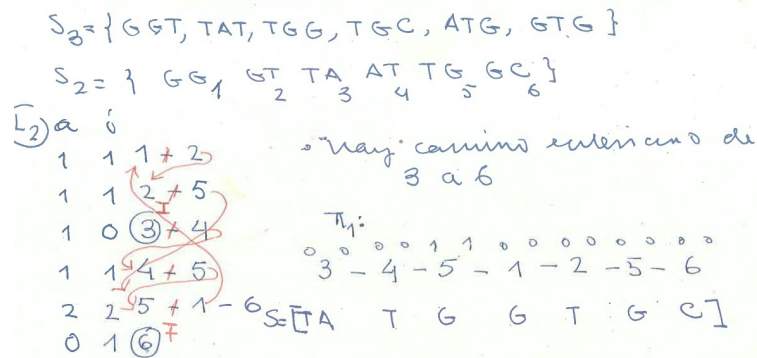
a	i		way circuits eulerian
1	1	2 + 6	$\pi_2: 2-6-7-2$
1	1	3 - 4	
1	1	4 - 7	
1	1	6 + 7	
2	2	7 + 2 - 3	
a	i		way circ. $\pi_3: 7-3-4-7$
1	1	3 + 4	todo junto:
1	1	4 + 7	1-2
1	1	7 + 3	2-5
			6-7
			3-4

Eulerian Sequencing

- Example: consider again $s = [\text{TATGGTGC}]$ and

$$\text{sp}(S, 2) = \{\text{TA}, \text{AT}, \text{TG}, \text{GG}, \text{GT}, \text{GC}\}$$

- Applying the Euler algorithm we obtain



4.2 From Reads to the Genome

The Shortest Superstring Problem (SSP)

- Given a string set $\{s_1, \dots, s_M\}$, find the **shortest superstring** S that contains the M substrings s_j
- Recall that the **overlap** $\omega_{i,j}$ between strings s_i, s_j is the length of the longest suffix of s_i that is also a prefix of s_j
- Notice that the length of the shortest string containing first s_i and then s_j is $|s_i| + |s_j| - \omega_{i,j}$ where $|s|$ denotes the length of s
- If we add another string s_k after s_j , the extra added length is $|s_k| - \omega_{j,k}$
- Thus, if we get S' collating the ordering $\{s_{i_1}, \dots, s_{i_M}\}$ then

$$\begin{aligned} |S'| &= |s_{i_1}| + |s_{i_2}| - \omega_{i_1, i_2} + |s_{i_3}| - \omega_{i_2, i_3} + \dots \\ &= \sum_1^M |s_j| - \sum_1^{M-1} \omega_{i_j, i_{j+1}} \end{aligned}$$

The Longest Path Problem (LPP)

- As an example, if we collate this way the strings
 $'ATGGTAG'$, $'GTAGACTA'$, $'CTAGGTATT'$
 we get the sequence $'ATGGTAGACTAGGTATT'$ of length $17 = 7 + 8 + 9 - 4 - 3$
- Clearly $|S'|$ will be minimal iff $\sum_1^{M-1} \omega_{i_j, i_{j+1}}$ is maximal
- Consider the complete graph G over $V = \{s_1, \dots, s_M\}$ and with cost $\omega_{i,j}$
- Solving the Shortest Superstring Problem is thus equivalent to solving the **Longest Path Problem (LPP)** in (G, ω) : to find a cycle-free path of maximal length

LPP and Hamiltonian Paths

- A Hamiltonian path (HP) is a path that passes over all the $|V|$ vertices once
- It can be shown that HC reduces polynomially to HP (the argument is easy but not totally obvious)
- Consider LPP-d, the decision version of LPP: given a graph G and c , to decide whether there is an acyclic path in G with length $\geq c$
- It is easy to see that LPP-d is in NP
- It is also easy to see that HP reduces to LPP-d: there is a HP in G iff LPP-d over $(G, |V| - 1)$ returns 1
- As a consequence, if HC is NP-complete, so is HP and so is LPP-d
- In general, if P_1 is NP-complete and it reduces polynomially to P_2 , P_2 is also NP-complete

