SFWR ENG 2C03 Summary

Author: Kemal Ahmed

Instructor: Dr. Karakostas

Date: Winter 2014

*Math objects made using* [*MathType*](http://www.dessci.com/en/products/mathtype/)*; graphs made using* [*Winplot*](http://math.exeter.edu/rparris/winplot.html)*.*

# Chapter 1

## Algorithm correctness

3 things to prove:

* Initialization
* Maintenance
* Termination

An algorithm is **correct** if its output is correct. This can be measured by looking at the loop invariant.

**Loop invariant**: a property that will be true through all iterations; it proves correctness in algorithms; initialization, maintenance, termination; e.g. Insertion Sort is always sorted on the left side

For a given array, a,

loop invariant: a[1: j − 1] is going to be sorted

# Chapter 2

## Insertion Sort

[Insertion sort](https://upload.wikimedia.org/wikipedia/commons/0/0f/Insertion-sort-example-300px.gif): know the pseudo-code

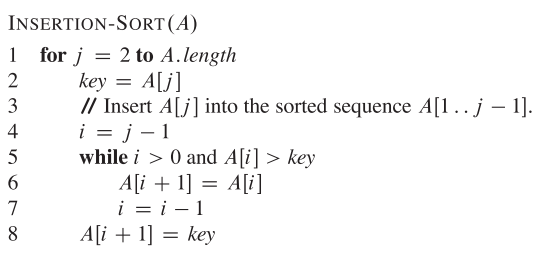
e.g. stack of cards with numbers: 6, 8, 4, 7, 3

j = current card

i = previous card

key = value at current card

Never taking array index of 0



T(n) = an2 + bn + c’

BC = 3 4 6 7 8 O(n)

WC = 8 7 6 4 3 O(n2)

n = # of elements in array

Running times by line:

C1: n

C2: n – 1

C4: n − 1

C5: 

C6: 

C7: 

C8: n – 1

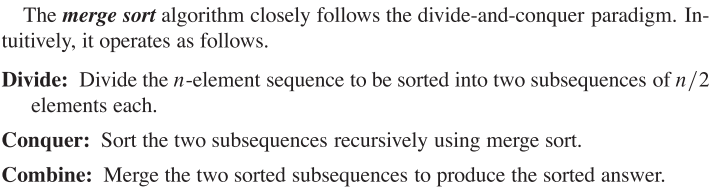
For this course, log2 *x* = log *x* = lg *x*

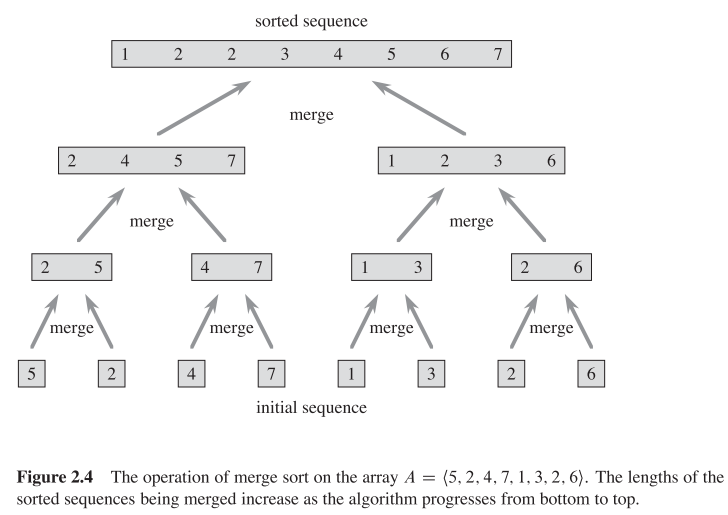
Better than Merge sort for smaller array sizes. It also uses less memory space than merge sort.

## 2.3 − Merge Sort

[Demonstration with dance](http://youtu.be/XaqR3G_NVoo).

[Gif Demonstration](https://upload.wikimedia.org/wikipedia/commons/c/cc/Merge-sort-example-300px.gif).





If the number is odd, split it...

Worst case = best = average = O(nlgn)

Slower than [quicksort](#_Chapter_7:_Quicksort), until a certain *n*, (1billion elements?).

**Auxiliary Space**: space required to run an algorithm

Merge sort is quite memory intensive because it is not an **in-place sort**; merge sort requires space for all its copies of the array, while in-place sorts require space, since they don’t make copies of the array.

Insertion sort is an in-place sort.

## Recurrence Relationships

The general equation for a recurrence relationship is:

* The top line defines when the run time of a function is linear. For example, in merge sort, the run time will always be the same when the size is 2. So *c* is 2.
* *a* is the number of divisions per recurrence
* *b* is the size of the next recursion
* is the time it takes to combine each sub-problem
* is the time it takes to divide each sub-problem

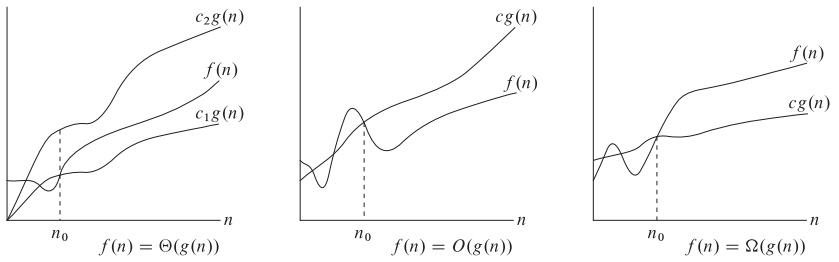
# Chapter 3

## Asymptotic notations

**Asymptotic bounds**: the behaviour of a function as functions get very large and approach infinity

Note: disregard any constants, since they become negligible as functions get very large

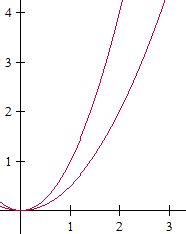
* Big Omicron: O ≤
* Big Theta: Θ =
* Big Omega: Ω ≥
* Small o: o <
* Small omega: ω >



