DESIGN AND ANALYSIS OF ALGORITHMS (Common to CSE & ISE)

Subject Code: 10CS43 Hours/Week: 04 **Total Hours : 52**

PART – A

UNIT – 1

INTRODUCTION: Notion of Algorithm, Review of Asymptotic Notations, Mathematical Analysis of Non-Recursive and Recursive AlgorithmsBrute Force Approaches: Introduction, Selection Sort and Bubble Sort, Sequential Search and Brute Force String Matching. **6** Hours

UNIT - 2

DIVIDE AND CONQUER: Divide and Conquer: General Method, Defective Chess Board, Binary Search, Merge Sort, Quick Sort and its performance.

UNIT - 3

THE GREEDY METHOD: The General Method, Knapsack Problem, Job Sequencing with Deadlines, Minimum-Cost Spanning Trees: Prim's Algorithm, Kruskal's Algorithm; Single Source Shortest Paths. **UNIT - 4**

DYNAMIC PROGRAMMING: The General Method, Warshall's Algorithm, Floyd's Algorithm for the All-Pairs Shortest Paths Problem, Single-Source Shortest Paths: General Weights, 0/1 Knapsack, The Traveling Salesperson problem.

PART – B

UNIT - 5

DECREASE-AND-CONQUER APPROACHES, SPACE-TIMETRADEOFFS:

Decrease-and-Conquer Approaches: Introduction, Insertion Sort, Depth First Search and Breadth First Search, Topological Sorting Space-Time Tradeoffs: Introduction, Sorting by Counting, Input Enhancement in String Matching.

UNIT - 6

LIMITATIONS OF ALGORITHMIC POWER AND COPING WITH THEM:

Lower-Bound Arguments, Decision Trees, P, NP, and NP-Complete Problems, Challenges of Numerical Algorithms.

UNIT - 7

COPING WITH LIMITATIONS OF ALGORITHMIC POWER:

Backtracking: n - Queens problem, Hamiltonian Circuit Problem, Subset -Sum Problem.Branch-and-Bound: Assignment Problem, Knapsack Problem, Traveling Salesperson Problem. Approximation Algorithms for NP-Hard Problems - Traveling Salesperson Problem, Knapsack Problem

UNIT - 8

PRAM ALGORITHMS: Introduction, Computational Model, Parallel Algorithms for Prefix Computation, List Ranking, and Graph Problems,

Text Books:

1. Anany Levitin: Introduction to The Design & Analysis of Algorithms, 2nd Edition, Pearson Education, 2007. (Listed topics only from the Chapters 1, 2, 3, 5, 7, 8, 10, 11).

2. Ellis Horowitz, Sartaj Sahni, Sanguthevar Rajasekaran: Fundamentals of Computer Algorithms, 2nd Edition, UniversitiesPress, 2007.(Listed topics only from the Chapters 3, 4, 5, 13)

Reference Books:

1. Thomas H. Cormen, Charles E. Leiserson, Ronal L. Rivest, Clifford Stein: Introduction to Algorithms, 3rd Edition, PHI. 2010.

2. R.C.T. Lee, S.S. Tseng, R.C. Chang & Y.T.Tsai: Introduction to the Design and Analysis of Algorithms A Strategic Approach, Tata McGraw Hill, 2005.

6 Hours

7 Hours

I.A. Marks : 25

Exam Hours: 03

Exam Marks: 100

7 Hours

6 Hours

7 Hours

6 Hours

7 Hours

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UNIT – 1

INTRODUCTION

1.1 Notion of Algorithm

- 1.2 Review of Asymptotic Notation
- 1.3 Mathematical Analysis of Non-Recursive and Recursive Algorithms
- 1.4 Brute Force Approaches: Introduction

1.5 Selection Sort and Bubble Sort

1.6 Sequential Search and Brute Force String Matching.

1.1 Notion of Algorithm

Need for studying algorithms:

The study of algorithms is the cornerstone of computer science. It can be recognized as the core of computer science. Computer programs would not exist without algorithms. With computers becoming an essential part of our professional & personal life's, studying algorithms becomes a necessity, more so for computer science engineers. Another reason for studying algorithms is that if we know a standard set of important algorithms, They further our analytical skills & help us in developing new algorithms for required applications

Algorithm

An algorithm is finite set of instructions that is followed, accomplishes a particular task. In addition, all algorithms must satisfy the following criteria:

- 1. Input. Zero or more quantities are externally supplied.
- 2. Output. At least one quantity is produced.
- 3. Definiteness. Each instruction is clear and produced.
- 4. Finiteness. If we trace out the instruction of an algorithm, then for all cases, the algorithm terminates after a finite number of steps.
- 5. Effectiveness. Every instruction must be very basic so that it can be carried out, in principal, by a person using only pencil and paper. It is not enough that each operation be definite as in criterion 3; it also must be feasible.



An algorithm is composed of a finite set of steps, each of which may require one or more operations. The possibility of a computer carrying out these operations necessitates that certain constraints be placed on the type of operations an algorithm can include. The fourth criterion for algorithms we assume in this book is that they terminate after a finite number of operations.

Criterion 5 requires that each operation be effective; each step must be such that it can, at least in principal, be done by a person using pencil and paper in a finite amount of time. Performing arithmetic on integers is an example of effective operation, but arithmetic with real numbers is not, since some values may be expressible only by infinitely long decimal expansion. Adding two such numbers would violet the effectiveness property.

- Algorithms that are definite and effective are also called computational procedures.
- The same algorithm can be represented in same algorithm can be represented in several ways
- Several algorithms to solve the same problem
- Different ideas different speed

Example:

Problem:GCD of Two numbers m,n

Input specifiastion :Two inputs, nonnegative, not both zero

Euclids algorithm

 $-gcd(m,n)=gcd(n,m \mod n)$

Untill m mod n =0,since gcd(m,0) =m

Another way of representation of the same algorithm

Euclids algorithm

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Step1:if n=0 return val of m & stop else proceed step 2

Step 2:Divide m by n & assign the value of remainder to r

Step 3: Assign the value of n to m,r to n,Go to step 1.

Another algorithm to solve the same problem

Euclids algorithm

Step1:Assign the value of min(m,n) to t Step 2:Divide m by t.if remainder is 0,go to step3 else goto step4

Step 3: Divide n by t.if the remainder is 0, return the value of t as the answer and

stop, otherwise proceed to step4

Step4 :Decrease the value of t by 1. go to step 2

1.1 Review of Asymptotic Notation

Fundamentals of the analysis of algorithm efficiency

- Analysis of algorithms means to investigate an algorithm's efficiency with respect to resources:
- running time (time efficiency)
- memory space (space efficiency)

Time being more critical than space, we concentrate on Time efficiency of algorithms. The theory developed, holds good for space complexity also.

Experimental Studies: requires writing a program implementing the algorithm and running the program with inputs of varying size and composition. It uses a function, like the built-in clock() function, to get an accurate measure of the actual running time, then analysis is done by plotting the results.

Limitations of Experiments

- It is necessary to implement the algorithm, which may be difficult
- Results may not be indicative of the running time on other inputs not included in the experiment.
- In order to compare two algorithms, the same hardware and software environments must be used

Theoretical Analysis: It uses a high-level description of the algorithm instead of an implementation. Analysis characterizes running time as a function of the input size, n, and takes into account all possible inputs. This allows us to evaluate the speed of an

algorithm independent of the hardware/software environment. Therefore theoretical analysis can be used for analyzing any algorithm

Framework for Analysis

We use a hypothetical model with following assumptions

- Total time taken by the algorithm is given as a function on its input size
- Logical units are identified as one step
- Every step require ONE unit of time
- Total time taken = Total Num. of steps executed

Input's size: Time required by an algorithm is proportional to size of the problem

instance. For e.g., more time is required to sort 20 elements than what is required to sort 10 elements.

Units for Measuring Running Time: Count the number of times an algorithm's **basic operation** is executed. (**Basic operation**: The most important operation of the algorithm, the operation contributing the most to the total running time.) For e.g., The basic operation is usually the most time-consuming operation in the algorithm's innermost loop.

Consider the following example:

ALGORITHM sum_of_numbers (A[0... n-1])

// Functionality : Finds the Sum
// Input : Array of n numbers
// Output : Sum of 'n' numbers
i 0
sum 0
while i < n
sum sum + A[i] n
i i + 1
return sum</pre>

Total number of steps for basic operation execution, C (n) = n

NOTE:

Constant of fastest growing term is insignificant: Complexity theory is an Approximation theory. We are not interested in exact time required by an algorithm to solve the problem. Rather we are interested in order of growth. i.e How much faster will algorithm run on computer that is twice as

fast? How much longer does it take to solve problem of double input size? We can crudely estimate running time by

 $T(n) \approx Cop \clubsuit C(n)$

Where,

T (n): running time as a function of n.

Cop : running time of a single operation.

C (n): number of basic operations as a function of n.

Order of Growth: For order of growth, consider only the leading term of a formula and ignore the constant coefficient. The following is the table of values of several functions important for analysis of algorithms.

| n | $\log_2 n$ | n | $n \log_2 n$ | n^2 | n^3 | 2^n | n! |
|----------|------------|----------|--------------------|-----------|-----------|---------------------|----------------------|
| 10 | 3.3 | 101 | $3.3 \cdot 10^{1}$ | 10^{2} | 10^{3} | 10^{3} | $3.6 \cdot 10^{6}$ |
| 10^2 | 6.6 | 10^{2} | $6.6 \cdot 10^{2}$ | 10^{4} | 10^{6} | $1.3 \cdot 10^{30}$ | $9.3 \cdot 10^{157}$ |
| 10^{3} | 10 | 10^{3} | $1.0.10^{4}$ | 10^{6} | 10^{9} | | |
| 10^{4} | 13 | 10^{4} | $1.3 \cdot 10^{5}$ | 10^{8} | 10^{12} | | |
| 10^{5} | 17 | 10^{5} | $1.7 \cdot 10^{6}$ | 1010 | 1015 | | |
| 10^{6} | 20 | 10^{6} | $2.0 \cdot 10^{7}$ | 10^{12} | 10^{18} | | |

Worst-case, Best-case, Average case efficiencies

Algorithm efficiency depends on the **input size n.** And for some algorithms efficiency depends on **type of input**. We have best, worst & average case efficiencies.

- Worst-case efficiency: Efficiency (number of times the basic operation will be executed) for the worst case input of size n. *i.e.* The algorithm runs the longest among all possible inputs of size n.
- **Best-case efficiency:** Efficiency (number of times the basic operation will be executed) for the **best case input of size n.** *i.e.* The algorithm runs the fastest among all possible inputs of size n.
- Average-case efficiency: Average time taken (number of times the basic operation will be executed) to solve all the possible instances (random) of the input. NOTE: NOT the average of worst and best case

Asymptotic Notations

Asymptotic notation is a way of comparing functions that ignores constant factors and small input sizes. Three notations used to compare orders of growth of an algorithm's basic operation count are:

O, Ω , Θ notations

Big Oh- O notation

Definition:

A function t(n) is said to be in O(g(n)), denoted t(n) O(g(n)), if t(n) is bounded above by some constant multiple of g(n) for all large n, i.e., if there exist some positive constant c and some nonnegative integer n0 such that



Big-oh notation: $t(n) \in O(g(n))$

Big Omega- Ω notation

Definition:

A function t (n) is said to be in Ω (g(n)), denoted t(n) $\diamondsuit \Omega$ (g (n)), if t (n) is bounded below by some constant multiple of g (n) for all large n, i.e., if there exist some positive constant c and some nonnegative integer n0 such that

 $t(n) \geq cg(n) \text{ for all } n \geq n0$



Big-omega notation: $t(n) \in \Omega(g(n))$

Big Theta- Θ notation

Definition:

A function t (n) is said to be in $\Theta(g(n))$, denoted t(n) $\mathbf{O}(g(n))$, if t (n) is bounded both above and below by some constant multiple of g (n) for all large n, i.e., if there exist some positive constant c1 and c2 and some nonnegative integer n0 such that



Big-theta notation: $t(n) \in \Theta(g(n))$

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Basic Efficiency classes

The time efficiencies of a large number of algorithms fall into only a few classes.

| £ | constant | High time efficiency |
|------|----------------------------|----------------------|
| iast | Constant | |
| | log <i>n</i> logarithmic | |
| | n linear | |
| | $n \log n$ $n \log n$ | |
| | n ² quadratic | |
| | n ³ cubic | |
| | 2 ⁿ exponential | |
| | n! factorial | |
| alow | | low time efficiency |
| SIOW | $\mathbf{\nabla}$ | |

1.2 Mathematical Analysis of Non-Recursive and Recursive Algorithms

Mathematical analysis (Time Efficiency) of Non-recursive Algorithms

General plan for analyzing efficiency of non-recursive algorithms:

- 1. Decide on parameter n indicating **input size**
- 2. Identify algorithm's basic operation
- Check whether the number of times the basic operation is executed depends only on the input size n. If it also depends on the type of input, investigate worst, average, and best case efficiency separately.
- 4. Set up **summation** for C(n) reflecting the number of times the algorithm's basic operation is executed.
 - 5. Simplify summation using standard formulas

Example: Finding the largest element in a given array

ALOGORITHM MaxElement(A[0..n-1])

//Determines the value of largest element in a given array

//nput: An array A[0..n-1] of real numbers

//Output: The value of the largest element in A

currentMax $\leftarrow A[0]$

for $i \leftarrow 1$ to n - 1 do

currentMax $\leftarrow A[i]$

return currentMax

Analysis:

- 1. Input size: number of elements = n (size of the array)
- 2. Basic operation:
- a) Comparison
- b) Assignment
- 3. NO best, worst, average cases.

Let C (*n*) denotes number of comparisons: Algorithm makes one comparison on each execution of the loop, which is repeated for each value of the loop's variable i within the bound between 1 and n - 1.

$$C(n) = \sum_{i=1}^{n-1} 1$$

5. Simplify summation using standard formulas

 $C(n) = \sum_{i=1}^{n-1} 1 \qquad \begin{array}{l} 1+1+1+\ldots+1\\ [(n-1) \text{ number of times}] \end{array}$ C(n) = n-1 $C(n) \square \Theta(n)$

Example: Element uniqueness problem

Algorithm UniqueElements (A[0..n-1])

//Checks whether all the elements in a given array are distinct

//Input: An array A[0..n-1]

//Output: Returns true if all the elements in A are distinct and false otherwise

for i 0 to n - 2 do

for j = i + 1 to n - 1 do

if A[i] = = A[j]

return *false*

return *true*

Analysis

- 1. Input size: number of elements = n (size of the array)
- 2. Basic operation: Comparison
- 3. Best, worst, average cases EXISTS.