How to Create New NP-complete Problems

- Assume T is the reduction operator from P_1 to P_2 with cost $O(|I|^p)$ when applied to an input I of P_1
- If P' is another problem that reduces polynomially to P_1 via T' , $T \circ T'$ **reduces P' to P_2 with polynomial cost**
- In fact, if $I = T'(I')$ with I' an input of P' with size n and T' has cost $O(n^q)$, we have $|I| = O(n^q)$, for the size of the outputs of T' cannot be larger than the cost of computing them
- But the cost of applying $T \circ T'$ to I' is $O(n^q + (n^q)^p) = O(n^{pq})$
- Thus, if a NP problem reduces polynomially to P_1 , so it does to P_2
- Thus, if P_1 is NP-complete, so is P_2
- Thus, if we manage to prove HC to be NP-complete, so is HP and so is LPP-d. And so is TSP-d

5 Depth First Search and Connectivity

5.1 Depth First Search

Breadth First Search (BFS)

- Recall the general pseudocode for BFS

```
def BFS(u, G):
    s[ ] = False; p[ ] = NULL; Q = pq()
    d[u] = 0; Q.put( (d[u], u) )
    do_something(u)
    while not Q.empty():
        _, v = Q.get()
        if not s[v]:
            s[v] = True
            for all z adjacent to v:
                do_something(z) #perhaps change d[z]
                if d[z] smaller:
                    p[z] = v; Q.put( (d[z], z) )
    return p
```

- If the cost of `do_something` is $O(1)$ and we work with a PQ, the cost of BFS is $O(|E| \log |V|)$ (which can be improved using more sophisticated PQ implementations)
- If we only need simple queue, we get a linear cost $O(|E|)$
- If needed, we add a driver to restart BFS at unseen nodes

Depth First Search (DFS)

- The alternative to BFS is recursive DFS from a starting node u

```
def DFS(u, G):
    s[u] = True
    do_something_before_DFS(u)
    for all w adjacent to u:
        if s[w] == False:
            p[w] = u
            DFS(w, G)
    do_something_after_DFS(u)
```

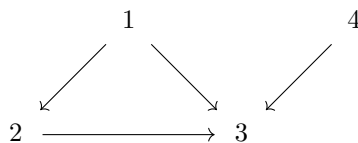
- The table `p[]` defines the DFS tree (or forest)

Depth First Search II

- We may have to restart DFS if not all nodes have been processed, for which we need a driver for DFS

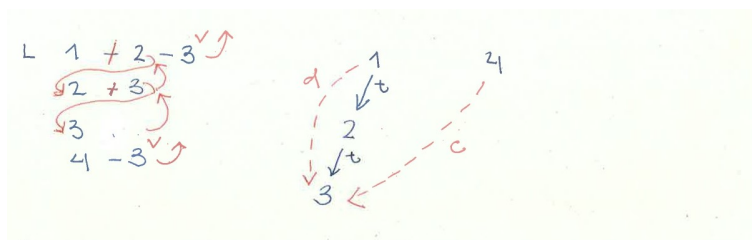
```
def driver_DFS(G):
    s[ ] = False; p[ ] = NULL
    for all u in V:
        if s[u] == False:
            DFS(u, G)
```

- If doing something has cost $O(1)$, the joint cost of `driver_DFS` and `DFS` is clearly $O(|E|)$
- An example:



Applying DFS

- The DFS evolution is



Edge Classification by DFS

- DFS induces a classification on the edges of a directed graph G
 - **Tree edges:** (u, v) where $u = p[v]$
 - **Back (ascending) edges:** (u, v) where $v = p[\dots p[u] \dots]$ (one or more p)
 - **Forward (descending) edges:** (u, v) where $u = p[\dots p[v] \dots]$ (with at least 2 $p[\]$)
 - **Cross edges:** any other $(u, v) \in E$
- If G is undirected and (u, v) is a forward edge, then (v, u) is a back edge
 - Thus, we will not distinguish then between forward and back edges
 - Also we require at least 2 $p[\]$ for back edges
- We prove next that if G is undirected there are no cross edges