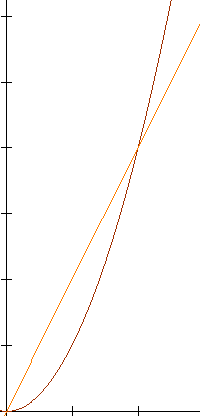


g(n) is going to be much bigger than f(n)

o(n):



2n = O(n2)



for ω(g(n) = f(n),



## Asymptotic Upper Bounds

Page 47: given a function g(n)

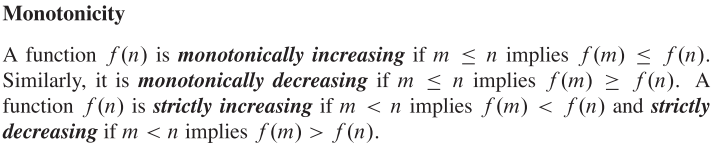




Generally, we focus on the O(n) because it models how the algorithm increases

If you want to find out information about an algorithm, use the Ω

Page 53: **Monotonicity**



Strictly: can only be going up or down

To figure out how long an algorithm takes, find the number of iterations.

If you are given, multiply both sides by n2 to get: (i.e. get rid of fractions)

O(n) is sometimes only true between a given range. For example, f(n) = 3x2 – 5x + 4, O(f(n)) = x2, but only between …(?)

# Chapter 4: Divide and Conquer [the world? no.]

Maximum sub-array problem

## 4.3 - Substitution Method

e.g. Determine upper bound on recurrence: T(n) = 2T() + n (e.g. 4.19 page 83)

T(n) = O(*n*lg*n*)

T(n) ≤ *cn*lg*n*

Assume holds for all positive m < n

m =  = floor(n/2)

T() ≤ c()lg(), sub into the original, since it’s a floor

T(n) ≤ 2(*c*()lg() = *n*

≤ 2*c*()lg() + *n*

≤ *cn*lg() + *n*

≤ , ≤ 0

c ≥ 1

residual ≤ 0

 ≤ 

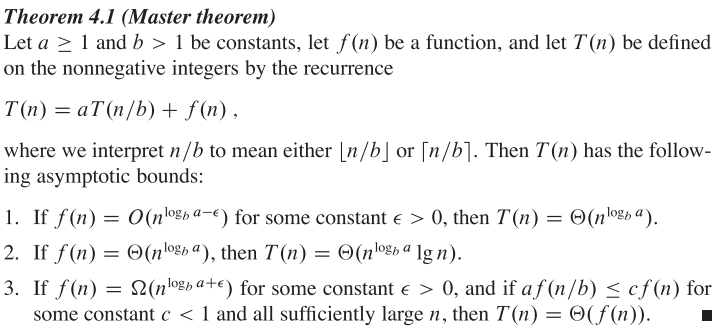
**Root**: node at top level of recursion tree

No parents

Batman

## 4.5 - Master Method

Page 94: **Master theorem**



a = # of sub-problems; n/b = size of the sub-problem; f(n) = base case, most of the time, it’s O(1)

ϵ = used to make the exponent of tn the same as that of f(n). Pg 94 paragraph under the cases

Compare  to f(n) to decide which case you are going to use

* Case 1:  > f(n)
* Case 2:  = f(n)
* Case 3:  < f(n)

### e.g.

T(n) = 9T() + n, a = 9, b = 3, f(n) = n

Choose ϵ = 1



T(n) = Θ(n2)

### e.g.

T(n) = 4T() + , a = 4, b = 2, f(n) = 



Case 3: f(n) = Ω(n2+ϵ), ϵ > 0

f(n) ≥ c ∙ n2+ϵ

 ≥ c ∙ n2+ϵ

e.g. ϵ ≤ ⅓, c = ½

Therefore, T(n) = Θ()

2)

a∙f() ≤ c ∙ f(n)

 ≤ c ∙ 

 ≤ c ∙ 

 ≤ c

e.g. c = 1

### e.g.

T(n) = 2T() + *n*lg*n*, a = 2, b = 2, f(n) = *n*lg*n*



Looks like *n*lg*n* = Ω()

But for any ϵ > 0, = ω(*n*lg*n*)

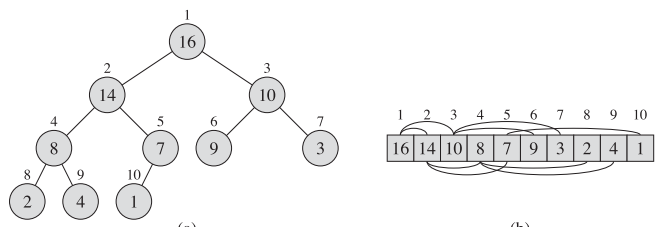
So *n*lg*n* asymptotically larger than n

BUT not *polynomially* larger (between case 2 & 3)

Thus, does not fit in the Master method.

# Chapter 6: Heaps

Page 152:



Parent (i) = 

left(i) = 2i

right(i) = 2i + 1

Useful for priority queues

Root might not be larger than its children.

Max-Heapify puts the root in its place

**Heap**

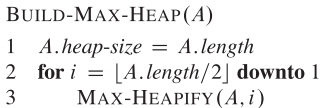
* A.K.A. array tree
* Binary tree in the sense that each node has 0-2 children and 1 parent and an example implementation of a priority queue
* Implements a set, *S*, of elements associated with a key.
* Operations: pick max/min priority, delete, insert, change priorities
* height = lg*n*; min for a heap
* max for a heap is 2h
* root = I, left child = 2i, right child = 2i + 1
* key of a node = S[node]
* Cost: O(*n*lg*n*)

methods:

* insert(S,x) = inserts element, x into set, S
* max(S) = returns value of largest key, usually means highest priority in S
* extract-max(S) = max(S) + deletes it from S
* increasekey(S,x,k) = set key of S(x) to a a high

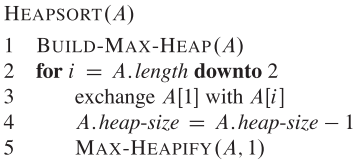
You can only delete the maximum value in a max-heap

Max-heap: key ≥ keys of children; max-heapify(A,i) = sorts heap into a max-heap; run time = O(lg*n*) for each iteration, so O(*n*lg*n*) for the whole tree