Worst case input is an array giving largest comparisons.

- Array with no equal elements
- Array with last two elements are the only pair of equal elements
- 4. Let C (n) denotes number of comparisons in worst case: Algorithm makes one

comparison for each repetition of the innermost loop i.e., for each value of the

loop's variable j between its limits i + 1 and n - 1; and this is repeated for each value of the outer loop i.e, for each value of the loop's variable i between its limits 0 and n - 2

C (n) =
$$\sum_{i=0}^{n-2} \sum_{j=i+1}^{n-1} 1$$

5. Simplify summation using standard formulas

$$C(n) = \sum_{i=0}^{n-2} ((n-1) - (i+1) + 1)$$

$$C(n) = \sum_{i=0}^{n-2} (n-1 - i)$$

$$C(n) = \sum_{i=0}^{n-2} (n-1) - \sum_{i=0}^{n-2} i$$

$$C(n) = (n-1) \sum_{i=0}^{n-2} 1 - \sum_{i=0}^{n-2} i$$

$$C(n) = (n-1) \sum_{i=0}^{n-2} 1 - \frac{(n-2)(n-1)}{2}$$

$$C(n) = (n-1)(n-1) - \frac{(n-2)(n-1)}{2}$$

$$C(n) = (n-1)((n-1) - \frac{(n-2)}{2})$$

$$C(n) = (n-1)(2n-2-n+2)$$

$$C(n) = (n-1)(n)/2$$

$$= (n^2 - n)/2$$

$$= (n^2/2 - n/2)$$

$$C(n) \Box \Theta(n^2)$$

Mathematical analysis (Time Efficiency) of recursive Algorithms

General plan for analyzing efficiency of recursive algorithms:

1. Decide on parameter n indicating input size

2. Identify algorithm's basic operation

3. Check whether the number of times the basic operation is executed depends only

on the input size n. If it also depends on the type of input, investigate worst,

average, and best case efficiency separately.

4. Set up **recurrence relation**, with an appropriate initial condition, for the number of times the algorithm's basic operation is executed.

5. Solve the recurrence.

Example: Factorial function

ALGORITHM Factorial (n)

//Computes n! recursively
//Input: A nonnegative integer n
//Output: The value of n!
if n = = 0

return 1 else **return** Factorial (n-1) * n**Analysis:** 1. Input size: given number = n2. Basic operation: multiplication 3. NO best, worst, average cases. 4. Let M (n) denotes number of multiplications. M(n) = M(n-1) + 1for n > 0M(0) = 0initial condition Where: M (n-1): to compute Factorial (n-1)1 :to multiply Factorial (n - 1) by n5. Solve the recurrence: Solving using "Backward substitution method": M (*n*) = M (n - 1) + 1 = [M(n-2) + 1] + 1= M (n - 2) + 2 = [M(n-3) + 1] + 3= M (n - 3) + 3. . . In the ith recursion, we have $= \mathbf{M} (n-i) + i$ When i = n, we have = M (n - n) + n = M (0) + nSince M(0) = 0= *n* $\mathbf{M}(n) \diamondsuit \Theta(\mathbf{n})$

Example: Find the number of binary digits in the binary representation of a positive

decimal integer

ALGORITHM BinRec (n) //Input: A positive decimal integer n //Output: The number of binary digits in n's binary representation if n = = 1return 1

else

return $BinRec (\lfloor n/2 \rfloor) + 1$

Analysis:

- 1. Input size: given number = n
- 2. Basic operation: addition
- 3. NO best, worst, average cases.
- 4. Let A (*n*) denotes number of additions.

A $(n) = A (\lfloor n/2 \rfloor) + 1$ for n > 1

A (1) = 0 initial condition

Where: A ($\lfloor n/2 \rfloor$) : to compute *BinRec* ($\lfloor n/2 \rfloor$)

1 : to increase the returned value by 1

5. Solve the recurrence:

A (*n*) = A ($\lfloor n/2 \rfloor$) + 1 for *n* > 1

Assume $n = 2^k$ (smoothness rule)

A $(2^k) = A(2^{k-1}) + 1$ for k > 0; A (20) = 0

Solving using "Backward substitution method":

A
$$(2^{k}) = A (2^{k-1}) + 1$$

= $[A (2^{k-2}) + 1] + 1$
= $A (2^{k-2}) + 2$
= $[A (2^{k-3}) + 1] + 2$
= $A (2^{k-3}) + 3$
...
In the ith recursion, we have
= $A (2^{k-i}) + i$

When i = k, we have $= A (2^{k-k}) + k = A (2^{0}) + k$ Since $A (2^{0}) = 0$ $A (2^{k}) = k$ Since $n = 2^{k}$, HENCE $k = \log_{2} n$ $A (n) = \log_{2} n$ $A (n) \diamondsuit \Theta (\log n)$

1.2 Brute Force Approaches:

Introduction

Brute force is a straightforward approach to problem solving, usually directly based on the problem's statement and definitions of the concepts involved. Though rarely a source of clever or efficient algorithms, the brute-force approach should not be overlooked as an important algorithm design strategy. Unlike some of the otherstrategies, **brute force** is applicable to a very wide variety of problems. For some important problems (e.g., sorting, searching, string matching), the brute-force approach yields reasonable algorithms of at least some practical value with no limitation on instance size Even if too inefficient in general, a brute-force algorithm can still be useful for solving small-size instances of a problem. A brute-force algorithm can serve an important theoretical or educational purpose.

1.3 Selection Sort and Bubble Sort

Problem:Given a list of *n* orderable items (e.g., numbers, characters from some alphabet, character strings), rearrange them in nondecreasing order.

Selection Sort

ALGORITHM *SelectionSort*(*A*[0..*n* - 1])

//The algorithm sorts a given array by selection sort //Input: An array A[0..n - 1] of orderable elements //Output: Array A[0..n - 1] sorted in ascending order for i=0 to n - 2 do min=ifor j=i + 1 to n - 1 do if A[j] < A[min] min=j swap A[i] an

Example:

| 89 | 45 | 68 | 90 | 29 | 34 | 17 |
|----|----|----|----|----|----|----|
| 17 | 45 | 68 | 90 | 29 | 34 | 89 |
| 17 | 29 | 68 | 90 | 45 | 34 | 89 |
| 17 | 29 | 34 | 90 | 45 | 68 | 89 |
| 17 | 29 | 34 | 45 | 90 | 68 | 89 |
| 17 | 29 | 34 | 45 | 68 | 90 | 89 |
| 17 | 29 | 34 | 45 | 68 | 89 | 90 |
| | | | | | | |

Selection sort's operation on the list 89, 45, 68, 90, 29, 34, 17. Each line corresponds to one iteration of the algorithm, i.e., a pass through the list's tail to the right of the vertical bar; an element in bold indicates the smallest element found. Elements to the left of the vertical bar are in their final positions and are not considered in this and subsequent iterations.

Performance Analysis of the selection sort algorithm:

The input's size is given by the number of elements n.

The algorithm's basic operation is the key comparison A[j] < A[min]. The number of times it is executed depends only on the array's size and is given by

$$C(n) = \sum_{i=0}^{n-2} \sum_{j=i+1}^{n-1} 1 = \sum_{i=0}^{n-2} [(n-1) - (i+1) + 1] = \sum_{i=0}^{n-2} (n-1-i).$$

Thus, selection sort is a $O(n^2)$ algorithm on all inputs. The number of key swaps is only O(n) or, more precisely, *n*-1 (one for each repetition of the *i* loop). This property distinguishes selection sort positively from many other sorting algorithms.

Bubble Sort

Compare adjacent elements of the list and exchange them if they are out of order. Then we repeat the process, By doing it repeatedly, we end up 'bubbling up' the largest element to the last position on the list

ALGORITHM

BubbleSort(A[0..n - 1]) //The algorithm sorts array A[0..n - 1] by bubble sort //Input: An array A[0..n - 1] of orderable elements //Output: Array A[0..n - 1] sorted in ascending order for i=0 to n - 2 do for j=0 to n - 2 - i do if A[j + 1] < A[j]swap A[j] and A[j + 1]

Example

| 89 | $\stackrel{?}{\leftrightarrow}$ | 45 | - | 68 | | 90 | | 29 | | 34 | | 17 |
|----|---------------------------------|----|---|----|----|------|---|----|---------------------------------|-----|---|-----|
| 45 | | 89 | 4 | 68 | 22 | 90 | | 29 | | 34 | | 17 |
| 45 | | 68 | | 89 | 4 | 90 | ÷ | 29 | | 34 | | 17 |
| 45 | | 68 | | 89 | | 29 | | 90 | \leftrightarrow | 34 | | 17 |
| 45 | | 68 | | 89 | | 29 | | 34 | | 90 | 4 | 17 |
| 45 | | 68 | | 89 | | 29 | | 34 | | 17 | | 90 |
| 45 | .?. | 68 | 4 | 89 | 4 | 29 | | 34 | | 17 | | 190 |
| 45 | | 68 | | 29 | | 89 | ÷ | 34 | 120 | 17 | | 190 |
| 45 | | 68 | | 29 | | 34 | | 89 | $\stackrel{?}{\leftrightarrow}$ | 17 | | 190 |
| 45 | | 68 | | 29 | | 34 | | 17 | | 189 | | 90 |
| | | | | | | etc. | | | | | | |

The first 2 passes of bubble sort on the list 89, 45, 68, 90, 29, 34, 17. A new line is shown after a swap of two elements is done. The elements to the right of the vertical bar are in their final positions and are not considered in subsequent iterations of the algorithm

Bubble Sort the analysis

Clearly, the outer loop runs *n* times. The only complexity in this analysis in the inner loop. If we think about a single time the inner loop runs, we can get a simple bound by noting that it can never loop more than *n* times. Since the outer loop will make the inner loop complete *n* times, the comparison can't happen more than $O(n^2)$ times.

The number of key comparisons for the bubble sort version given above is the same for all arrays of size *n*.

$$C(n) = \sum_{i=0}^{n-2} \sum_{j=0}^{n-2-i} 1 = \sum_{i=0}^{n-2} [(n-2-i) - 0 + 1]$$
$$= \sum_{i=0}^{n-2} (n-1-i) = \frac{(n-1)n}{2} \in \Theta(n^2).$$

The number of key swaps depends on the input. For the worst case of decreasing arrays, it is the same as the number of key comparisons.

$$S_{worst}(n) = C(n) = \frac{(n-1)n}{2} \in \Theta(n^2).$$

Observation: if a pass through the list makes no exchanges, the list has been sorted and we can stop the algorithm Though the new version runs faster on some inputs, it is still in $O(n^2)$ in the worst and average cases. Bubble sort is not very good for big set of input. How ever bubble sort is very simple to code.

General Lesson From Brute Force Approach

A first application of the brute-force approach often results in an algorithm that can be improved with a modest amount of effort. Compares successive elements of a given list with a given search key until either a match is encountered (successful search) or the list is exhausted without finding a match (unsuccessful search)

1. 4 Sequential Search and Brute Force String Matching.

Sequential Search

ALGORITHM SequentialSearch2(A [0..n], K)

```
//The algorithm implements sequential search with a search key as a sentinel
//Input: An array A of n elements and a search key K
//Output: The position of the first element in A[0..n - 1] whose value is
// equal to K or -1 if no such element is found
A[n]=K
i=0
while A[i] = K do
i=i+1
if i < n return i
else return
```

Brute-Force String Matching

Given a string of *n* characters called the *text* and a string of *m* characters (m = n) called the *pattern*, find a substring of the text that matches the pattern. To put it more precisely, we want to find *i*—the index of the leftmost character of the first matching

substring in the text—such that

 $t_i = p_0, \dots, t_{i+j} = p_j, \dots, t_{i+m-1} = p_{m-1}$: $t_0 \dots t_i \dots t_{i+j} \dots t_{i+m-1} \dots t_{n-1}$ text *T* $p_0 \dots p_j \dots p_{m-1}$ pattern *P* 1. Pattern: 001011 Text: 1001010110100110101111010 2. Pattern: happy Text: It is never too late to have a happy childho



The algorithm shifts the pattern almost always after a single character comparison. in the worst case, the algorithm may have to make all *m* comparisons before shifting the pattern, and this can happen for each of the n - m + 1 tries. Thus, in the worst case, the algorithm is in $\theta(nm)$.

UNIT - 2 DIVIDE & CONQUER

1.1 Divide and Conquer

1.2 General Method

- 1.3 Binary Search
- 1.4 Merge Sort
- 1.5 Quick Sort and its performance

1.1 Divide and Conquer

Definition:

Divide & conquer is a general algorithm design strategy with a general plan as follows:

1. DIVIDE:

A problem's instance is divided into several smaller instances of the same problem, ideally of about the same size.

2. RECUR:

Solve the sub-problem recursively.

3. CONQUER:

If necessary, the solutions obtained for the smaller instances are combined to get a solution to the original instance.

Diagram 1 shows the general divide & conquer plan



NOTE:

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The base case for the recursion is sub-problem of constant size.

Advantages of Divide & Conquer technique:

- For solving conceptually difficult problems like Tower Of Hanoi, divide & conquer is a powerful tool
- Results in efficient algorithms
- Divide & Conquer algorithms are adapted foe execution in multi-processor machines
- Results in algorithms that use memory cache efficiently.

Limitations of divide & conquer technique:

- Recursion is slow
- Very simple problem may be more complicated than an iterative approach. Example: adding n numbers etc

1.2 General Method

General divide & conquer recurrence:

An instance of size n can be divided into b instances of size n/b, with "a" of them needing to be solved. [$a \ge 1, b > 1$].

Assume size n is a power of b. The recurrence for the running time T(n) is as follows:

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

where:

f(n) – a function that accounts for the time spent on dividing the problem into smaller ones and on combining their solutions

Therefore, the order of growth of T(n) depends on the values of the constants a & b and the order of growth of the function f(n).

Master theorem

Theorem: If $f(n) \in \Theta(n^d)$ with $d \ge 0$ in recurrence equation

$$T(n) = aT(n/b) + f(n),$$

then

$$T(n) = \begin{cases} \Theta \ (n^d) & \text{ if } a < b^d \\ \Theta \ (n^d \text{log } n) & \text{ if } a = b^d \\ \Theta \ (n^{\log_{a} a}) & \text{ if } a > b^d \end{cases}$$

Example:

Let T(n) = 2T(n/2) + 1, solve using master theorem. **Solution:**

Here: a = 2b = 2 $f(n) = \Theta(1)$ $\mathbf{d} = \mathbf{0}$

Therefore:

bre: $a > b^d$ i.e., $2 > 2^0$ Case 3 of master theorem holds good. Therefore: $T(n) \in \Theta(n^{\log_b a})$ $\in \Theta(n^{\log_2 2})$ $\in \Theta(n)$

1.3Binary Search

Description:

Binary tree is a dichotomic divide and conquer search algorithm. Ti inspects the middle element of the sorted list. If equal to the sought value, then the position has been found. Otherwise, if the key is less than the middle element, do a binary search on the first half, else on the second half.