Parenthesis Theorem

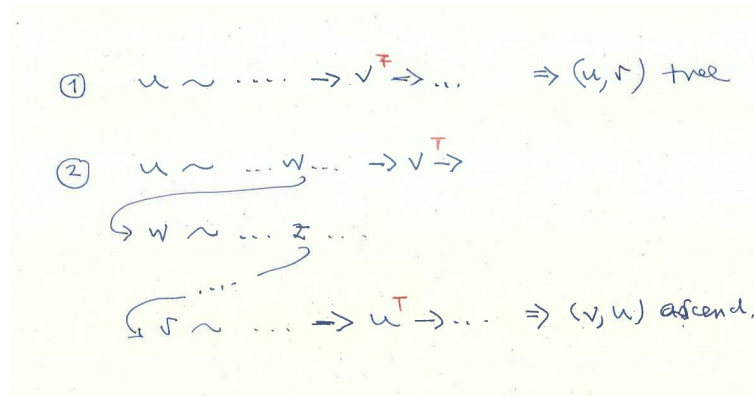
- Assume we have a counter c in DFS and consider 2 time-stamps:
 - **Discovery:** $d[u] = c; c++$, updated when DFS **starts** on u
 - **Finish:** $f[u] = c; c++$, updated when DFS **ends** on u
- Obviously $d_u < f_u$
- **Parenthesis Theorem.** For a graph G and $u, v \in V$, consider the intervals $I_u = (d_u, f_u)$, $I_v = (d_v, f_v)$. Assuming $d_u < d_v$ we either have $I_v \subset I_u$, or $I_u \cap I_v = \emptyset$
- **Proof sketch:** Assume $d_u < d_v$;
 - If $f_u < d_v$, obviously $I_u \cap I_v = \emptyset$
 - And if $f_u > d_v$, DFS started recursively on v before finishing with u ; thus the recursion on v must finish before that of u and $f_v < f_u$
 - Thus, $I_v \subset I_u$

No Cross Edges in Undirected Graphs

- **Corollary.** If G is undirected there are no cross edges
- **Proof sketch:** Take $(u, v) \in E$:
 - Assume $d_u < d_v$; then we have $f_v < f_u$ for v is adjacent to u
 - If $s[v] = F$ when we arrive at v , then (u, v) is a tree edge
 - And if $s[v] = T$ when we arrive at v , we have processed $L[v] \Rightarrow$ we have processed $(v, u) \Rightarrow (v, u)$ must be either a tree or a back edge
 - Thus, (u, v) is either a tree or a forward edge
- Thus, in no case is (u, v) a cross edge

No Cross Edges in Undirected Graphs II

- We sketch the previous arguments.



5.2 Biconnected Graphs

Undirected Graph Connectivity

- Recall that an undirected graph $G = (V, E)$ is connected if **for every pair** $u, v \in V$ **there is a path** π in E **from** u **to** v
- **Connected component**: a maximal connected subgraph of G
- If $G_i = (V_i, E_i)$ are the connected components of G , the V_i are a **partition** of V and the E_i of E
- If we order the vertices of G as $V = V_1 \cup \dots \cup V_K$, then the adjacency matrix M is **block diagonal** with the blocks M_k being the adjacency matrices of the G_k
- DFS can be used to give the connected components of G through the table $p[]$ and restarting DFS as needed
 - BFS and its driver can also be used to give the connected components of G through the table $p[]$

An Aside: Directed Graph Connectivity

- A directed graph $G = (V, E)$ is **weakly connected** if its extension to an undirected graph is connected
- A directed graph $G = (V, E)$ is **strongly connected** if for every pair $u, v \in V$ there is a path π in E from u to v
- DFS is also used in **Tarjan's Algorithm** to obtain the strong components of a graph
- Tarjan's algorithm basically obtains the strong components by
 - Computing DFS's ending times on G and

- Applying again DFS to the transpose graph G^T in the order inverse to the ending times

Articulation Points

- If G is undirected and connected, a **cut vertex** or **articulation point (AP)** is a vertex u such that

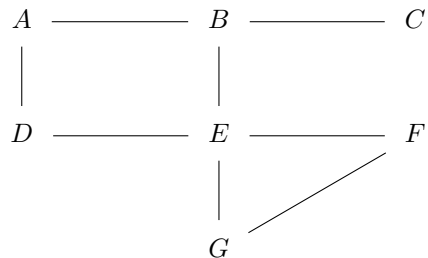
$$G' = (V - \{u\}, E - \{(u, z) \in E\})$$

is no longer connected

- An undirected and connected graph G is **biconnected** if it has no articulation points
- Biconnected graphs are desirable in computer networks, as they are more robust against router failures
- Q: how we detect APs?

How to Detect APs?