Fully-filled in tree has 2n nodes, where n = the level

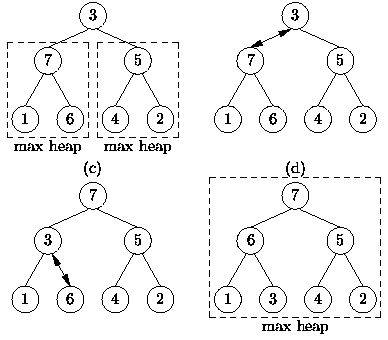
Array, A

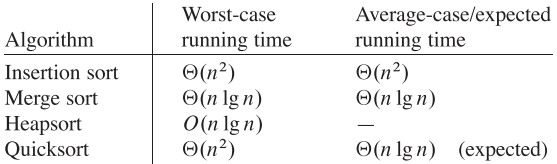


We defined heap size as the array length.

[Example of heap-sort](https://upload.wikimedia.org/wikipedia/commons/4/4d/Heapsort-example.gif).

[Example of Max-heapify](http://www.brpreiss.com/books/opus5/html/page501.html).





# Chapter 7: Quicksort

[Demonstration through a dance](http://youtu.be/ywWBy6J5gz8).

[Quicksort gif](https://upload.wikimedia.org/wikipedia/commons/9/9c/Quicksort-example.gif).

Pointer: address of a particular variable/byte

Pivot point: point in the middle

Best case: splitting into 2 equal sections

To left of pivot is left partition; right, right partition.

You have to change the pivot each layer.

Run time: O(nlgn)

Worst: O(n2)

Faster than [merge sort](#_2.3_−_Merge) below one billion elements.

Balanced?

Worst case is when it’s in decreasing order because the default pivot is always on the right-most element.



# Chapter 10: Elementary Data Structures

**Array**: no book-keeping info

**Stacks**: LIFO (Last-In, First-Out)

Analogy: when you clean your dishes, say, plates, the first plates you’ll pick up when it comes time to use them will be the top one, the most-recently washed one.

Analogy 2: a stack of cups at a fast food restaurant

**Queue**: FIFO (First-in, First Out)

Analogy: line-up

**Head**: First element of a queue

**Tail**: last element in a queue

Stack under/overflow are for stacks & queue’s

Stack underflow: empty stack, but command been sent

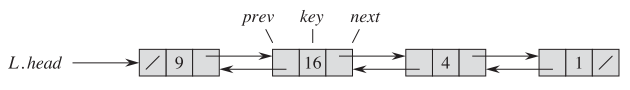
Stack overflow: call a location over the size of the stack

**Linked List**:

Data structure where objects arranged in linear order that is determined by pointer in each object; contain a dummy first and last item, called the sentinel, which indicates the beginning/end when splitting

Linked list vs. doubly-linked list: doubly-linked list has a pointer. What this does

Doubly-linked list (figure 10.3)



Problem 10-1:

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | Unsorted, singly linked | Sorted, singly linked | unsorted, doubly linked | sorted, doubly linked |
| Search(L,k) | O(n)  gotta search thru whole array of elements | O(n) | O(n) | O(n) |
| Insert(L,x) | O(1) | O(1) | O(1) | O(1) |
| Delete(L,x) | O(1)  Can remove element and point to the one after it | 1 | 1 | 1 |
| Successor(L,x) | 1  Just follow the link | 1 | 1 | 1 |
| Predecessor(L,x) | n  There is no link back in singly linked. Must search entire list. | n | 1  There is a link back. | 1 |
| Minimum(L) | n  Isn’t sorted, have to search whole list. | 1  Assumed to be sorted lowest→highest, first element is lowest | n | 1 |
| Maximum(L) | n  Have to search whole list. | n  Know where it is, but have to follow path there. | n  Have to search whole list. | 1  Know where it is, can jump right to it? |

# Chapter 11: Hash Tables

**Hashing**: a.k.a. hash table, hash map; keys are values at given points, e.g. storing songs, key would be song name;

**Chaining**: type of hashing; when the end of the first array is “chained” to the beginning of a another array; pointers necessary to point to the next address; worst-case searching time for hashing with chaining is Θ(n); you need this because you don’t know how many elements you are going to have at each entry

**Double-hashing**: δ(u,v) ≤ δ(s,u) –π

**Direct Access Table**:

# Chapter 12: Binary Search Trees (BSTs)

?

# Chapter 15: Dynamic Programming (DP)

## Dynamic Programming

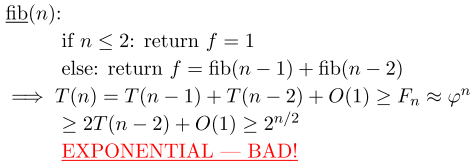
* “careful brute force”
* **Memoize** and reuse solutions to sub-problems that help solve original problems

It’s a way to get away from greedy algorithms. It can be done with bottom-up filling in of the table, but is usually top-down or **memoization**; both have linear time

Focus on the Fibonacci example, instead of the rod-cutting example, which is super confusing

Memoize: remember each step of the recursion

Naive Recursive



Methods:

1) Memoized

memo = {}

fib(n):

if n in memo: return memo[n] ← if key is already in dictionary, return value (do not compute again)

if n ≤ 2: f = 1

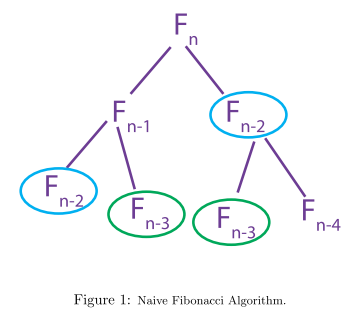
else f = fib(n − 1) + fib (n − 2)

memo[n] = f

return f

* fib(k) only recurses first time it’s called
* memoized calls take Θ(1) (basically free)//I think they would still take Θ(n) because you need to search a memoize table for them, but that’s better than 2n
* # of non-memoized calls is n: fib(1),...,fib(n)
  + non-recursive work per call = Θ(1)