Algorithm:

Algorithm can be implemented as recursive or non-recursive algorithm.

ALGORITHM BinSrch (A[0 ... n-1], key)

//implements non-recursive binary search//i/p: Array A in ascending order, key k//o/p: Returns position of the key matched else -1

```
1
    0
r
    n-1
while l \leq r do
       m
            (1+r)/2
       if key = = A[m]
              return m
       else
              if key < A[m]
                          m-1
                      r
              else
                      1
                          m+1
```

return -1

Analysis:

- Input size: Array size, n
- **Basic operation:** key comparison
- Depend on

Best – key matched with mid element

Worst - key not found or key sometimes in the list

• Let C(n) denotes the number of times basic operation is executed. Then $C_{worst}(n) = Worst$ case efficiency. Since after each comparison the algorithm divides the problem into half the size, we have $C_{worst}(n) = C_{worst}(n/2) + 1$ for n > 1

 $C_{\text{worst}}(n) = C_{\text{worst}}(n/2) + 1$ for n > 1

C(1) = 1

• Solving the recurrence equation using master theorem, to give the number of times the search key is compared with an element in the array, we have: C(n) = C(n/2) + 1

$$a = 1$$

$$b = 2$$

$$f(n) = n^{0}; d = 0$$

$$case \ 2 \ holds:$$

$$C(n) = \Theta \ (n^{d} \log n)$$

$$= \Theta (n^0 \log n)$$

$$= \Theta (\log n)$$

Applications of binary search:

- Number guessing game
- Word lists/search dictionary etc

Advantages:

- Efficient on very big list
- Can be implemented iteratively/recursively

Limitations:

- Interacts poorly with the memory hierarchy
- Requires given list to be sorted
- Due to random access of list element, needs arrays instead of linked list.

1.4Merge Sort

Definition:

Merge sort is a sort algorithm that splits the items to be sorted into two groups, recursively sorts each group, and merges them into a final sorted sequence.

Features:

- Is a comparison based algorithm
- Is a stable algorithm
- Is a perfect example of divide & conquer algorithm design strategy
- It was invented by John Von Neumann

Algorithm:

ALGORITHM Mergesort (A[0... n-1])

//sorts array A by recursive mergesort //i/p: array A //o/p: sorted array A in ascending order

if n > 1

```
copy A[0... (n/2 - 1)] to B[0... (n/2 - 1)]
copy A[n/2... n -1)] to C[0... (n/2 - 1)]
Mergesort ( B[0... (n/2 - 1)] )
Mergesort ( C[0... (n/2 - 1)] )
Merge ( B, C, A )
```

ALGORITHM Merge (B[0... p-1], C[0... q-1], A[0... p+q-1]) //merges two sorted arrays into one sorted array //i/p: arrays B, C, both sorted //o/p: Sorted array A of elements from B & C

```
\begin{split} I &\rightarrow 0 \\ j &\rightarrow 0 \\ k &\rightarrow 0 \\ \text{while } i
```

copy B [$i \dots p-1$] to A [$k \dots (p+q-1)$]

Example:

Apply merge sort for the following list of elements: 6, 3, 7, 8, 2, 4, 5, 1



Analysis:

- Input size: Array size, n
- **Basic operation:** key comparison
- Best, worst, average case exists: Worst case: During key comparison, neither of the two arrays becomes empty before the other one contains just one element.
- Let C(n) denotes the number of times basic operation is executed. Then $C(n) = 2C(n/2) + C_{merge}(n)$ for n > 1C(1) = 0

where, $C_{merge}(n)$ is the number of key comparison made during the merging stage. In the worst case:

$$\begin{split} C_{\text{merge}}(n) &= 2 \ C_{\text{merge}}(n/2) + n\text{-}1 \qquad \quad \text{for } n > 1 \\ C_{\text{merge}}(1) &= 0 \end{split}$$

• Solving the recurrence equation using master theorem:

```
C(n) = 2C(n/2) + n-1 \quad \text{for } n > 1

C(1) = 0

Here a = 2

b = 2

f(n) = n; d = 1

Therefore 2 = 2^1, case 2 holds

C(n) = \Theta (n^d \log n)

= \Theta (n^1 \log n)

= \Theta (n \log n)
```

Advantages:

- Number of comparisons performed is nearly optimal.
- Mergesort will never degrade to $O(n^2)$
- It can be applied to files of any size

Limitations:

• Uses O(n) additional memory.

1.6 Quick Sort and its performance Definition:

Quick sort is a well –known sorting algorithm, based on divide & conquer approach. The steps are:

- 1. Pick an element called pivot from the list
- 2. Reorder the list so that all elements which are less than the pivot come before the pivot and all elements greater than pivot come after it. After this partitioning, the pivot is in its final position. This is called the partition operation
- 3. Recursively sort the sub-list of lesser elements and sub-list of greater elements.

Features:

- Developed by C.A.R. Hoare
- Efficient algorithm
- NOT stable sort
- Significantly faster in practice, than other algorithms

ALGORITHM Quicksort (A[l ...r])

//sorts by quick sort //i/p: A sub-array A[1..r] of A[0..n-1], defined by its left and right indices 1 and r //o/p: The sub-array A[l..r], sorted in ascending order if l < rPartition (A[1..r]) // s is a split position Quicksort(A[1..s-1]) Quicksort(A[s+1..r] **ALGORITHM** Partition (A[1..r]) //Partitions a sub-array by using its first element as a pivot //i/p: A sub-array A[l..r] of A[0..n-1], defined by its left and right indices l and r (l < r) //o/p: A partition of A[l..r], with the split position returned as this function's value $p \rightarrow A[1]$ i→l $j \rightarrow r + 1;$ Repeat repeat $i \rightarrow i + 1$ until A[i] >= p //left-right scan //right-left scan repeat $j \rightarrow j - 1$ until A[j] < p if (i < j) //need to continue with the scan swap(A[i], a[j]) until $i \ge j$ //no need to scan swap(A[1], A[j])return j

Example: Sort by quick sort the following list: 5, 3, 1, 9, 8, 2, 4, 7, show recursion tree.



Analysis:

- Input size: Array size, n
- **Basic operation:** key comparison
- Best, worst, average case exists: Best case: when partition happens in the middle of the array each time. Worst case: When input is already sorted. During key comparison, one half is empty, while remaining n-1 elements are on the other partition.
- Let C(n) denotes the number of times basic operation is executed in worst case: Then

 $C(n)=C(n\mathchar`-1)+(n\mathchar`-1)$ for n>1 (2 sub-problems of size 0 and n-1 respectively) C(1)=1

Best case: $C(n) = 2C(n/2) + \Theta(n)$ (2 sub-problems of size n/2 each)

• Solving the recurrence equation using backward substitution/ master theorem, we have:

$$\begin{split} C(n) &= C(n\text{-}1) + (n\text{+}1) \ \text{for } n > 1; \ C(1) = 1 \\ C(n) &= \ \Theta \ (n^2) \end{split}$$

$$\begin{split} C(n) &= 2C(n/2) + \Theta(n). \\ &= \Theta(n^1 \log n) \\ &= \Theta(n \log n) \end{split}$$

NOTE:

The quick sort efficiency in average case is $\Theta(n \log n)$ on random input.

UNIT - 3 THE GREEDY METHOD

- 3.1 The General Method
- 3.2 Knapsack Problem
- **3.3 Job Sequencing with Deadlines**
- 3.4 Minimum-Cost Spanning Trees
- 3.5 Prim'sAlgorithm
- 3.6 Kruskal's Algorithm
- **3.7** Single Source Shortest Paths.

3.1The General Method

Definition:

Greedy technique is a general algorithm design strategy, built on following elements:

- configurations: different choices, values to find
- **objective function**: some configurations to be either maximized or minimized

The method:

- Applicable to **optimization problems** ONLY
- Constructs a solution through a sequence of steps
- Each step expands a partially constructed solution so far, until a complete solution to the problem is reached.
 - On each step, the choice made must be
- Feasible: it has to satisfy the problem's constraints
- Locally optimal: it has to be the best local choice among all feasible choices available on that step
- Irrevocable: Once made, it cannot be changed on subsequent steps of the algorithm

NOTE:

- Greedy method works best when applied to problems with the **greedy-choice** property
- A globally-optimal solution can always be found by a series of local improvements from a starting configuration.

Greedy method vs. Dynamic programming method:

- LIKE dynamic programming, greedy method solves optimization problems.
- LIKE dynamic programming, greedy method problems exhibit optimal substructure
- UNLIKE dynamic programming, greedy method problems exhibit the greedy choice property -avoids back-tracing.

Applications of the Greedy Strategy:

• Optimal solutions:

Change making Minimum Spanning Tree (MST) Single-source shortest paths Huffman codes

• Approximations:

Traveling Salesman Problem (TSP) Fractional Knapsack problem

3.2 Knapsack problem

- One wants to pack n items in a luggage
 - The ith item is worth v_i dollars and weighs w_i pounds
 - Maximize the value but cannot exceed W pounds
 - v_i, w_i, W are integers
- 0-1 knapsack \rightarrow each item is taken or not taken
- Fractional knapsack \rightarrow fractions of items can be taken
- Both exhibit the optimal-substructure property
 - 0-1: If item j is removed from an optimal packing, the remaining packing is an optimal packing with weight at most W-w_i
 - Fractional: If w pounds of item j is removed from an optimal packing, the remaining
 packing is an optimal packing with weight at most W-w that can be taken from other n-1
 items plus w_i w of item j

Greedy Algorithm for Fractional Knapsack problem

- Fractional knapsack can be solvable by the greedy strategy
 - Compute the value per pound v_i/w_i for each item
 - Obeying a greedy strategy, take as much as possible of the item with the greatest value per pound.
 - If the supply of that item is exhausted and there is still more room, take as much as possible of the item with the next value per pound, and so forth until there is no more room
 - O(*n* lg *n*) (we need to sort the items by value per pound)

O-1 knapsack is harder

• knapsack cannot be solved by the greedy strategy

- Unable to fill the knapsack to capacity, and the empty space lowers the effective value per pound of the packing
- We must compare the solution to the sub-problem in which the item is included with the solution to the sub-problem in which the item is excluded before we can make the choice



select a subset of the three items shown whose weight must not exceed 50 pounds. (b) The optimal subset includes items 2 and 3. Any solution with item 1 is suboptimal, even though item 1 has the greatest value per pound. (c) For the fractional knapsack problem, taking the items in order of greatest value per pound yields an optimal solution.

3.3 Job sequencing with deadlines

The problem is stated as below.

- There are n jobs to be processed on a machine.
- Each job i has a deadline $d_i\!\!\geq\!0$ and profit $p_i\!\!\geq\!\!0$.
- Pi is earned iff the job is completed by its deadline.
- The job is completed if it is processed on a machine for unit time.
- Only one machine is available for processing jobs.
- Only one job is processed at a time on the machine
- A feasible solution is a subset of jobs J such that each job is completed by its deadline.
- An optimal solution is a feasible solution with maximum profit value.

Example : Let n = 4, $(p_1, p_2, p_3, p_4) = (100, 10, 15, 27)$, $(d_1, d_2, d_3, d_4) = (2, 1, 2, 1)$

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|--------|
|--------|

| Sr.No. | Feasible | Processing | Profit value |
|--------|----------|----------------|------------------------|
| | Solution | Sequence | |
| (i) | (1,2) | (2,1) | 110 |
| (ii) | (1,3) | (1,3) or (3,1) | 115 |
| (iii) | (1,4) | (4,1) | 127 is the optimal one |
| (iv) | (2,3) | (2,3) | 25 |
| (v) | (3,4) | (4,3) | 42 |
| (vi) | (1) | (1) | 100 |
| (vii) | (2) | (2) | 10 |
| (viii) | (3) | (3) | 15 |
| (ix) | (4) | (4) | 27 |

- Consider the jobs in the non increasing order of profits subject to the constraint that the resulting job sequence J is a feasible solution.
- In the example considered before, the non-increasing profit vector is

 $(100 \ 27 \ 15 \ 10) \qquad (2 \ 1 \ 2 \ 1)$ $p_1 \ p_4 \ p_3 \ p_2 \qquad d_1 \ d \ d_3 \ d_2$ $J = \{ \ 1\} \text{ is a feasible one}$ $J = \{ \ 1, 4\} \text{ is a feasible one with processing sequence (4,1)}$ $J = \{ \ 1, 3, 4\} \text{ is not feasible}$ $J = \{ \ 1, 2, 4\} \text{ is not feasible}$ $J = \{ \ 1, 4\} \text{ is optimal}$

Theorem: Let J be a set of K jobs and

 $\Sigma=(i_1,i_2,\ldots,i_k) \text{ be a permutation of jobs in J such that } di_1 \leq di_2 \leq \ldots \leq di_k.$

J is a feasible solution iff the jobs in J can be processed in the order Σ without violating any deadly.

Proof:

• By definition of the feasible solution if the jobs in J can be processed in the order without violating any deadline then J is a feasible solution.

- So, we have only to prove that if J is a feasible one, then Σ represents a possible order in which the jobs may be processed.
- Suppose J is a feasible solution. Then there exists $\Sigma^1 = (r_1, r_2, ..., r_k)$ such that

$$d_{rj} \geq j, \qquad 1 \leq j <\!\! k$$

i.e.
$$d_{r1} \ge 1, d_{r2} \ge 2, \dots, d_{rk} \ge k.$$

each job requiring an unit time.

- $\Sigma = (i_1, i_2, ..., i_k)$ and $\Sigma^1 = (r_1, r_2, ..., r_k)$
- Assume $\Sigma^{-1} \neq \Sigma$. Then let a be the least index in which Σ^{-1} and Σ differ. i.e. a is such that $r_a \neq i_a$.
- Let $r_b = i_a$, so b > a (because for all indices j less than a $r_j = i_j$).
- In Σ^{1} interchange r_{a} and r_{b} .