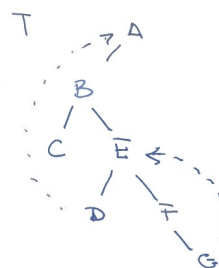
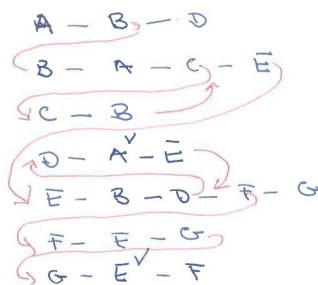
- An example: the graph below has two APs, B and E



- We next apply DFS.

DFS To Detect APs

- We show DFS evolution of the adjacency list and the edges on the DFS tree:



DFS To Detect APs II

- From this “top-down” view of the graph we can more easily detect APs:
 - A is not AP: it does not unhook any vertex as it has just one child
 - B is AP: it unhooks C
 - C is not AP: it has no children
 - E is AP: it unhooks F and G (but not D)
 - D is not AP: it has no children
 - F is not AP: G can reach E without F
 - G is not AP: it has no children it does not unhook any vertex
- The example shows that the DFS tree gives a “top-bottom” view of a graph in which
 - APs other than the root disconnect lower parts of the graph
 - An AP at the root disconnects subtrees

DFS and Articulation Points

- We can use two auxiliary tables that can be computed by DFS to detect articulation points
 - The **order** table $o[\]$ that contains the order in which DFS arrives at a node u .
 - The **ascent** table $a[\]$ which is defined as $a[u] = \min\{o[v]\}$ where v is any node that can be accessed from u by
 - * Going “down” through 0, 1 or more tree edges, and then
 - * Going “up” through a single back edge
- The o, a tables for the previous example are

v	A	B	C	D	E	F	G
o	1	2	3	5	4	6	7
a	1	1	3	1	1	4	4

Detecting Articulation Points

- Clearly if we remove a non root node u from the DFS tree, it will disconnect one of its children v unless v can go “above” $o[u]$ using back edges,
- In other words, u **will be an AP if for some child v we have $o[u] \leq a[v]$**
 - Notice that a larger number means a “lower” node
- Since there are no cross edges on the DFS tree, **the root node will be an AP if it has two or more children**
- It is also clear that these sufficient conditions are also necessary

- A single root node cannot be an AP
- If the ascents of all children of u bypass it, u cannot be an AP

Computing $o[]$ and $a[]$

- We compute $o[u]$ **before** DFS explores u 's adjacency list
- We can use two auxiliary tables to compute the table $a[]$
- The **direct ascent** table $o'[u]$ that contains the order of highest node accessible from u by an ascending edge

$$o'[u] = \min\{o[v] : (v, u) \text{ is a back edge}\}$$

- $o'[u]$ can be computed **before** DFS looking at the w adjacent to u s.t. $s[w] == \text{True}$

- The **ascent by children** table $a'[u]$ that contains the order of highest node accessible from any of the children of u

$$a'[u] = \min\{a[v] : u = p[v]\}$$

- $a'[u]$ can be computed **after** the recursive call to DFS returns
- We then have $a[u] = \min\{o[u], o'[u], a'[u]\}$

The DFS Auxiliary Tables

- The o, o', a', a tables for the previous example are

	o	o'	a'	a
A	1	∞	1	1
B	2	∞	1	1
C	3	1	∞	1
D	5	∞	∞	5
E	4	∞	1	1
F	6	∞	4	4
G	7	4	∞	4

Computing $o[], o'[], a'[]$ and $a[]$

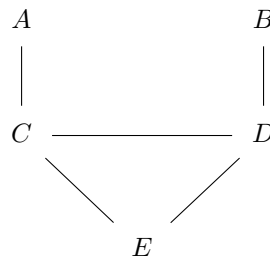
- Assume the DFS driver has initialized $o[]$ and $a[]$ to ∞ and a counter c to 0
- We compute $o[]$ and $a[]$ recursively as follows

```
def ap_tables(u, G):
    s[u] = True; o[u] = c; a[u] = o[u]; c += 1
    for all w adjacent to u: # direct ascent
        if s[w] == True and w != p[u] and o[w] < a[u]:
            a[u] = o[w]
    for all w adjacent to u:
        if s[w] == False:
            p[w] = u; ap_tables(w, G)
    for all w adjacent to u: # ascent by children
        if p[w] == u and a[u] > a[w]:
            a[u] = a[w]
```