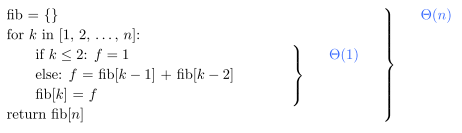
Run-time = (# of subproblems) ∙ (time/subproblem)

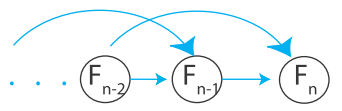


For the memoized algorithm, you’d stop going down on the blue circle on the right, so you wouldn’t even see the right green circle

Same! i.e. already in dictionary. Therefore left continues; right stops

2) Bottom-Up DP

[](http://ocw.mit.edu/courses/electrical-engineering-and-computer-science/6-006-introduction-to-algorithms-fall-2011/lecture-videos/MIT6_006F11_lec19.pdf)

[](http://ocw.mit.edu/courses/electrical-engineering-and-computer-science/6-006-introduction-to-algorithms-fall-2011/lecture-videos/MIT6_006F11_lec19.pdf)

Same as method 1, different wayto think about it, often more efficient (saves space, only need to remember last 2 numbers)

Topological sort of subproblem dependency DAG

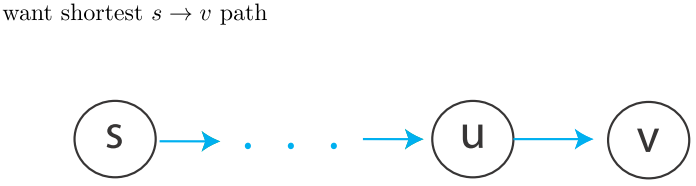
In summary: do basic stuff first that helps you compute the complicated stuff later on

## Shortest Paths

δ(s,n)v

− Guess: try all guesses, pick best one

e.g. shortest path from s to v



* last edge must come from u, try all
* δ(s,n) = min(δ(s,n) + w(u,v)) ← will choose best/shortest path
* takes −∞ time with negative weight cyclic graphs
* ignores a positive weight cycle
* DAG: O(V + E)

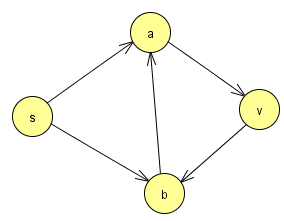
time = indegree(v) + 1 ← in case indegree(V) = 0

total time () + 1 = O(E)

\*Subproblem dependences must be acyclic\*

Why?

e.g. δ(s,s) = 0





## 5 Basic Steps:

1. Define subproblems − # of subproblems
2. Guess part of the solution
3. Relate subproblem solutions with recurrence
4. Build algorithm − check subproblem is acyclic (has topological order)
5. Solve original problem

|  |  |  |
| --- | --- | --- |
| Examples | Fibonacci | Shortest Paths |
| 1) Subproblem:  # of subproblems: | Fk for k = 1,...,n  n | δk(s,v) for vϵV,  0 ≤ k ≤ |v|,  V2 |
| 2) Guess:  # of choices | Nothing  1 | edge info v (if any)  indegree(v) + 1 ← when no edges occur |
| 3) Recurrence:  Time/sub-problem: | Fk = Fk−1 + Fk−2  Θ(1) | δk(s,v) = min{δk−1(s,n)|(u,v) ϵ E}  Θ(indegree(v) + 1) |
| 4) Topological Order:  time: | k = 1,...n  Θ(n) | for k = 0,1,...|v| − 1 for vϵV  Θ(VE) |
| 5) Solve:  extra time: | Fn  Θ(1) | δ|v|−1(s,v)  Θ(v) |

### e.g. Text Justification

split text into “good” lines (align nicely)

text = list of words



→ use words [i, j] as line

|  |  |
| --- | --- |
| 1) Subproblems:  # subproblems: | suffixes words [i,j]  n |
| 2) Guess:  Choices: | where to start 2nd line  ≤ n − i = O(n) |
| 3) Recurrence: | DP[i]  = min(DP(j) + badness(i,j))  time/subproblem = O(n)  for j in range (i + 1, n + 1):   * i is first word of first line * j is first word of second line * DP(n) = 0 |
| 4) Topological order:  Total time: | i = n, n − 1,...0  Θ(n2) |
| 5) Original problem gets solved | DP(0) |

Parent Pointers: remember which choice was best

### e.g.

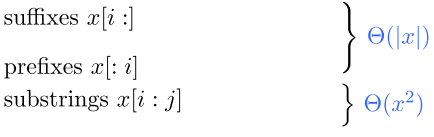
Perfect-Information, Black-jack

* deck = c0, c1,...,cn−1
* 1 player vs dealer
* 1 bet/hand

1. Sub-problems: cards remaining
2. Guess: how many times should I hit per hand?

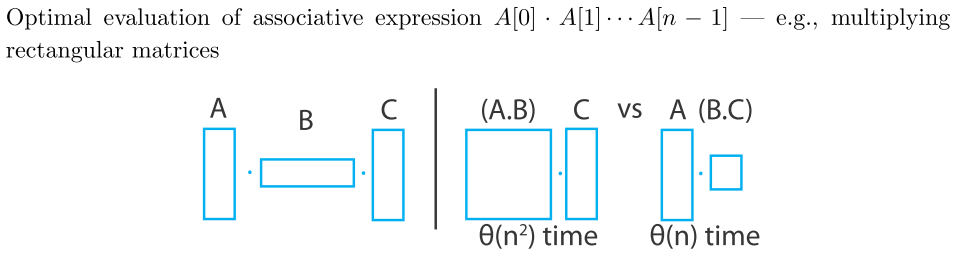
|  |  |
| --- | --- |
| 1) Subproblems:  # subproblems: | Suffix ci  n |
| 2) Guess:  # choices: | How many times hit?  ≤ n |
| 3) Recurrence: | BJ(i) = max(  outcome ϵ {−1,0,1}←$ earned / lost  +BJ(j)→j = [i +4+#hits + #dealer hits]  for #hits in range (0,n)  if valid play) |

### Sub-problems for strings / sequences



Note: never use both prefix + suffix in same sub-problem (use substring instead)

### Parenthesization



Θ(n), therefore linear, better parenthesization

For associative expressions it doesn’t matter where parameters are, but some will be “cheaper” than others. For example, matrix multiplication.