•
$$\Sigma = (i_1, i_2, \dots i_a \quad i_b \quad i_k)$$
 [r_b occurs before r_a

• in i₁,i₂,...,i_k]

•
$$\Sigma^{1} = (\mathbf{r}_{1}, \mathbf{r}_{2}, \dots, \mathbf{r}_{a} | \mathbf{r}_{b} \dots \mathbf{r}_{k})$$

•
$$i_1=r_1, i_2=r_2, \dots i_{a-1}=r_{a-1}, i_a \neq r_b \text{ but } i_a = r_b$$

- We know $di_1 \le di_2 \le \dots di_a \le di_b \le \dots \le di_k$.
- Since $i_a = r_b$, $dr_b \le dr_a$ or $dr_a \ge dr_b$.
- In the feasible solution $dr_a \ge a \ dr_b \ge b$
- So if we interchange r_a and r_b , the resulting permutation $\Sigma^{11} = (s_1, \dots, s_k)$ represents an order with the least index in which Σ^{11} and Σ differ is incremented by one.
- Also the jobs in Σ^{11} may be processed without violating a deadline.
- Continuing in this way, Σ^1 can be transformed into Σ without violating any deadline.
- Hence the theorem is proved

GREEDY ALGORITHM FOR JOB SEQUENSING WITH DEADLINE

| Procedure greedy job (D, J, n) | J may be represented by |
|--|---------------------------------------|
| // J is the set of n jobs to be complete | ed // one dimensional array J (1: K) |
| // by their deadlines // | The deadlines are |
| J ← {1} | $D(J(1)) \le D(J(2)) \le \le D(J(K))$ |
| for I \leftarrow 2 to n do | To test if JU {i} is feasible, |

| If all jobs in $JU{i}$ can be completed | we insert i | into J and verify |
|---|-------------------------|--------------------------|
| by their deadlines | $D(J\mathbb{R}) \leq r$ | $1 \leq r \leq k{+}1$ |
| then $J \leftarrow JU{I}$ | | |
| end if | | |
| repeat | | |
| end greedy-job | | |
| Procedure JS(D,J,n,k) | | |
| // D(i) \geq 1, 1 \leq i \leq n are the deadlines // | | |
| $/\!/$ the jobs are ordered such that $/\!/$ | | |
| $/\!/ p_1 \!\geq\! p_2 \!\geq\! \dots \dots \geq\! p_n /\!/$ | | |
| // in the optimal solution , $D(J(i) \ge D(J(i - i)))$ | +1)) // | |
| // $1 \le i \le k$ | c // | |
| integer D(o:n), J(o:n), i, k, n, r | | |
| $D(0) \leftarrow J(0) \leftarrow 0$ | | |
| // J(0) is a fictious job with D(0) = 0 // | | |
| $K \leftarrow 1; J(1) \leftarrow 1 // job one is inserted into$ | to J // | |
| for i \leftarrow 2 to do // consider jobs in non inc | creasing order of | of pi // |
| // find the position of i and check feasib | oility of insertio | on // |
| $r \leftarrow k // r$ and k are indices for existing | ng job in J // | |
| // find r such that i can be inserted after | r // | |
| while $D(J(r)) > D(i)$ and $D(i) \neq r$ do | | |
| // job r can be processed after i and // | | |
| // deadline of job r is not exactly r // | | |
| r \leftarrow r-1 // consider whether job r-1 car | n be processed | after i // |
| repeat | | |
| if $D(J(r)) \ge d(i)$ and $D(i) > r$ then | | |
| // the new job i can come after existing | job r; insert i i | nto J at position r+1 // |
| for I \leftarrow k to r+1 by -1 do | | |
| $J(I+1) \leftarrow J(l) // \text{ shift jobs(r+1) to k right}$ | by// | |
| //one position // | | |

repeat

if $D(J(r)) \ge d(i)$ and D(i) > r then

// the new job i can come after existing job r; insert i into J at position r+1 //

for I \leftarrow k to r+1 by -1 do

 $J(I+1) \leftarrow J(l) // shift jobs(r+1) to k right by//$

//one position //

Repeat

COMPLEXITY ANALYSIS OF JS ALGORITHM

- Let n be the number of jobs and s be the number of jobs included in the solution.
- The loop between lines 4-15 (the for-loop) is iterated (n-1)times.
- Each iteration takes O(k) where k is the number of existing jobs.
- ∴ The time needed by the algorithm is $0(sn) s \le n$ so the worst case time is $0(n^2)$.

If $d_i = n - i + 1$ $1 \le i \le n$, JS takes $\theta(n^2)$ time

D and J need $\theta(s)$ amount of space.

3.4 Minimum-Cost Spanning Trees

Spanning Tree

Spanning tree is a connected acyclic sub-graph (tree) of the given graph (G) that includes all of G's vertices

Example: Consider the following graph



The spanning trees for the above graph are as follows:


Minimum Spanning Tree (MST)

Definition:

MST of a weighted, connected graph G is defined as: A spanning tree of G with **minimum total weight**.

Example: Consider the example of spanning tree:

For the given graph there are three possible spanning trees. Among them the spanning tree with the minimum weight 6 is the MST for the given graph

Question: Why can't we use **BRUTE FORCE** method in constructing MST ?

Answer: If we use Brute force method-

- Exhaustive search approach has to be applied.
- Two serious obstacles faced:
 - 1. The number of spanning trees grows exponentially with graph size.
 - 2. Generating all spanning trees for the given graph is not easy.

MST Applications:

- **Network design.** Telephone, electrical, hydraulic, TV cable, computer, road
- Approximation algorithms for NP-hard problems. Traveling salesperson problem, Steiner tree
- Cluster analysis.
- Reducing data storage in sequencing amino acids in a protein
- Learning salient features for real-time face verification
- Auto config protocol for Ethernet bridging to avoid cycles in a network, etc

3.5 Prim's Algorithm

Some useful definitions:

- **Fringe edge:** An edge which has one vertex is in partially constructed tree Ti and the other is not.
- Unseen edge: An edge with both vertices not in Ti

Algorithm:

ALGORITHM Prim (G)

//Prim's algorithm for constructing a MST

//Input: A weighted connected graph G = { V, E }

//Output: ET the set of edges composing a MST of G

// the set of tree vertices can be initialized with any vertex

$$V_T \rightarrow \{ v_0 \}$$

 $E_T \rightarrow \qquad Ø$

for $i \rightarrow 1$ to |V| - 1 do

Find a minimum-weight edge e^* = $(v^*,\,u^*)$ among all the edges $(v,\,u)$ such that v is in V_T and u is in V - V_T

 $V_T \rightarrow V_T U \{ u^* \}$

 $E_T \rightarrow E_T U \{ e^* \}$

return E_T

STEP 1: Start with a tree, T₀, consisting of one vertex

STEP 2: "Grow" tree one vertex/edge at a time

- Construct a series of expanding sub-trees $T_1, T_2, \dots T_{n-1}$.
- At each stage construct T_{i+1} from T_i by adding the minimum weight edge connecting a vertex in tree (Ti) to one vertex not yet in tree, **choose from** "fringe" edges (this is the "greedy" step!)

Algorithm stops when all vertices are included

Example:

Apply Prim's algorithm for the following graph to find MST.



Solution:

| Tree vertices | Remaining vertices | Graph |
|------------------|---|-------------|
| a (-, -) | $b(a,3)c(-,\infty)d(-,\infty)e(a,6)f(a,5)$ | a b |
| b (a, 3) | c (b, 1) d (-,∞) e (a, 6) f (b, 4) | a b 1 c |
| c(b,1) | d(c,6) e(a,6) f(b,4) | a b 1 c c f |



Efficiency:

Efficiency of Prim's algorithm is based on data structure used to store priority queue.

- Unordered array: Efficiency: $\Theta(n^2)$
- Binary heap: **Efficiency:** $\Theta(m \log n)$
- Min-heap: For graph with n nodes and m edges: Efficiency: $(n + m) \log n$

Conclusion:

- Prim's algorithm is a "vertex based algorithm"
- Prim's algorithm "Needs priority queue for locating the nearest vertex." The choice of priority queue matters in Prim implementation.
 - Array optimal for dense graphs
 - Binary heap better for sparse graphs
 - Fibonacci heap best in theory, but not in practice

3.6 Kruskal's Algorithm

Algorithm:

ALGORITHM Kruskal (G)

//Kruskal's algorithm for constructing a MST
//Input: A weighted connected graph G = { V, E }
//Output: ET the set of edges composing a MST of G

Sort E in ascending order of the edge weights

// initialize the set of tree edges and its size $E_T \rightarrow \emptyset$ edge_counter $\rightarrow 0$

//initialize the number of processed edges $K \rightarrow 0$

```
while edge_counter < |V| - 1

k \rightarrow k + 1

if E_T \cup \{ e_{i \ k} \} is acyclic

E_T \rightarrow E_T \cup \{ e_{i \ k} \}

edge_counter \rightarrow edge_counter + 1
```

return E_T

The method:

STEP 1: Sort the edges by increasing weight

STEP 2: Start with a forest having |V| number of trees.

- **STEP 3**: Number of trees are reduced by ONE at every inclusion of an edge At each stage:
 - Among the edges which are not yet included, select the one with minimum weight AND which does not form a cycle.
 - the edge will reduce the number of trees by one by combining two trees of the forest

Algorithm stops when |V| -1 edges are included in the MST i.e : when the number of trees in the forest is reduced to ONE.

Apply Kruskal's algorithm for the following graph to find MST.



Solution:

The list of edges is:

| Edge | ab | af | ae | bc | bf | cf | cd | df | de | ef |
|--------|----|----|----|----|----|----|----|----|----|----|
| Weight | 3 | 5 | 6 | 1 | 4 | 4 | 6 | 5 | 8 | 2 |

Sort the edges in ascending order:

| Edge | bc | ef | ab | bf | cf | af | df | ae | cd | de |
|--------|----|----|----|----|----|----|----|----|----|----|
| Weight | 1 | 2 | 3 | 4 | 4 | 5 | 5 | 6 | 6 | 8 |



| Edge | ab | 3 b 1 c |
|---------------------|-------------|---|
| Weight | 3 | |
| Insertion status | YES | |
| Insertion order | 3 | e |
| Edge | bf | 3 (b) 1 (c) |
| Weight | 4 | a 4 f d |
| Insertion status | YES | |
| Insertion order | 4 | e |
| Edge | cf | |
| Weight | 4 | |
| Insertion status | NO | |
| Insertion order | - | |
| | | |
| Edge | af | |
| Weight | 5 | |
| Insertion status | NO | |
| Insertion order | - | |
| Edge | df | 3 (b) 1 |
| Weight | 5 | |
| Insertion status | YES | $\begin{array}{cccc} a & 4 & f \\ \hline & 5 \\ \hline & 2 \end{array}$ |
| Insertion order | 5 | e |
| Algorithm | stops as V | / -1 edges are included in the MST |

Efficiency:

Efficiency of Kruskal's algorithm is based on the time needed for sorting the edge weights of a given graph.

• With an efficient sorting algorithm: Efficiency: $\Theta(|\mathbf{E}| \log |\mathbf{E}|)$

Conclusion:

- Kruskal's algorithm is an "edge based algorithm"
- Prim's algorithm with a heap is faster than Kruskal's algorithm.

3.7 Single Source Shortest Paths.

Some useful definitions:

• Shortest Path Problem: Given a connected directed graph G with non-negative weights on the edges and a root vertex r, find for each vertex x, a directed path P (x) from r to x so that the sum of the weights on the edges in the path P (x) is as small as possible.

Algorithm

- By Dutch computer scientist Edsger Dijkstra in 1959.
- Solves the single-source shortest path problem for a graph with nonnegative edge weights.
- This algorithm is often used in routing.

E.g.: Dijkstra's algorithm is usually the working principle behind link-state routing protocols

ALGORITHM Dijkstra(G, s)

//Input: Weighted connected graph G and source vertex s

//Output: The length Dv of a shortest path from s to v and its penultimate vertex Pv for every vertex v in V $% \left(V_{x}^{2}\right) =0$

//initialize vertex priority in the priority queue Initialize (Q)

for every vertex v in V do

 $D_v \rightarrow \infty$; $P_v \rightarrow null$ // Pv, the parent of v

insert(Q, v, D_v) //initialize vertex priority in priority queue

 $d_s \rightarrow 0$

//update priority of s with ds, making ds, the minimum $Decrease(Q, s, d_s)$



for $i \rightarrow 0$ to |V| - 1 do $u^* \rightarrow DeleteMin(Q)$ //expanding the tree, choosing the locally best vertex $V_T \rightarrow V_T \ U \{u^*\}$ for every vertex u in $V - V_T$ that is adjacent to u* do if $Du^* + w (u^*, u) < Du$ $Du \rightarrow Du + w (u^*, u)$; $Pu \quad u^*$ Decrease(Q, u, Du)

The method

Dijkstra's algorithm solves the single source shortest path problem in 2 stages. **Stage 1**: A greedy algorithm computes the shortest distance from **source to all other nodes** in the graph and saves in a data structure.

Stage 2 : Uses the data structure for finding a shortest path from source to any vertex v.

- At each step, and for each vertex x, keep track of a "distance" D(x) and a directed path P(x) from root to vertex x of length D(x).
- Scan first from the root and take initial paths P(r, x) = (r, x) with D(x) = w(rx) when rx is an edge,
 D(x) = ∞ when rx is not an edge.
 For each temporary vertex y distinct from x, set
 D(y) = min{ D(y), D(x) + w(xy) }

Example:

Apply Dijkstra's algorithm to find Single source shortest paths with vertex \mathbf{a} as the source.



Solution:

Length Dv of shortest path from source (s) to other vertices v and Penultimate vertex Pv for every vertex v in V:

Da = 0, Pa = null $Db = \infty, Pb = null$ $Dc = \infty, Pc = null$ $Dd = \infty, Pd = null$ $De = \infty, Pe = null$ $Df = \infty, Pf = null$

| Tree | RemainingDistance & Path | Graph |
|------------|---|--|
| vertices | vertices vertex | |
| a (-, 0) | Da = 0 Pa = a $b(a, \mathbf{D}_{b} = 3 \mathbf{Pb} = [\mathbf{a}, \mathbf{b}]$ $c(-, \alpha D)c = \mathbf{\infty} Pc = null$ $d(-, \alpha D)d = \mathbf{\infty} Pd = null$ $e(a, \mathbf{d})e = 6 Pe = [\mathbf{a}, \mathbf{e}]$ $f(a, 5)f = 5 Pf = [\mathbf{a}, f]$ | a b |
| b (a, 3) | $Da = 0 Pa = a$ $c (b, 3 + Db) = 3 Pb = [a, b]$ $d (-, \alpha Dc = 4 Pc = [a, b, c]$ $e (a, 6Dd = \infty Pd = null$ $f (a, 5De = 6 Pe = [a, e]$ $Df = 5 Pf = [a, f]$ | a b 1 c |
| c (b, 4) | Da = 0 	Pa = a Db = 3 	Pb = [a, b] d (c, 4+6) = 4 	Pc = [a,b,c] e (a, 6bd=10 	Pd = [a,b,c,d] f (a, 5) 	De = 6 	Pe = [a, e] Df = 5 	Pf = [a, f] | af |
| f (a, 5) | Da = 0 	Pa = a Db = 3 	Pb = [a, b] Dc = 4 	Pc = [a,b,c] d (c , 0bd=0 	Pd = [a,b,c,d] e (a , 0be = 6 	Pe = [a, e] Df = 5 	Pf = [a, f] | a 6 e |
| e (a, 6) | Da = 0 Pa = a Db = 3 Pb = [a, b] Dc = 4 Pc = [a,b,c] $d (c, 1 \bigoplus d = 10 Pd = [a,b,c,d]$ De = 6 Pe = [a, e] Df = 5 Pf = [a, f] | $ \begin{array}{c} $ |
| d(c, 10) | ed | Algorithm stops since no ges to scan |

Conclusion:

- Doesn't work with negative weights
- Applicable to both undirected and directed graphs
- Use unordered array to store the priority queue: Efficiency = $\Theta(\mathbf{n}^2)$
- Use min-heap to store the priority queue: **Efficiency = O(m log n)**

UNIT - 4 Dynamic Programming

4.1 The General Method

4.2 Warshall's Algorithm

4.3 Floyd's Algorithm for the All-Pairs Shortest Paths Problem

4.4 Single-Source Shortest Paths

4.5 General Weights 0/1 Knapsack

4.6 The Traveling Salesperson problem.

4.1 The General Method

Definition

Dynamic programming (DP) is a general algorithm design technique for solving problems with overlapping sub-problems. This technique was invented by American mathematician "Richard Bellman" in 1950s.