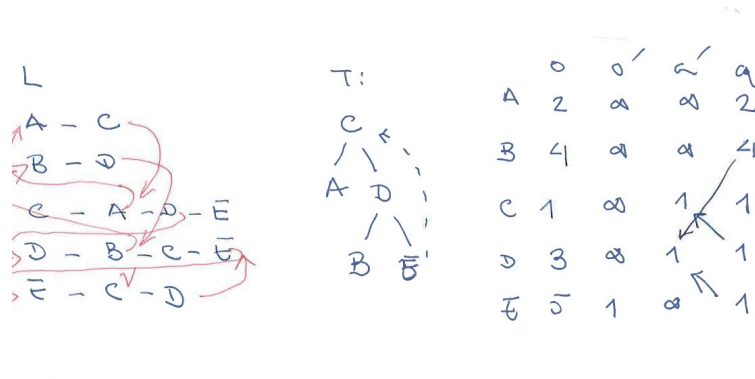
- The cost of `ap_tables` is clearly $O(|E|)$

Algorithm Application

- A second example:

**Algorithm Application II**

- We compute o, o' before DFS and a', a after DFS

**Analyzing the Tables**

- A is not AP: it has no children
- B is not AP: it has no children
- C is AP: root with 2 children
- D is AP: $a[B]4 \geq 3 = o[D]$
- E is not AP: it has no children

5.3 DAGs and Topological Sort

Directed Acyclic Graphs

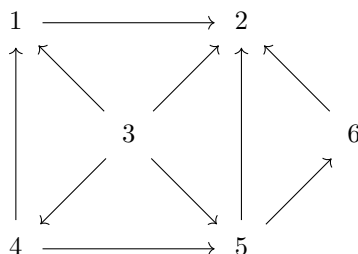
- A **directed acyclic graph** (DAG) is a directed graph without cycles
- **Proposition:** G is a DAG iff there are no ascending edges in G
 - If (v, u) is ascending, there is a path from u to v in the DFS forest, and adding (v, u) results in a cycle
 - Conversely, assume π is a cycle, $u \in V_\pi$ is the first node processed in DFS and $\pi = (u, \dots, v, u)$
 - Then v descends from u in the DFS forest and, thus, (v, u) is ascending
- DFS can be used to detect cycles in a graph by slightly modifying our previous AP algorithm
- DAGs can be used to model many other problems of interest, such as topological node ordering

Topological Sort

- Recall: \leq is a **total order** if either $u \leq v$ or $v \leq u$ or both
- A **topological sort** in a DAG $G = (V, E)$ is any total ordering of its vertices s.t. if $(u, v) \in E$, then $u \leq v$
- If G is a DAG, a topological sort can be obtained
 - Applying DFS starting at u with $inc[u] = 0$ (there is always one) and
 - Adding a vertex u at the beginning of a linked list after DFS ends its process
- We end up with a topological sort of G :
 - Since DFS ended at v **after** all the vertices w adjacent to v have been processed, then these w are in the list **after** v
- The cost of TS on DAGs is thus $O(|E|)$

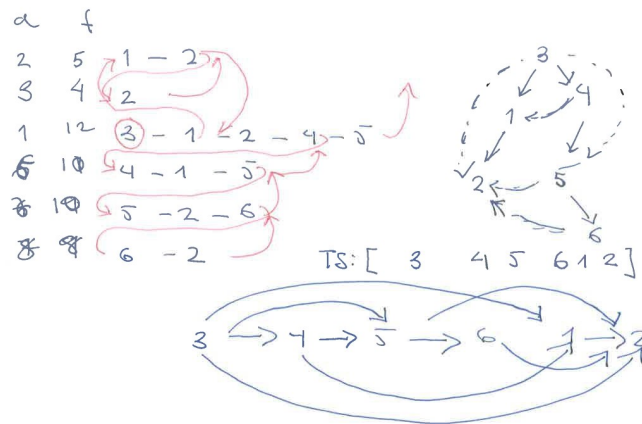
Applying Topological Sort

- An example:



Applying Topological Sort II

- We apply DFS computing the discovery and finish times



- TS can also be obtained by reversed finish times

Applying Topological Sort II

- TS can also be obtained by reversed finish times