1) Subproblem: optimal evaluation of Ak...Aj−1

# subproblems: n2

2) Guess: outermost / last multiplication

e.g. (A0...Ak−1)∙(Ak...An−1)

prefix suffix

(A0...Ak−1)∙(Ak...Ak−1)

substring

(Aj...Ak−1)∙(Ak...Aj−1)

#choices: O(j − i + 1) = O(n)

3) Recurrence: DP(i,j) = min(DP(i,k) + DP(k,j) + cost(Ai:k)(Ak∙j) for k in range(i+1,j))

time/subproblem: O(n)

4)Total time: Θ(n3)

Topological: Increasing substring size

# Chapter 16: Greedy algorithms

Think of booze. E.g. At a party with free booze. If you want to get drunk fast drink whatever’s in front of you.

Elements of a greedy algorithm:

1. Greedy choices made so far are part of a universal, optimal solution, S (*greedy choice property*).

2. The optimal solution to the sub-problem is part of a universal, optimal solution, S (*optimal substructure*).

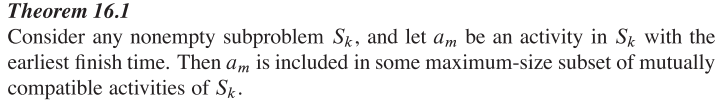
3. Can be a recursive *or* iterative solution

Imagine that you’re robbing a gold vault and trying to fill your backpack with soft gold bars; edge is item, node is state

Fractional Knapsack: you may pick up parts of items, thus you look for the item with the highest money:weight ratio

0-1 Knapsack: only pick the whole items that fit; take it or leave it − no middle way, Buddhist crap; think of it like binary

Greedy: you always pick the most valuable item, the shortest edge, so you’re greedy, like the thief in the gold vault

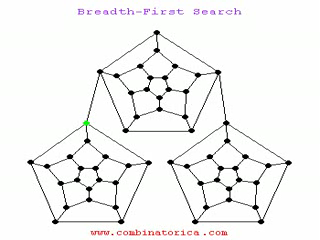


Stupid greedy algorithm: according to your requirements, you go for the higher priority; very dumbed-down example of what a greedy algorithm is; for example, looking for the best money:weight ratio, which isn’t always optimal, but can be

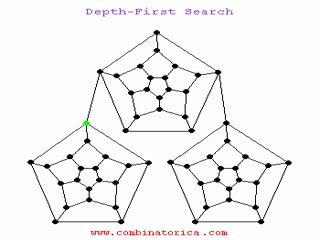
If you have negative weights and you can create a cycle that has a net negative weight (*negative cycle*), you can have a net weight as negative as you want.

# Chapter 22: BFS & DFS

[BFS](https://upload.wikimedia.org/wikipedia/commons/4/46/Animated_BFS.gif) - Breadth first search - undirected graphs



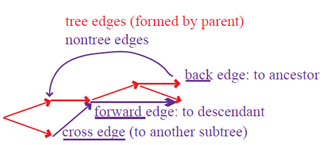
[DFS](https://upload.wikimedia.org/wikipedia/commons/7/7f/Depth-First-Search.gif) - Depth first search - directed graphs

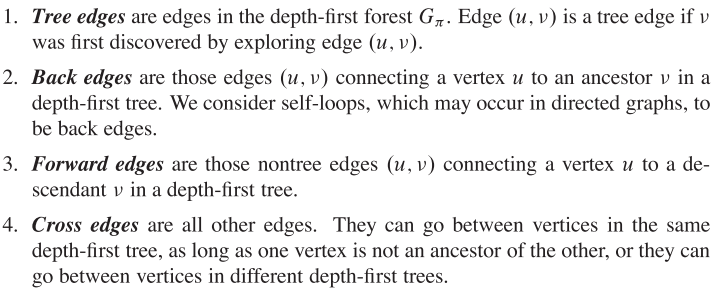


Types of graphs:

-Directed - Can only travel in the direction of the pointer

-Undirected - Can travel through any line any direction





**Acyclic**: no cycles

**Strongly-connected component** (SCC): a part of a directed graph where there exists a path to get from every vertex to every other vertex

**Multi-graph**: graph with multiple edges between a pair of vertices; can have self-loops. There exists both directed and undirected multi-graphs.

# Chapter 23: Minimum Spanning Trees

**Spanning tree**: acyclic; connects all vertices; not necessarily a complete graph

**Complete graph**: everything connected by unique path; all the nodes are directly connected to every other node.



**bottleneck**: heaviest edge in a tree

**Minimum Bottleneck Spanning Tree** (MBST): the tree in a graph with the lowest bottleneck; no cycles; linear time to find an MBST; came up in tutorial

**Safe edge**: when connecting multiple trees, adding it will result in a new forest that is STILL a part/subset of an MST; Invariant, A, is part of a bigger MST

Graph, G(V,E) ← V = set of vertices; E = set of edges

edge(u,v), where u = , and v = vertex

Digraph = directed graph

Prim’s and Kruskal’s Algorithms are greedy algorithms that find a minimum spanning tree for a connected, weighted graph. This means it finds a subset of the edges that forms a tree that includes every vertex, where the total weight of all the edges in the tree is minimized. Both run inif you use binary heaps.