Key Idea

The key idea is to save answers of overlapping smaller sub-problems to avoid recomputation.

Dynamic Programming Properties

- An instance is solved using the solutions for smaller instances.
- The solutions for a smaller instance might be needed multiple times, so store their results in a table.
- Thus each smaller instance is solved only once.
- Additional space is used to save time.

Dynamic Programming vs. Divide & Conquer

LIKE divide & conquer, dynamic programming solves problems by combining solutions to sub-problems. UNLIKE divide & conquer, sub-problems are NOT independent in dynamic programming.

| Divide & Conquer | Dynamic Programming |
|--|---|
| 1. Partitions a problem into independent smaller sub-problems | 1. Partitions a problem into overlapping sub-problems |
| 2. Doesn't store solutions of sub- problems. (Identical sub-problems may arise - results in the same computations are performed repeatedly.) | 2. Stores solutions of sub- problems: thus avoids calculations of same quantity twice |
| 3. Top down algorithms: which logically progresses from the initial instance down to the smallest sub-instances via intermediate sub-instances. | 3. Bottom up algorithms: in which the smallest sub-problems are explicitly solved first and the results of these used to construct solutions to progressively larger sub-instances |

Dynamic Programming vs. Divide & Conquer: EXAMPLE Computing Fibonacci Numbers

1. Using standard recursive formula:

$$F(n) = \begin{cases} 0 & \text{if } n=0 \\ 1 & \text{if } n=1 \\ F(n-1) + F(n-2) & \text{if } n >1 \end{cases}$$

Algorithm F(n)

// Computes the nth Fibonacci number recursively by using its definitions

- // Input: A non-negative integer n
- // Output: The nth Fibonacci number

if $n==0 \parallel n==1$ then

return n

else

return F(n-1) + F(n-2)

Algorithm F(n): Analysis

- Is too expensive as it has repeated calculation of smaller Fibonacci numbers.
- Exponential order of growth.



2. Using Dynamic Programming:

Algorithm F(n)

// Computes the nth Fibonacci number by using dynamic programming method

// Input: A non-negative integer n

// Output: The nth Fibonacci number

A[0] 0 A[1] 1

for i 2 to n do

 $A[i] \quad A[i-1] + A[i-2]$

return A[n]

Algorithm F(n): Analysis

- Since it caches previously computed values, saves time from repeated computations of same sub-instance
- Linear order of growth

Rules of Dynamic Programming

- 1. **OPTIMAL SUB-STRUCTURE:** An optimal solution to a problem contains optimal solutions to sub-problems
- 2. **OVERLAPPING SUB-PROBLEMS:** A recursive solution contains a "small" number of distinct sub-problems repeated many times
- 3. **BOTTOM UP FASHION:** Computes the solution in a bottom-up fashion in the final step

Three basic components of Dynamic Programming solution

The development of a dynamic programming algorithm must have the following three basic components

- 1. A recurrence relation
- 2. A tabular computation
- 3. A backtracking procedure

Example Problems that can be solved using Dynamic Programming method

- 1. Computing binomial co-efficient
- 2. Compute the longest common subsequence
- 3. Warshall's algorithm for transitive closure
- 4. Floyd's algorithm for all-pairs shortest paths
- 5. Some instances of difficult discrete optimization problems like knapsack problem And traveling salesperson problem

4.2 Warshall's Algorithm

Some useful definitions:

- **Directed Graph:** A graph whose every edge is directed is called directed graph OR digraph
- Adjacency matrix: The adjacency matrix $A = \{a_{ij}\}$ of a directed graph is the boolean matrix that has
 - \circ 1 if there is a directed edge from ith vertex to the jth vertex
 - \circ 0 Otherwise
- Transitive Closure: Transitive closure of a directed graph with n vertices can be defined as the n-by-n matrix T={tij}, in which the elements in the ith row (1≤ i ≤ n) and the jth column(1≤ j ≤ n) is 1 if there exists a nontrivial directed path (i.e., a directed path of a positive length) from the ith vertex to the jth vertex, otherwise tij is 0. The transitive closure provides reach ability information about a digraph.

Computing Transitive Closure:

- We can perform DFS/BFS starting at each vertex
 - Performs traversal starting at the ith vertex.
 - Gives information about the vertices reachable from the ith vertex
 - **Drawback:** This method traverses the same graph several times.
 - **Efficiency** : (O(n(n+m)))
- Alternatively, we can use dynamic programming: the Warshall's Algorithm

Underlying idea of Warshall's algorithm:

- Let A denote the initial boolean matrix.
- The element r(k) [i, j] in ith row and jth column of matrix Rk (k = 0, 1, ..., n) is equal to 1 if and only if there exists a directed path from ith vertex to jth vertex with intermediate vertex if any, numbered not higher than k

Recursive Definition: •

• **Case 1:** A path from *vi* to *vj* restricted to using only vertices from $\{v_1, v_2, \dots, v_k\}$ as intermediate vertices does not use v_k . Then R

$$(k) [i, j] = \mathbf{R}(k-1) [i, j].$$

Case 2: A path from vi to vj restricted to using only vertices from $\{v_1, v_2, \dots, v_k\}$ as intermediate vertices do use vk. Then

R(k) [i, j] = R(k-1) [i, k]**AND** R(k-1) [k, j].

R(k)[i, j] = R(k-1)[i, j] OR (R(k-1)[i, k] AND R(k-1)[k, j])

Algorithm:

Algorithm Warshall(A[1..n, 1..n])

// Computes transitive closure matrix

// Input: Adjacency matrix A

// Output: Transitive closure matrix R

R(0) Α

for $k \rightarrow 1$ to n do

for $i \rightarrow 1$ to n do

```
for j \rightarrow 1 to n do
```

$$R(k)[i, j] \rightarrow R(k-1)[i, j] OR (R(k-1)[i, k] AND R(k-1)[k, j])$$

return R(n)

Find Transitive closure for the given digraph using Warshall's algorithm.



Solution:

| | | | Α | В | С | D | |
|--------------|---|---|---|---|---|---|--|
| K (0) | = | Α | 0 | 0 | 1 | 0 | |
| | | В | 1 | 0 | 0 | 1 | |
| | | С | 0 | 0 | 0 | 0 | |
| | | D | 0 | 1 | 0 | 0 | |

| | | | | р | C | D | | | | р | C | D |
|-----------------|-----------------|--|---------------------------------|------------------|---------------|----|--|----|------|--------|---|---|
| R(0) | k = 1 | | Α | В | C | D | | | A | В | C | D |
| | Vertex 1 | A | 0 | 0 | 1 | 0 | | A | 0 | 0 | 1 | 0 |
| | can be | B | 1 | 0 | 0 | 1 | | B | 1 | 0 | 1 | 1 |
| | intermediate | C | 0 | 0 | 0 | 0 | | C | 0 | 0 | 0 | 0 |
| | | n | Ň | 1 | Ň | Ň | | D | Ň | 1 | Ň | Ň |
| | node | | U | I | U | U | | D | U | 1 | U | U |
| | | R1[2, = R0 R0 = 0 C = 1 | ,3] [2,3] [2,1])R (1 | OR AND AND | R0[1, 0 1) | 3] | | | | | | |
| R(1) | k = 2 | | Α | В | C | D | | | Α | В | С | D |
| | Vertex | A | 0 | 0 | 1 | 0 | | Α | 0 | 0 | 1 | 0 |
| | $\{1,2\}$ can | B | 1 | 0 | 1 | 1 | | B | 1 | 0 | 1 | 1 |
| | be | C | 0 | 0 | 0 | 0 | | C | 0 | 0 | 0 | 0 |
| | intermediate | D | 0 | 1 | 0 | 0 | | D | 1 | 1 | 1 | 1 |
| | nodes | D 0 1 0 0 R2[4,1] = R1[4,1] OR R1[4,2] AND R1[2,1] = 0 OR (1 AND 1) = 1 R2[4,3] = R1[4,3] OR R1[4,2] AND R1[2,3] = 0 OR (1 AND 1) = 1 R2[4,4] = R1[4,4] OR R1[4,2] AND R1[2,4] = 0 OR (1 AND 1) = 1 | | | | | | | | | | |
| $\mathbf{P}(2)$ | k - 3 | | Α | B | С | D | | | Α | B | С | D |
| K(2) | N - J Vortov | Δ | | ñ | 1 | Õ | | Δ | 0 | ñ | 1 | 0 |
| | vertex | R | 1 | ñ | 1 | 1 | | R | 1 | ñ | 1 | 1 |
| | $\{1,2,3\}$ can | | | U A | 1 | 1 | | | | U A | 1 | 1 |
| | be | | U | U | U | 0 | | | U | U | U | 0 |
| | intermediate | D | 1 | 1 | 1 | 1 | | D | 1 | 1 | 1 | 1 |
| | nodes | | | | | | | NO | CHAI | NGE | | |

| R(3) | k = 4 | | A | B | С | D | | | A | B | С | D |
|------|--------------|-------|--|-----|--------------|----|-------|--------|--------|---------|------|-----|
| | Vertex | A | 0 | 0 | 1 | 0 | | Α | 0 | 0 | 1 | 0 |
| | {1.2.3.4 } | B | 1 | 0 | 1 | 1 | | B | 1 | 1 | 1 | 1 |
| | can be | C | 0 | 0 | 0 | 0 | | С | 0 | 0 | 0 | 0 |
| | intermediate | D | 1 | 1 | 1 | 1 | | D | 1 | 1 | 1 | 1 |
| | nodes | | | | | | ' | | I | | | |
| | nodes | R4[2. | .21 | | | | | | | | | |
| | | = R3 | [2.2] | OR | | | | | | | | |
| | | R3 | $R_{3}[2, 4] \text{ AND } R_{3}[4, 2]$ | | | | | | | | | |
| | | = 0.0 | R (1 | ΑΝΓ | 1) | -1 | | | | | | |
| | | - 0 C | | | · 1) | | | | | | | |
| | | - 1 | | | | | | | | | | |
| | | | | | ~ | | | | | | | |
| R(4) | | | A | В | C | D | . - | | NOT | | | |
| | | A | 0 | 0 | 1 | 0 | | IKA | NSII | IVE | CLOS | URE |
| | | B | 1 | 1 | 1 | 1 | f | tor th | ie giv | ven gra | aph | |
| | | C | 0 | 0 | 0 | 0 | | | | | | |
| | | D | 1 | 1 | 1 | 1 | | | | | | |

Efficiency:

- Time efficiency is $\Theta(n^3)$
- Space efficiency: Requires extra space for separate matrices for recording intermediate results of the algorithm.

4.3Floyd's Algorithm to find -ALL PAIRS SHORTEST PATHS Some useful definitions:

- Weighted Graph: Each edge has a weight (associated numerical value). Edge weights may represent costs, distance/lengths, capacities, etc. depending on the problem.
- Weight matrix: W(i,j) is
 - \circ 0 if i=j
 - $\circ \quad \infty$ if no edge b/n i and j.
 - \circ "weight of edge" if edge b/n i and j.

Problem statement:

Given a weighted graph G(V, Ew), the all-pairs shortest paths problem is to find the shortest path between every pair of vertices (vi, vj) \in V.

Solution:

A number of algorithms are known for solving All pairs shortest path problem

- Matrix multiplication based algorithm
- Dijkstra's algorithm
- Bellman-Ford algorithm
- Floyd's algorithm

Underlying idea of Floyd's algorithm:

- Let W denote the initial weight matrix.
- Let D(k) [i, j] denote cost of shortest path from i to j whose intermediate vertices are a subset of $\{1, 2, \dots, k\}$.
- Recursive Definition

Case 1:

A shortest path from vi to vj restricted to using only vertices from $\{v_1, v_2, \dots, v_k\}$ as intermediate vertices does not use vk. Then

D(k) [i, j] = D(k-1) [i, j].

Case 2:

A shortest path from vi to vj restricted to using only vertices from $\{v_1, v_2, \dots, v_k\}$ as intermediate vertices do use vk. Then

$$D(k) [i, j] = D(k-1) [i, k] + D(k-1) [k, j].$$

We conclude:

 $D(k)[i, j] = \min \{ D(k-1) [i, j], D(k-1) [i, k] + D(k-1) [k, j] \}$

Algorithm:

Algorithm Floyd(W[1..n, 1..n])

// Implements Floyd's algorithm // Input: Weight matrix W // Output: Distance matrix of shortest paths' length D W for k \rightarrow 1 to n do for $i \rightarrow 1$ to n do for $j \rightarrow 1$ to n do $D[i, j] \rightarrow min \{ D[i, j], D[i, k] + D[k, j] \}$

return D

Example:

Find All pairs shortest paths for the given weighted connected graph using Floyd's algorithm.



Solution:

D(0) =B С Α Α 5 2 0 B 4 ∞ 0 С ∞ 3 0

| D(0) | k = 1 | | | | | | | | | |
|--------|------------------|-------------|--|-------------------------|---------------|--------------------|------------------------|------------------|--------------|---|
| | Vertex 1 | | | | | | | | | |
| 1 | can be | | | | | I. | | | | |
| | intermediate | | Α | В | С | | Α | В | С | |
| | node | A | 0 | 2 | 5 | A | 0 | 2 | 5 | |
| | | В | 4 | 0 | ∞ | B | 4 | 0 | 9 | |
| | | С | ∞ | 3 | 0 | C C | ∞ | 3 | 0 | |
| | | | | | | | I | | | ' |
| | | D1[2.3 | 5] | | | | | | | |
| | | = min | { D 0 [: | 2.31. | | | | | | |
| | | | D0 [2 | -,e,, .1] + D | 0[1.3] | L I | | | | |
| D(1) | 1 0 | = min | {∞.(4 | (1+5) | • [_,•]] | | | | | |
| D(1) | $\mathbf{k} = 2$ | - 9 | ι ,(- | T (C)] | | | | | | |
| | Vertex 1,2 | | | | | | | | | |
| | can be | | • | P | C | | • | P | C | |
| | intermediate | | A | В | C - | | A | B | C F | |
| | nodes | | 0 | 2 | 5 | | 0 | 2 | 5 | |
| | | B | 4 | 0 | 9 | B | 4 | 0 | 9 | |
| | | C | ∞ | 3 | 0 | C | 7 | 3 | 0 | |
| | | | _ | | | | | | | |
| | | D2[3,1 | .] | | | | | | | |
| | | = min | { D1 [3 | 3,1], | | | | | | |
| | | | D1 [3 | ,2] + D | 01 [2,1] } | } | | | | |
| D(2) | k = 3 | = min | {∞,(4 | 4 + 3) } | ł | | | | | |
| = (=) | Vertex 1.2.3 | = 7 | | | | | | | | |
| | can be | | | | | | | | | |
| | intermediate | | A | B | С | | Α | В | С | |
| | nodes | A | 0 | 2 | 5 | A | 0 | 2 | 5 | |
| | | В | 4 | 0 | 9 | B | 4 | 0 | 9 | |
| | | C | 7 | 3 | 0 | | 7 | 3 | 0 | |
| D(3) | | | | C | v | | | C | v | |
| - (-) | | | | | | NO | Change | | | |
| | | | | | | | ~ | | | |
| | | | | В | С | | | | | |
| | | | A | | | | ALL PAIRS SHORTEST | | | |
| | | A | | $\overline{2}$ | 5 | | , PAIR | S SHO | RTEST | |
| | | A B | $\begin{vmatrix} 0 \\ 4 \end{vmatrix}$ | 2 0 | 5 9 | ALL PAT | / PAIR: HS for | S SHO the giv | RTEST 'en | |
| | | A B C | A 0 4 7 | 2 0 3 | 5 9 0 | ALL PAT grap | / PAIRS HS for h | S SHO the giv | RTEST 'en | |
| | | A B C | A 0 4 7 | 2 0 3 | 5 9 0 | ALL PAT grap | / PAIR: HS for h | 5 SHO the giv | RTEST 7en | |
| | | A B C | A 0 4 7 | 2 0 3 | 5 9 0 | ALL PAT grap | / PAIR: HS for h | 5 SHO the giv | RTEST 'en | |

4.4 0/1 Knapsack Problem Memory function Definition:

Given a set of n items of known weights w1,...,wn and values v1,...,vn and a knapsack of capacity W, the problem is to find the most valuable subset of the items that fit into the knapsack.

Knapsack problem is an OPTIMIZATION PROBLEM

Dynamic programming approach to solve knapsack problem

Step 1:

Identify the smaller sub-problems. If items are labeled 1..n, then a sub-problem would be to find an optimal solution for $Sk = \{items | abeled 1, 2, ... k\}$

Step 2:

Recursively define the value of an optimal solution in terms of solutions to smaller problems.

Initial conditions:

 $\begin{array}{ll} V[\ 0, j \] = 0 & \mbox{ for } j \geq 0 \\ V[\ i, 0 \] = 0 & \mbox{ for } i \geq 0 \end{array}$

Recursive step:

| ſ | - max { V[i-1, | j], vi +V[i-1, j - wi] } |
|---------------------|-----------------|-----------------------------|
| $V[i, j] = \langle$ | | if j - wi ≥ 0 |
| l | V[i-1, j] | if j - wi < 0 |

Step 3:

Bottom up computation using iteration

Question:

Apply bottom-up dynamic programming algorithm to the following instance of the knapsack problem Capacity W= 5

| Item # | Weight (Kg) | Value (Rs.) |
|--------|-------------|-------------|
| 1 | 2 | 3 |
| 2 | 3 | 4 |
| 3 | 4 | 5 |
| 4 | 5 | 6 |

Solution:

Using dynamic programming approach, we have:

| Step | Calculation | Table | | | | | | |
|------|-----------------------------------|---------------|------------|---|---|---|---|---|
| 1 | Initial conditions: | | | | | | | |
| | $V[0, j] = 0$ for $j \ge 0$ | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | $V[i, 0] = 0$ for $i \ge 0$ | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | | 1 | 0 | | | | | |
| | | 2 | 0 | | | | | |
| | | 3 | 0 | | | | | |
| | | 4 | 0 | | | | | |
| 2 | W1 = 2, | | | | | | | |
| | Available knapsack capacity = 1 | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | W1 > WA, CASE 1 holds: | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | V[i, j] = V[i-1, j] | 1 | 0 | 0 | | | | |
| | V[1,1] = V[0,1] = 0 | 2 | 0 | | | | | |
| | | 3 | 0 | | | | | |
| | | 4 | 0 | | | | | |
| 3 | W1 = 2, | | | | | | | |
| | Available knapsack capacity $= 2$ | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | W1 = WA, CASE 2 holds: | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | $V[i, j] = max \{ V[i-1, j],$ | 1 | 0 | 0 | 3 | | | |
| | vi +V[i-1, j - wi] } | 2 | 0 | | | | | |
| | $V[1,2] = \max \{ V[0,2],$ | 3 | 0 | | | | | |
| | 3 + V[0, 0] | 4 | 0 | | | | | |
| 1 | $= \max \{0, 3+0\} = 3$ W1 - 2 | | I | | | | | |
| - | Available knapsack capacity = | V[i,j] | i=0 | 1 | 2 | 3 | 4 | 5 |
| | 3,4,5 | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | W1 < WA, CASE 2 holds: | 1 | 0 | 0 | 3 | 3 | 3 | 3 |
| | $V[i, j] = max \{ V[i-1, j],$ | 2 | 0 | | | | | |
| | vi +V[i-1, j - wi] } | 3 | 0 | | | | | |
| | $V[1,3] = \max \{ V[0,3],$ | 4 | 0 | | | | | |
| | 3 + V[0, 1] | L | 1 | 1 | 1 | | 1 | |
| 5 | $W_2 = 3.$ | | | | | | | |
| | Available knapsack capacity = 1 | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | W2 >WA, CASE 1 holds: | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | V[i, j] = V[i-1, j] | 1 | 0 | 0 | 3 | 3 | 3 | 3 |
| | V[2,1] = V[1,1] = 0 | 2 | 0 | 0 | | | | |
| | | 3 | 0 | | | | | |
| 1 | | - | | | | | | |

Analysis and Design of Algorithm

| 6 | W2 = 3, | | | | | | | |
|----|---|---------------|-------------|---|---|---|---|---|
| | Available knapsack capacity = 2 | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | W2 >WA, CASE 1 holds: | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | V[i, j] = V[i-1, j] | 1 | 0 | 0 | 3 | 3 | 3 | 3 |
| | V[2,2] = V[1,2] = 3 | 2 | 0 | 0 | 3 | | | |
| | | 3 | 0 | | | | | |
| | | 4 | 0 | | | | | |
| | | | | | | | | |
| 7 | W2 = 3, | | | | | | | |
| | Available knapsack capacity = 3 | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | W2 = WA, CASE 2 holds: | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | $V[1, j] = \max \{ V[1-1, j],$ | 1 | 0 | 0 | 3 | 3 | 3 | 3 |
| | V1 + V[1-1,] - W1] | 2 | 0 | 0 | 3 | 4 | | |
| | $v[2,3] = \max \{ v[1,3], 4 \mid V[1,0] \}$ | 3 | 0 | | | | | |
| | 4 + v[1, 0] | 4 | 0 | | | | | |
| 8 | $-\max\{3, 4+0\} = 4$ W/2 = 3 | | | | | | | |
| 0 | 4 vailable knapsack capacity -4 | VI; j] | j_0 | 1 | 2 | 2 | 1 | 5 |
| | $W^2 < WA$ CASE 2 holds: | <u> </u> | J -0 | | 0 | 0 | 0 | 0 |
| | $V[i i] = max \{ V[i-1 i] \}$ | 1-0 | 0 | 0 | 3 | 3 | 3 | 3 |
| | vi +V[i-1, i - wi] } | | 0 | 0 | 2 | 3 | 3 | 5 |
| | $V[2,4] = \max \{ V[1,4], \dots \}$ | | 0 | 0 | 5 | 4 | 4 | |
| | 4 +V[1,1]} | | 0 | | | | | |
| | $= \max \{ 3, 4+0 \} = 4$ | 4 | 0 | | | | | |
| | | | | | | | | |
| 9 | W2 = 3, | | | | | | | |
| | Available knapsack capacity = 5 | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | W2 < WA, CASE 2 holds: | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | $V[i, j] = max \{ V[i-1, j],$ | 1 | 0 | 0 | 3 | 3 | 3 | 3 |
| | vi +V[i-1, j - wi] } | 2 | 0 | 0 | 3 | 4 | 4 | 7 |
| | $V[2,5] = max \{ V[1,5],$ | 3 | 0 | | | | | |
| | 4 +V[1,2]} | 4 | 0 | | | | | |
| | $= \max \{ 3, 4+3 \} = 7$ | | | | 1 | | 1 | I |
| 10 | | | | | | | | |
| 10 | $W_{3} = 4,$ | | | | | | | |
| | Available knapsack capacity = $1.2.2$ | V[i,j] | j=0 | | | 3 | 4 | 5 |
| | $ 1,2,3 \rangle = 0$ | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | $W_{3} > WA, CASE I HOLDS:$ | | 0 | 0 | 3 | 3 | 3 | 3 |
| | v [1, J] — v [1-1, J] | 2 | 0 | 0 | 3 | 4 | 4 | 7 |
| | | 3 | 0 | 0 | 3 | 4 | | |
| | | 4 | 0 | | | | | |

| 11 | W3 = 4, | _ | | | | | | |
|----|---------------------------------|---------------|----------------|-------|---|---|---|---|
| | Available knapsack capacity = 4 | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | W3 = WA, CASE 2 holds: | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | $V[i, j] = max \{ V[i-1, j],$ | 1 | 0 | 0 | 3 | 3 | 3 | 3 |
| | vi +V[i-1, j - wi] } | 2 | 0 | 0 | 3 | 4 | 4 | 7 |
| | $V[3,4] = \max \{ V[2,4], $ | 3 | 0 | 0 | 3 | 4 | 5 | |
| | 5+V[2,0]} | 4 | 0 | | | | | |
| | $= \max \{ 4, 5 + 0 \} = 5$ | L | 1 | I | | | | |
| 12 | W3 = 4, | | | | | | | |
| | Available knapsack capacity = 5 | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | W3 < WA, CASE 2 holds: | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | $V[i, j] = max \{ V[i-1, j],$ | 1 | 0 | 0 | 3 | 3 | 3 | 3 |
| | vi +V[i-1, j - wi] } | 2 | 0 | 0 | 3 | 4 | 4 | 7 |
| | $V[3,5] = \max \{ V[2,5],$ | 3 | 0 | 0 | 3 | 4 | 5 | 7 |
| | 5 + V[2, 1] | 4 | 0 | | | | | |
| | $= \max\{7, 5+0\} = 7$ | | | | | | | |
| 13 | W4 = 5. | | | | | | | |
| | Available knapsack capacity = | V[i,i] | i=0 | 1 | 2 | 3 | 4 | 5 |
| | 1,2,3,4 | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | W4 < WA, CASE 1 holds: | 1 | 0 | 0 | 3 | 3 | 3 | 3 |
| | V[i, j] = V[i-1, j] | 2 | 0 | 0 | 3 | 4 | 4 | 7 |
| | | 3 | 0 | 0 | 3 | 4 | 5 | 7 |
| | | 4 | 0 | 0 | 3 | 4 | 5 | |
| 14 | W4 = 5, | | | | | | | |
| | Available knapsack capacity = 5 | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | W4 = WA, CASE 2 holds: | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | $V[i, j] = max \{ V[i-1, j],$ | 1 | 0 | 0 | 3 | 3 | 3 | 3 |
| | vi +V[i-1, j - wi] } | 2 | 0 | 0 | 3 | 4 | 4 | 7 |
| | $V[4,5] = \max \{ V[3,5],$ | 3 | 0 | 0 | 3 | 4 | 5 | 7 |
| | 6 + V[3, 0] | 4 | 0 | 0 | 3 | 4 | 5 | 7 |
| | $- \max\{ 1, 0 + 0 \} = 1$ | | | | | | | |
| | Maximal valu | e is V [4 | I, 5]= | - 7/- | | | | |

What is the composition of the optimal subset?