Kruskal’s algorithm: greedy algorithm that finds minimum spanning tree that is both iterative and recursive methods; 2 nodes become 1; choose lightest edge in the whole graph, then next lightest (doesn’t have to be connected as you are working through it as long as it joins up in the end), until all vertices are connected; looks at the smallest weight on the graph and draws a edge, if the nodes have already been “explored” and the edge is useless it skips it, it does this till every node has been “explored”

**Kruskal**

T = Kruskal(G) //Greedy Algorithms

If G = v, then return v,

Find min weight edge, e. (1)

Contract e à G’ = G \ {e} /\*e = weight of the edge. By definition, removing the weight of an edge means removing the edge itself\*/

T’ = Kruskal(G’) (2)

T(n+m) = (1) + T(n−1, m−1)

Return (T’ + e)

Prim’s minimum spanning tree algorithm: a greedy algorithm; choose any starting vertex and look for edge with lowest weight and add it to the tree. Next, choose edge with lowest weight connected to the tree. This tree spans all vertices, so continually repeat this until you’ve covered them all; similar to Dijikstra’s

[Trees] Cut: the splitting of the nodes in a graph into 2 parts to get 2 minimum-spanning trees. If there are any cycles, you can use it to break up the cycles. It is also useful if you know that the item you are looking for is in a specific part of a big tree, so you don’t have to spend as much time looking through the whole tree. After cutting a minimum spanning tree, the lightest edge (weight) is a safe edge; notation for a cut (S, V−S), where S = the connected nodes. A set of edges, A, is a forest, s.t. , T, that satisfies the invariant

**Summary**:

Kruskal's adds one edge at a time. The next line is always the shortest (minimum weight) ONLY if it does NOT create a cycle.

Prim’s adds one vertex at a time. The next vertex to be added is always the one nearest to a vertex already on the graph.

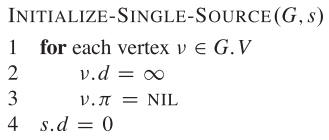
# Chapter 24: Single-Source Shortest Path

Shortest path from s to every other node, but edges are weighted with no loss of generality.

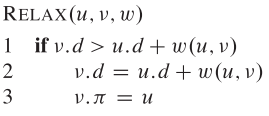
Shortest path weight from u to v: δ(u,v)

Negative weight edges are a problem. None can deal with negative-weighted cycles.

Can a graph have a cycle at all for a shortest path? If no weight, all positive, simple path.

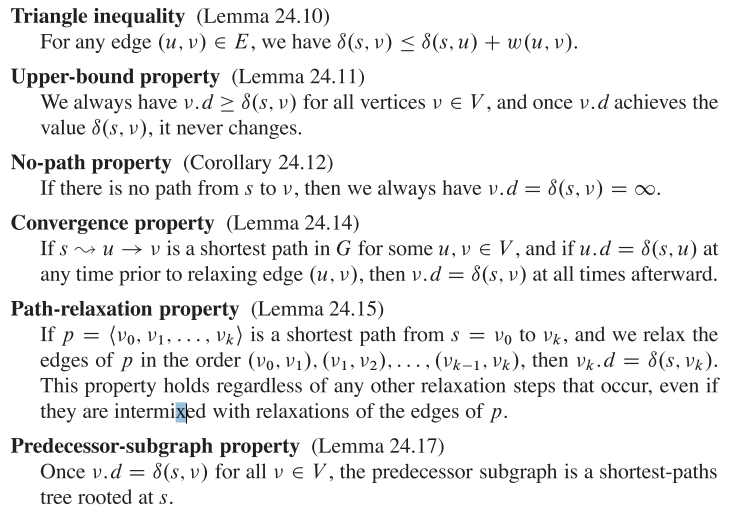


**Relaxation**



Line 2: You realize there’s a shorter path than your estimate

There are 6 properties of shortest paths and relaxation: (page 650)



Predecessor: once all estimates are correct, you have a subgraph that is all shortest paths from s.

## 24.1 Bellman-Ford

More a checker for shortest path than a finder. Also checker for negative cycles.

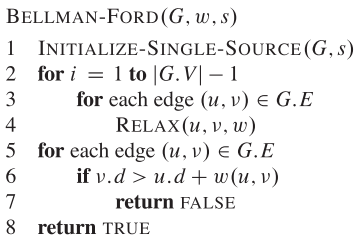
The tree is represented by an adjacency list, which gives us the order to iterate from; s is the first element in the adjacency list.

O(V∙E)

E = O(V2)

Therefore, O(V3)

### Page 651: Bellman-Ford

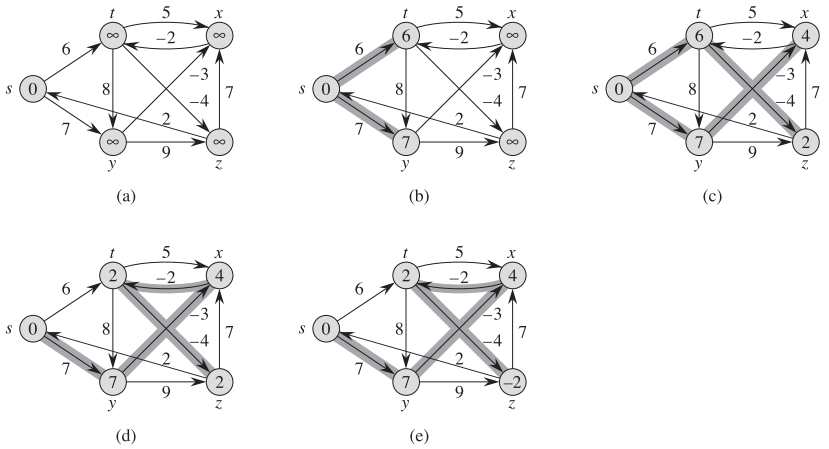


Line 2: iterate V − 1 times (where huge run time comes from)

Line 5-7: Checks if negative weights. If negative weights, then return false.

Once you relax ALL your predecessors, later relaxations WILL result in a shortest path. Relax everything, everything times. Each edge is being visited V−1 times.

### e.g. Page 652



Explanation: It’s brute force and isn’t greedy, so even if you have the most relaxed possible value, you still have to relax it, until the end even though nothing will happen to the edges that are already good. It doesn’t necessarily start where you want it to start. First you do correctness distance 1, then correctness distance 2, etc, like a BFS.

a) Initial

b) First iteration of relaxing edges; correctness level 1

δ(s,t) = 6, but t.d = ∞, so make t.d ≤ 6

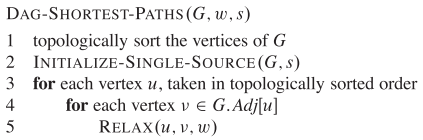
c) Correctness level 2

d) Iterating through all the edges

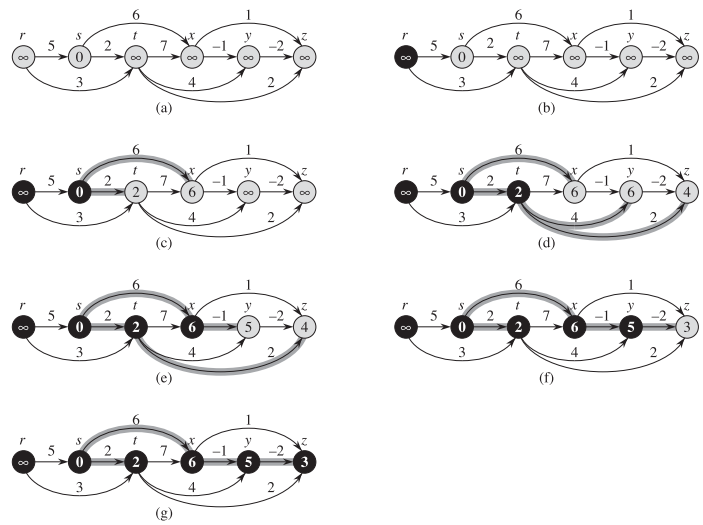
e) Finished!

## 24.2: Single-Shortest path in DAG’s \*woof\*

You know what order because you have sorted, so it is a lot faster, since you don’t ever move backwards.



### e.g. page 656



a) given graph

b) Select single source

c) Relax

d)

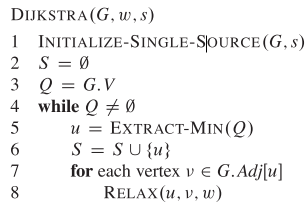
e)

f)

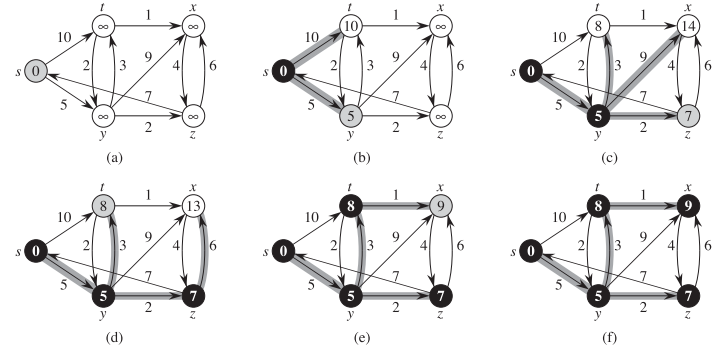
g)

## 24.3: Dijkstra’s

* NO negatively-weighted edges
* Uses min-heaps, probably a binary heap because it uses minimum priority queues
* brute force
* More inituition, than proof
* The top node of your priority queue is always correct at the time of extraction (i.e. taking the top node of the heap)



### e.g. Page 659



Black: node has been extracted

Grey: current node

a) Start!

b) Extract minimum node

c)

d)

e)

f)

# Chapter 25: All-Pairs Shortest Path

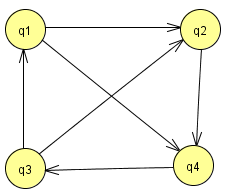
## 25.2: Floyd-Warshall

Floyd-Warshall shortest path algorithm: get from every vertex to every other vertex. There are multiple implementations. It is the bottom-up filling in of the table. There can be no negative cycles (although negative edges are OK). The predecessor matrix is Π. The weight matrix is D.

O(V3)

### Page 694:

https://lh6.googleusercontent.com/AlDo0AJ9L2bycdsX3NK2Dap9Auz8mt9w9VG1bEMYYDeXJb1v0VXsYlGz89jtclbhJm8pGPWTfOwgfXPkIyQFGcPxJjlnnPjlHxHapKxKWJAzFDmOe_BeTtG_





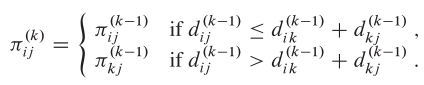
Since no negative edges, all edges are carried over to the new one

When k = 3, you may use the vertices numbered 1-3. So in the D(1), we are only allowed going through vertex number 1.

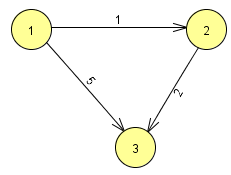
Transitive closure between a and b means there is path between the two vertices

Transitive relationship: whenever A > B and B > C, then also A > C

https://lh4.googleusercontent.com/MgeTgqLlA5YOTKV_yPDE73mrEbbOfGDYQ-9yulNluKU92YZfXGshLfQ2fVcNo-UDR5MfJNIrQ540NBMRw6b5F8JV3T2TCzjnITdcM8fNBxjZWfkbN-AwldBR



If the new path is shorter, then change it. If not, don’t.



All 0’s and ∞’s are NIL. Let’s represent NIL by just an “N”.

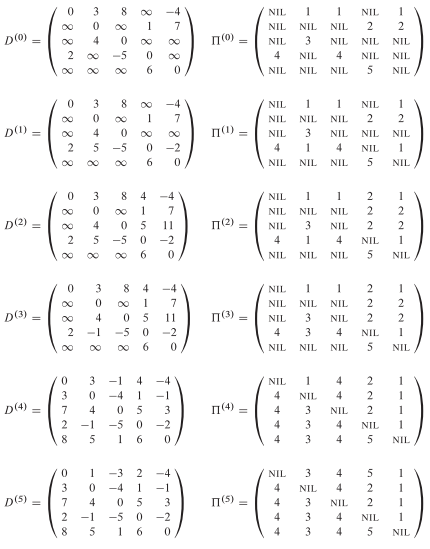


The weight diagonal is always 0, i.e. 

The predecessor diagonal is always NIL, i.e. 