The composition of the optimal subset if found by tracing back the computations for the entries in the table.

| Step | | |] | Table | | | | Remarks |
|------|----------|-------------------|----------------|------------|--------|--------|--------|------------------------------------|
| 1 | | | | | | | | |
| | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 | V[4,5] = V[3,5] |
| | i=0 | 0 | 0 | 0 | 0 | 0 | 0 | |
| | 1 | 0 | 0 | 3 | 3 | 3 | 3 | ITEM 4 NOT included in the |
| | 2 | 0 | 0 | 3 | 4 | 4 | 7 | subset |
| | 3 | 0 | 0 | 3 | 4 | 5 | 7 | |
| | 4 | 0 | 0 | 3 | 4 | 5 | | |
| 2 | | | | | | | | |
| | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 | V[3,5] = V[2,5] |
| | i=0 | 0 | 0 | 0 | 0 | 0 | 0 | |
| | 1 | 0 | 0 | 3 | 3 | 3 | 3 | ITEM 3 NOT included in the |
| | 2 | 0 | 0 | 3 | 4 | 4 | 7 | subset |
| | 3 | 0 | 0 | 3 | 4 | 5 | 7 | |
| | 4 | 0 | 0 | 3 | 4 | 5 | 7 | |
| 3 | | | | | | | | |
| | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 | V[2, 5] ≠ V[1, 5] |
| | i=0 | 0 | 0 | 0 | 0 | 0 | 0 | |
| | 1 | 0 | 0 | 3 | 3 | 3 | 3 | ITEM 2 included in the subset |
| | 2 | 0 | 0 | 3 | 4 | 4 | 7 | |
| | 3 | 0 | 0 | 3 | 4 | 5 | 7 | |
| | 4 | 0 | 0 | 3 | 4 | 5 | 7 | |
| 4 | Since it | em 2 i | s inclu | ıded i | n the | knaps | sack: | |
| | Weight | of | item | 2 is | 3kg, | , ther | efore, | |
| | remaini | ng cap | acity o | of the | knaps | ack is | | |
| | | (5 - 3 = | =) 2 kg | | | | | $V[1, 2] \neq V[0, 2]$ |
| | V[i i] | i-0 | 1 | ~ | 3 | 4 | 5 | ITEM 1 included in the subset |
| | i=0 | 0 | 0 | 0 | 0 | 0 | 0 | |
| | 1 | 0 | 0 | 3 | 3 | 3 | 3 | |
| | 2 | 0 | 0 | 3 | 4 | 4 | 7 | |
| | 3 | 0 | 0 | 3 | 4 | 5 | 7 | |
| | 4 | 0 | 0 | 3 | 4 | 5 | 7 | |
| 5 | Since it | em 1 ; | s inch | i hahi | n the | knong | sack. | Ontimal subset. ∫ item 1 item 2] |
| | Weight | of | item | 1 is | 2kg. | ther | efore. | Optimal subset. { Item 1, Item 2 } |
| | remaini | ng cap | acity of | of the | knaps: | ack is | , | Total weight is: 5kg (2kg + 3kg) |
| | | $(\tilde{2} - 2)$ | =) 0 kg | z . | T | | | Total profit is: 7/- (3/- + 4/-) |
| | | | - | | | | | |

Efficiency:

- Running time of Knapsack problem using dynamic programming algorithm is: O(n * W)
- Time needed to find the composition of an optimal solution is: $O(\ n+W\)$

Memory function

- Memory function combines the strength of top-down and bottom-up approaches
- It solves ONLY sub-problems that are necessary and does it ONLY ONCE.

The method:

- Uses top-down manner.
- Maintains table as in bottom-up approach.
- Initially, all the table entries are initialized with special "null" symbol to indicate that they have not yet been calculated.
- Whenever a new value needs to be calculated, the method checks the corresponding entry in the table first:
- If entry is NOT "null", it is simply retrieved from the table.
- Otherwise, it is computed by the recursive call whose result is then recorded in the table.

Algorithm:

Example:

Apply memory function method to the following instance of the knapsack problem Capacity W=5

| Item # | Weight (Kg) | Value (Rs.) |
|--------|-------------|-------------|
| 1 | 2 | 3 |
| 2 | 3 | 4 |
| 3 | 4 | 5 |
| 4 | 5 | 6 |

Solution:

Using memory function approach, we have:

| | Computation | | | Ren | narks | 5 | | |
|---|--|----------|-------|-----|-------|----|----|----|
| 1 | Initially, all the table entries are initialized | | | | | | | |
| | with special "null" symbol to indicate that | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | they have not yet been calculated. Here | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | null is indicated with -1 value. | 1 | 0 | -1 | -1 | -1 | -1 | -1 |
| | | 2 | 0 | -1 | -1 | -1 | -1 | -1 |
| | | 3 | 0 | -1 | -1 | -1 | -1 | -1 |
| | | 4 | 0 | -1 | -1 | -1 | -1 | -1 |
| 2 | MFKnap(4, 5) | | | | | | | |
| | | V[1, 5 |] = 3 | | | | | |
| | $\mathbf{MFKnap}(3,5) \qquad 6 + \mathbf{MFKnap}(3,0)$ | V[i,i] | i=0 | 1 | 2 | 3 | 4 | 5 |
| | \mathbf{k} | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | MFKnap $(2, 5)$ 5 + MFKnap $(2, 1)$ | 1 | 0 | -1 | -1 | -1 | -1 | 3 |
| | | 2 | 0 | -1 | -1 | -1 | -1 | -1 |
| | MFKnap(1, 5) 4 + MFKnap(1, 2) | 3 | 0 | -1 | -1 | -1 | -1 | -1 |
| | 0 3 | 4 | 0 | -1 | -1 | -1 | -1 | -1 |
| | | | _ | | | | | |
| | $MFKnap(0, 5) \qquad 3 + MFKnap(0, 3)$ | | | | | | | |
| | 0 3+0 | | | | | | | |
| 3 | MFKnap(4, 5) | | | | | | | |
| | | V[1, 2 |] = 3 | | | | | |
| | MFKnap(3,5) 6 + MFKnap(3,0) | | 1 | 1 | | | 1 | |
| | | V[i,j] | j=0 | 1 | 2 | 3 | 4 | 5 |
| | $MFKnap(2,5) \qquad 5 + MFKnap(2,1)$ | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | | 1 | 0 | -1 | 3 | -1 | -1 | 3 |
| | $MEK_{non}(1,5) \qquad \qquad 4 + MEK_{non}(1,2)$ | 2 | 0 | -1 | -1 | -1 | -1 | -1 |
| | $\frac{3}{2} \sim \frac{3}{2}$ | 3 | 0 | -1 | -1 | -1 | -1 | -1 |
| | | 4 | 0 | -1 | -1 | -1 | -1 | -1 |
| | MFKnap(0,2) 3 + MFKnap(0,0 |) | | | | | | |
| | 0 3+0 | | | | | | | |
| 4 | MFKnap(4, 5) | | | | | | | |
| | | V[2, 5 |] = 7 | | | | | |
| | $MFKnap(3,5) \qquad 6 + MFKnap(3,0)$ | V[i,i] | i=0 | 1 | 2 | 3 | 4 | 5 |
| | | i=0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | MFKnap(2,5) 5 + MFKnap(2,1) | 1 | 0 | -1 | 3 | -1 | -1 | 3 |
| | 3 7 | 2 | 0 | -1 | -1 | -1 | -1 | 7 |
| | $MEKnon(1.5) \qquad A + MEKnon(1.2)$ | 3 | 0 | -1 | -1 | -1 | -1 | -1 |
| | $\frac{1}{2} + \frac{1}{2} + \frac{1}$ | 4 | 0 | -1 | -1 | -1 | -1 | -1 |
| | 3 3 | <u> </u> | I | | 1 | 1 | I | |

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Efficiency:

- Time efficiency same as bottom up algorithm: O(n * W) + O(n + W)
- Just a constant factor gain by using memory function
- Less space efficient than a space efficient version of a bottom-up algorithm

UNIT-5

DECREASE-AND-CONQUER APPROACHES, SPACE-TIMETRADEOFFS

5.1DECREASE-AND-CONQUER APPROACHES: INTRODUCTION 5.2 INSERTION SORT 5.3 DEPTH FIRST SEARCH AND BREADTH FIRST SEARCH 5.4TOPOLOGICAL SORTING 5.5SPACE-TIME TRADEOFFS: INTRODUCTION 5.6SORTING BY COUNTING 5.7 INPUT ENHANCEMENT IN STRING MATCHING.

5.1 INTRODUCTION:

Decrease & conquer is a general algorithm design strategy based on exploiting the relationship between a solution to a given instance of a problem and a solution to a smaller instance of the same problem. The exploitation can be either top-down (recursive) or bottom-up (non-recursive).

The major variations of decrease and conquer are

- 1. Decrease by a constant :(usually by 1):
 - a. insertion sort
 - b. graph traversal algorithms (DFS and BFS)
 - c. topological sorting
 - d. algorithms for generating permutations, subsets
- 2. Decrease by a constant factor (usually by half)
 - a. binary search and bisection method
- 3. Variable size decrease
 - a. Euclid's algorithm

Decrease by a constant :(usually by 1):



Decrease by a constant factor (usually by half)



5.2 INSERTION SORT

Description:

Insertion sort is an application of decrease & conquer technique. It is a comparison based sort in which the sorted array is built on one entry at a time

$$A[0] \leq \ldots \leq A[j] < A[j+1] \leq \ldots \leq A[i-1] \mid A[i] \ldots A[n-1]$$

smaller than or equal to $A[i]$ greater than $A[i]$

Algorithm:

ALGORITHM Insertionsort(A [0 ... n-1])

//sorts a given array by insertion sort
//i/p: Array A[0...n-1]
//o/p: sorted array A[0...n-1] in ascending order

for i 1 to n-1 V A[i] j i-1

```
while j \ge 0 AND A[j] > V do

A[j+1] A[j]

j j-1

A[j+1] V
```

Analysis:

- Input size: Array size, n
- **Basic operation:** key comparison
- Best, worst, average case exists
 Best case: when input is a sorted array in ascending order:
 Worst case: when input is a sorted array in descending order:
- Let $C_{worst}(n)$ be the number of key comparison in the worst case. Then

$$C_{worst}(n) = \sum_{i=1}^{n-1} \sum_{j=0}^{i-1} 1 = \sum_{i=1}^{n-1} i = \frac{(n-1)n}{2} \in \Theta(n^2).$$

- Let $C_{\text{best}}(n)$ be the number of key comparison in the best case. Then

$$C_{best}(n) = \sum_{i=1}^{n-1} 1 = n - 1 \in \Theta(n).$$

Example:

Sort the following list of elements using insertion sort:

89, 45, 68, 90, 29, 34, 17

| 89 | 45 | 68 | 90 | 29 | 34 | 17 |
|----|----|----|----|----|----|----|
| 45 | 89 | 68 | 90 | 29 | 34 | 17 |
| 45 | 68 | 89 | 90 | 29 | 34 | 17 |
| 45 | 68 | 89 | 90 | 29 | 34 | 17 |
| 29 | 45 | 68 | 89 | 90 | 34 | 17 |
| 29 | 34 | 45 | 68 | 89 | 90 | 17 |
| 17 | 29 | 34 | 45 | 68 | 89 | 90 |

Advantages of insertion sort:

- Simple implementation. There are three variations
 - Left to right scan
 - Right to left scan
 - Binary insertion sort
- Efficient on small list of elements, on almost sorted list
- Running time is linear in best case
- Is a stable algorithm
- Is a in-place algorithm

5.3 DEPTH-FIRST SEARCH (DFS) AND BREADTH-FIRST SEARCH (BFS)

DFS and BFS are two graph traversing algorithms and follow decrease and conquer approach – decrease by one variation to traverse the graph

Some useful definition:

- Tree edges: edges used by DFS traversal to reach previously unvisited vertices
- **Back edges:** edges connecting vertices to previously visited vertices other than their immediate predecessor in the traversals
- **Cross edges:** edge that connects an unvisited vertex to vertex other than its immediate predecessor. (connects siblings)
- **DAG:** Directed acyclic graph

Depth-first search (DFS)

Description:

- DFS starts visiting vertices of a graph at an arbitrary vertex by marking it as visited.
- It visits graph's vertices by always moving away from last visited vertex to an unvisited one, backtracks if no adjacent unvisited vertex is available.
- Is a recursive algorithm, it uses a stack
- A vertex is pushed onto the stack when it's reached for the first time
- A vertex is popped off the stack when it becomes a dead end, i.e., when there is no adjacent unvisited vertex
- "Redraws" graph in tree-like fashion (with tree edges and back edges for undirected graph)

Algorithm:

ALGORITHM DFS (G)

//implements DFS traversal of a given graph
//i/p: Graph G = { V, E}
//o/p: DFS tree

Mark each vertex in V with 0 as a mark of being "unvisited" count 0for each vertex v in V do if v is marked with 0 dfs(v)

dfs(v)

count count + 1 mark v with count for each vertex w in V adjacent to v do if w is marked with 0 dfs(w)

Example:

Starting at vertex A traverse the following graph using DFS traversal method:



Solution:

| Step | Graph | Remarks |
|------|---|---------------------|
| 1 | | Insert A into stack |
| | | A(1) |
| | | |
| 2 | | Insert B into stack |
| | | |
| | | B (2) |
| | | A(1) |
| 3 | A B | Insert F into stack |
| | | hisort i mto stuck |
| | | F (3) |
| | (F) | B (2) |
| L | | A(1) |
| 4 | | Insert E into steek |
| | (A) (B) | Insert E into stack |
| | | E (4) |
| | | F (3) |
| | (E)(F) | B (2) |
| - | | A(1) |
| 5 | NO unvisited adjacent vertex for E, backfrack | Delete E from stack |
| | | E(4, 1) |
| | | $ \mathbf{F}(3) $ |
| | | B (2) |
| | | A(1) |

| 6 | NO unvisited adjacent vertex for F, backtrack | Delete F from stack |
|----|---|----------------------------------|
| | | E (4, 1) F (3, 2) |
| | | B (2) A(1) |
| 7 | | Insert G into stack |
| | | |
| | | F(4, 1) F(3, 2) G(5) |
| | (E) (F) (G) | B (2) A(1) |
| 8 | | Insert C into stack |
| | | E(4, 1) C(6) E(3, 2) G(5) |
| | E F C | B (2) A(1) |
| 9 | \bigcirc \bigcirc \bigcirc \bigcirc | Insert D into stack |
| | | D (7) |
| | | E (4, 1) C (6) |
| | | F (3, 2) G (5) B (2) |
| 10 | | A(1) |
| | | Insert H into stack H (8) |
| | | D (7) E (4, 1) C (6) |
| | | F(3,2) = G(5) F(2) |
| | | A(1) |
| | NO unvisited adjacent vertex for H, backtrack | Delete H from stack |
| | | H (8, 3) D (7) |
| | | E (4, 1) C (6) F (3, 2) G (5) |
| | | B (2) |
| L | | A(1) |

| 12 | NO unvisited adjacent vertex for D, backtrack | Delete D from stack |
|----|---|---------------------|
| | | H (8, 3) |
| | | D (7, 4) |
| | | E (4, 1) C (6) |
| | | F (3, 2) G (5) |
| | | B (2) |
| | | A(1) |
| 13 | NO unvisited adjacent vertex for C, backtrack | Delete C from stack |
| | | H (8, 3) |
| | | D (7, 4) |
| | | E (4, 1) C (6, 5) |
| | | F (3, 2) G (5) |
| | | B (2) |
| | | A(1) |
| 14 | NO unvisited adjacent vertex for G, backtrack | Delete G from stack |
| | | H (8, 3) |
| | | D (7, 4) |
| | | E (4, 1) C (6, 5) |
| | | F (3, 2) G (5, 6) |
| | | B (2) |
| | | A(1) |
| 15 | NO unvisited adjacent vertex for B, backtrack | Delete B from stack |
| | | H (8, 3) |
| | | D (7, 4) |
| | | E (4, 1) C (6, 5) |
| | | F (3, 2) G (5, 6) |
| | | B (2, 7) |
| | | A(1) |
| 16 | NO unvisited adjacent vertex for A, backtrack | Delete A from stack |
| | | H (8, 3) |
| | | D (7, 4) |
| | | E (4, 1) C (6, 5) |
| | | F (3, 2) G (5, 6) |
| | | B (2, 7) |
| | | A(1, 8) |
| | Stack becomes empty. Algorithm stops as all the | |
| | nodes in the given graph are visited | |