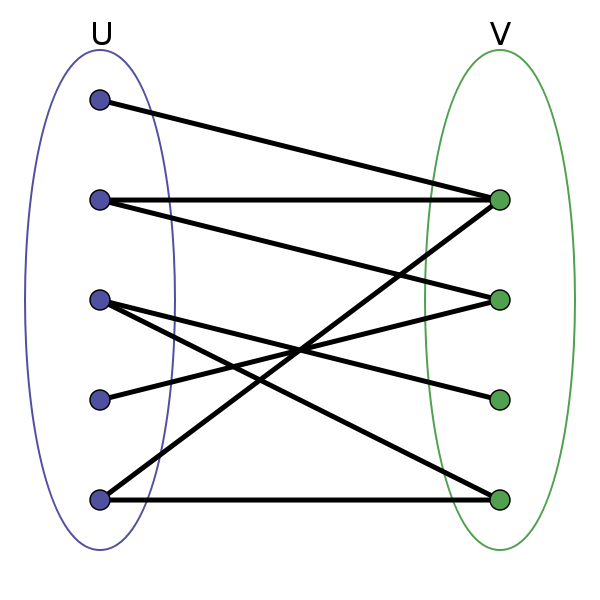


### e.g. 2 Page 696

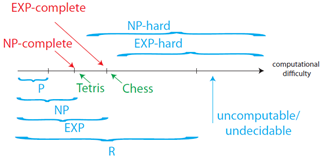


### e.g. 3)

Bipartite graph: 1-1 function

[](https://en.wikipedia.org/wiki/Bipartite_graph)

# Chapter 34: NP-Completeness

[](http://ocw.mit.edu/courses/electrical-engineering-and-computer-science/6-006-introduction-to-algorithms-fall-2011/lecture-videos/MIT6_006F11_lec23.pdf)

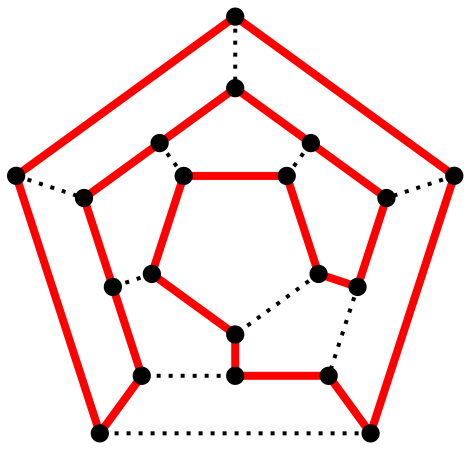
Languages in NP-hard are not verifiable in polynomial time, but are reducible.

Complexity Class P: problems solvable in polynomial time (run-time = O(nk))

P = 

Complexity Class NP: {x  {0,1}\*: there exists a certificate, y, |y| = 0 (|x|c), s.t. A(x,y) = 1}

e.g. Hamiltonian cycle: cycle covering all vertices

[](https://en.wikipedia.org/wiki/Hamiltonian_path)

Verification Algorithm: L = {xϵ{0,1}\*, there exists yϵ{0,1}\*, s.t. A(x,y) = 1

A language (think in terms of SE 2FA3), L  {0,1}\* NPC:f:

1) L ϵ NP

2) L’≤p L for every L’ϵNP

Polynomial reducibility:

ax + b = 0

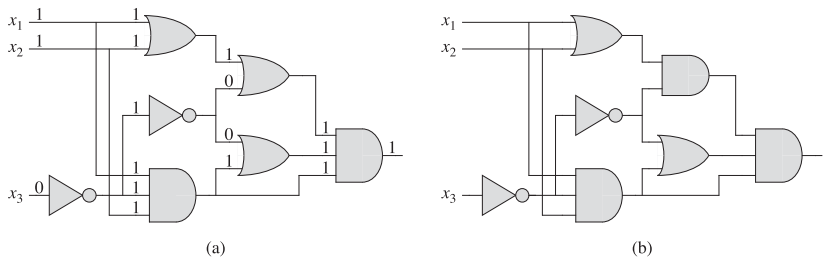
ax2 + ax + b = 0

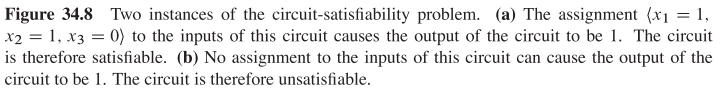


P = NP? P ≠ NP? Is NP is solvable in polynomial time?

NPC: NP-complete

examples of NP problems: circuit satisfiability (page 1070)





**Proving NPC**:

Find an alg, A, that can verify L in polynomial time.

NPC ≤p P

**Proof [problem-specific]**: (Page 1078)

Prove L ϵ NP

Select a known NPC-complete language, L’.

Find a poly-time reduction function, f, that reduces L’ to L. algorithm, A, that verifies L in polynomial time.

length (x1,...xn) => O(n)

e.g. Formula satisfiability (page 1079)

SAT ≤p P

1) Circuit-sat is NPC

circuit {x ϵ {0,1}\*, L satisfies the circuit

SAT{ϕ}: ϕ is satisfiable}

Page 1081

NP-Hard

Page 1069:

L ϵ NP

L’ ≤p P for every L’ ϵ NP

Co-NP: set of languages whose complement is in NP; it says yes whenever the NP checker says no.

Circle = language, red = co-NP, yellow = NP, orange = both

P = co-NP, P is in orange

MIPS - Microprocessor without Interlocked Pipeline Stages

Clique problem: finding particular complete sub-graphs ("cliques") in a graph, i.e., sets of elements where each pair of elements is connected.

Appendix B:

In a given digraph, an edge (u,v) is incident to or enters v and is incident from u.

Midterm:

Part I:

1)

3)

Part II:

2b)

23

15 17

14 12,13 11

9 7,4 18

Switch 12 and 18

Switch 18 and 15

2c) Make a new version of BUILD-MAX-HEAP

Recursion tree:

Height = logn

Width = n

Divide: RECURSIVE-BUILD(A) split into 2 sub-problems (first/last half of A)

Conquer: Recursion of RECURSIVE-BUILD until size 1

Combine: MAX-HEAPIFY between the sub-problems, height

height ,sub-problems → Total runtime: Θ()

3)

Part V:

Input: Adjacency-list

G = (V,E)

Output: starting soon

O(V + E)

Size of input: # one-way doors + 2 × # 2-way doors + # rooms

Use strongly-connected components

Topologically sort it

DFS on the very first node