The DFS tree is as follows: (dotted lines are back edges)



Applications of DFS:

- The two orderings are advantageous for various applications like topological sorting, etc
- To check connectivity of a graph (number of times stack becomes empty tells the number of components in the graph)
- To check if a graph is acyclic. (no back edges indicates no cycle)
- To find articulation point in a graph

Efficiency:

- Depends on the graph representation:
 - Adjacency matrix : $\Theta(n^2)$
 - Adjacency list: $\Theta(n + e)$

Breadth-first search (BFS) Description:

- BFS starts visiting vertices of a graph at an arbitrary vertex by marking it as visited.
- It visits graph's vertices by across to all the neighbors of the last visited vertex
- Instead of a stack, BFS uses a queue
- Similar to level-by-level tree traversal
- "Redraws" graph in tree-like fashion (with tree edges and cross edges for undirected graph)

Algorithm:

ALGORITHM BFS (G)

//implements BFS traversal of a given graph
//i/p: Graph G = { V, E}
//o/p: BFS tree/forest
Mark each vertex in V with 0 as a mark of being "unvisited"

add w to the queue

remove vertex v from the front of the queue

Example:

Starting at vertex A traverse the following graph using BFS traversal method:



Solution:

| Step | Graph | Remarks |
|------|--------------|--|
| 1 | A | Insert A into queue A(1) |
| 2 | A B E | Insert B, E into queue A(1), B (2), E(3) B (2), E(3) |
| 3 | A B E F G | Insert F, G into queue B(2), E(3), F(3), G(4) E(3), F(3), G(4) |

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| 4 | NO unvisited adjacent vertex for E, backtrack | Delete E from queue |
|---|--|--------------------------------|
| | | F(3), G(4) |
| 5 | NO unvisited adjacent vertex for F, backtrack | Delete F from queue |
| | | G(4) |
| 6 | | Insert C, H into queue |
| | | G(4), C(5), H(6) |
| | (E) (F) (G) (H) | C(5), H(6) |
| 7 | | Insert D into queue |
| | E F G H | C(5), H(6), D(7) H(6), D(7) |
| | NO unvisited adjacent vertex for H, backtrack | Delete H from queue |
| 8 | | D(7) |
| 9 | NO unvisited adjacent vertex for D, backtrack | Delete D from queue |
| | Queue becomes empty. Algorithm stops as all the nodes in the given graph are visited | |

The BFS tree is as follows: (dotted lines are cross edges)



Applications of BFS:

- To check connectivity of a graph (number of times queue becomes empty tells the number of components in the graph)
- To check if a graph is acyclic. (no cross edges indicates no cycle)
- To find minimum edge path in a graph

Efficiency:

- Depends on the graph representation:
 - Array : $\Theta(n^2)$
 - List: $\Theta(n+e)$

Difference between DFS & BFS:

| | DFS | BFS |
|-------------------------|---------------------|--------------------|
| Data structure | Stack | Queue |
| No. of vertex orderings | 2 orderings | 1 ordering |
| Edge types | Tree edge | Tree edge |
| | Back edge | Cross edge |
| Applications | Connectivity | Connectivity |
| | Acyclicity | Acyclicity |
| | Articulation points | Minimum edge paths |
| Efficiency for | $\Theta(n^2)$ | $\Theta(n^2)$ |
| adjacency matrix | | |
| Efficiency for | $\Theta(n+e)$ | $\Theta(n+e)$ |
| adjacency lists | | |

5.4 Topological Sorting

Description:

Topological sorting is a sorting method to list the vertices of the graph in such an order that for every edge in the graph, the vertex where the edge starts is listed before the vertex where the edge ends.

NOTE: There is no solution for topological sorting if there is a cycle in the digraph . [MUST be a DAG]

Topological sorting problem can be solved by using

- 1. DFS method
- 2. Source removal method

DFS Method:

- Perform DFS traversal and note the order in which vertices become dead ends (popped order)
- Reverse the order, yield the topological sorting.

Example:

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Apply DFS – based algorithm to solve the topological sorting problem for the given graph:



| Step | Graph | Remarks |
|------|-----------|----------------------|
| 1 | | Insert C1 into stack |
| | C1 | C1(1) |
| 2 | | Insert C2 into stack |
| | 63 | C2 (2) |
| | | C1(1) |
| 3 | C4 | Insert C4 into stack |
| | | C4 (3) |
| | | C2 (2) |
| |) | C1(1) |
| 4 | C4 | Insert C5 into stack |
| | | C5 (4) |
| | | C4 (3) |
| | | C2(2) |
| | | CI(1) |

| 5 | NO unvisited adjacent vertex for C5, backtrack | Delete C5 from stack |
|---|--|----------------------|
| | | |
| | | C5 (4, 1) |
| | | C4 (3) |
| | | C2 (2) |
| | | C1(1) |
| 6 | NO unvisited adjacent vertex for C4, backtrack | Delete C4 from stack |
| | | |
| | | C5 (4, 1) |
| | | C4 (3, 2) |
| | | C2 (2) |
| | | C1(1) |

| 7 | NO unvisited adjacent vertex for C3, backtrack | Delete C3 from stack |
|--------------|--|---|
| | | C5 (4, 1) |
| | | C4 (3,2) |
| | | C2(2,3) |
| | | C1(1) |
| 8 | NO unvisited adjacent vertex for C1, backtrack | Delete C1 from stack |
| | | C5(4,1) |
| | | $C_{3}(4, 1)$ |
| | | C4(3,2) |
| | | $C_2(2,3)$ |
| 0, 11 | | |
| Stack b | ecomes empty, but there is a node which is unvisited com arbitrarily selecting a unvisited node as source | , therefore start the DFS |
| again n Q | on arbitrarity selecting a unvisited node as source | Insert C2 into stack |
| | $\overline{(2)}$ | msert C2 mto stack |
| | CZ | C5(4, 1) |
| | | $C_{4}(3, 1)$ |
| | | $C_{4}(3,2)$ |
| | | $C_{2}(2, 3)$ $C_{1}(1, 4)$ $C_{2}(5)$ |
| | | C1(1, 4) $C2(3)$ |
| 10 | NO unvisited adjacent vertex for C2, backtrack | Delete C2 from stack |
| | | C5 (4, 1) |
| | | C4 (3,2) |
| | | $C_{2}(2,3)$ |
| | | C1(1, 4) $C2(5, 5)$ |
| Stack b | ecomes empty, NO unvisited node left, therefore algo | orithm stops. |
| The por | oping – off order is: | L |
| C5, C4 | C3, C1, C2, | |
| Topolo | gically sorted list (reverse of pop order): | |
| C^2 C^1 | $C_3 C_4 C_5$ | |

Source removal method:

- Purely based on decrease & conquer
- Repeatedly identify in a remaining digraph a source, which is a vertex with no incoming edges
- Delete it along with all the edges outgoing from it.

Example:

Apply Source removal – based algorithm to solve the topological sorting problem for the given graph:



Solution:



The topological order is C1, C2, C3, C4, C5

5.5 SPACE-TIME TRADEOFFS:

Introduction

Two varieties of space-for-time algorithms:

- <u>input enhancement</u> preprocess the input (or its part) to store some info to be used later in solving the problem
 - counting sorts
 - string searching algorithms
- <u>pre structuring</u> preprocess the input to make accessing its elements easier 1)hashing 2)indexing schemes (e.g., B-trees)

5.6 SORTING BY COUNTING

Assume elements to be sorted belong to a known set of small values between l and u, with potential duplication

Constraint: we cannot overwrite the original list Distribution Counting: compute the frequency of each

element and later accumulate sum of frequencies (distribution)

Algorithm:

```
for j \leftarrow 0 to u-1 do D[j] \leftarrow 0 // init frequencies
for i \leftarrow 0 to n-1 do D[A[i]-1] \leftarrow D[A[i] - 1] + 1 // compute frequencies
for j \leftarrow 1 to u-1 do D[j] \leftarrow D[j-1] + D[j] // reuse for distribution
for i \leftarrow n-1 downto 0 do
j \leftarrow A[i] - 1
S[D[j] - 1] \leftarrow A[i]
D[j] \leftarrow D[j] - 1
return S
```

| 100 | 11 | 12 | 13 | 12 | 2 12 | ā . | | | |
|--------|---|--|---|--|---|---|---|--|--|
| Arra | y Valu | es | 11 | 12 | 13 | | | | |
| Freq | Juenci | əs | 1 | 3 | 2 | | | | |
| Distri | bution | | 1 | 4 | 6 | | | | |
| D[0] | D[1] | D[| 2] | S[0] | S[1] | S[2] | S[3] | S[4] | S[5] |
| 1 | 4 | | 6 | | | | 12 | | |
| 12 | 3 | | 6 | | 4 | 12 | 4 | a. | |
| 12 | 2 | | 6 | | | | | | 13 |
| 13 | 2 | | 5 | | 12 | | | | |
| 12 | 1 | | 5 | 11 | | | | | |
| 10 | 1 | | 5 | | | | | 13 | |
| | Arra Frea Distri D[0] 1 12 13 13 14 13 14 13 14 13 13 13 13 13 13 13 14 13 14 14 14 14 14 14 14 14 14 14 | Array Valu Frequencie Distribution D[0] D[1] 1 4 12 3 12 1 13 1 10 1 | Array Values Frequencies Distribution D[0] D[1] D[0] D[1] 1 4 12 1 13 2 14 2 13 1 14 1 15 1 | Array Values 11 Frequencies 1 Distribution 1 D[0] D[1] D[2] 1 4 6 12 1 3 6 12 2 6 13 13 2 5 14 10 1 5 13 | Array Values 11 12 Frequencies 1 3 Distribution 1 4 D[0] D[1] D[2] S[0] 1 4 6 12 6 13 6 12 6 13 6 14 6 15 11 16 11 17 2 5 18 11 19 1 5 13 6 | Array Values 11 12 13 Frequencies 1 3 2 Distribution 1 4 6 D[0] D[1] D[2] S[0] S[1] 1 4 6 - 12 3 6 - 12 3 6 - 12 6 - - 13 6 - - 12 5 12 - 13 5 11 - 14 5 11 - | Array Values 11 12 13 Frequencies 1 3 2 Distribution 1 4 6 D[0] D[1] D[2] S[0] S[1] S[2] 1 4 6 12 12 1 4 6 12 12 1 3 6 12 12 12 5 12 12 13 1 5 11 12 13 6 12 12 12 13 1 5 11 12 14 5 11 11 12 14 5 11 12 13 | Array Values 11 12 13 Frequencies 1 3 2 Distribution 1 4 6 D[0] D[1] D[2] S[0] S[1] S[2] S[3] 1 4 6 12 12 12 1 3 6 12 12 1 3 6 12 12 1 3 6 12 13 6 12 12 12 13 6 12 12 12 13 1 5 11 11 14 1 5 11 11 14 1 5 11 12 14 1 5 11 11 15 11 11 11 11 | Array Values 11 12 13 Frequencies 1 3 2 Distribution 1 4 6 D[0] D[1] D[2] S[0] S[1] S[2] S[3] S[4] 1 4 6 12 12 1 4 6 12 12 1 3 6 12 12 1 3 6 12 12 12 5 12 12 12 13 6 12 12 13 12 5 12 13 13 6 11 13 |

Efficiency: $\Theta(n)$

Best so far but only for specific types of input

5.7 INPUT ENHANCEMENT IN STRING MATCHING.

Horspool's Algorithm

A simplified version of Boyer-Moore algorithm: preprocesses pattern to generate a shift table that determines how much to shift the patter when a mismatch occurs. Always makes a shift based on the text's character c aligned with the <u>last</u> compared (mismatched) character in the pattern according to the shift table's entry for c

```
void horspoolInitocc()
{
    int j;
    char a;
    for (a=0; a<alphabetsize; a++)
        occ[a]=-1;
    for (j=0; j<m-1; j++)
    {
        a=p[j];
        occ[a]=j;
    }
}</pre>
```

```
}
void horspoolSearch()
{
    int i=0, j;
    while (i<=n-m)
    {
        j=m-1;
        while (j>=0 && p[j]==t[i+j]) j--;
        if (j<0) report(i);
        i+=m-1;
        i-=occ[t[i]];
    }
}</pre>
```

Shift table

Shift sizes can be precomputed by the formula

distance from <u>c's</u> rightmost occurrence in pattern among its first <u>m-1</u> characters to its right end



pattern's length m, otherwise

by scanning pattern before search begins and stored in a table called *shift table*. After the shift, the right end of pattern is *t(c)* positions to the right of the last compared character in text.

 Shift table is indexed by text and pattern alphabet Eg, for BAOBAB:

| A | в | С | D | E | F | G | н | I | J | ĸ | L | м | N | 0 | Р | Q | R | s | т | υ | v | w | x | Y | z |
|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| 1 | 2 | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 | з | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 | 6 |

Example of Horspool's algorithm



Time complexity

- preprocessing phase in $O(m + \pi)$ time and $O(\pi)$ space complexity.
- searching phase in O(mn) time complexity.
- the average number of comparisons for one text character is between $1/\pi$ and $2/(\pi+1)$.

(π is the number of storing characters)

UNIT 6

LIMITATIONS OF ALGORITHMIC POWER AND COPING WITH THEM

6.1LOWER-BOUND ARGUMENTS

6.2DECISION TREES

6.3 P, NP, AND NP-COMPLETE PROBLEMS

Objectives

We now move into the third and final major theme for this course.

- 1. *Tools* for analyzing algorithms.
- 2. Design strategies for designing algorithms.
- 3. Identifying and coping with the *limitations* of algorithms.

Efficiency of an algorithm

• By establishing the asymptotic efficiency class

| Problem | Algorithm | Time efficiency |
|-----------------|----------------|-----------------|
| sorting | selection sort | $\Theta(n^2)$ |
| Towers of Hanoi | recursive | $\Theta(2^n)$ |

- The efficiency class for selection sort (quadratic) is lower. Does this mean that selection sort is a "better" algorithm?
 - Like comparing "apples" to "oranges"
- By analyzing how efficient a particular algorithm is compared to other algorithms for the same problem
 - It is desirable to know the best possible efficiency any algorithm solving This problem may have – establishing a *lower bound*

6.1 LOWER BOUNDS ARGUMENTS

Lower bound: an estimate on a minimum amount of work needed to solve a given problem Examples:

- number of comparisons needed to find the largest element in a set of *n* numbers
- number of comparisons needed to sort an array of size *n*
- number of comparisons necessary for searching in a sorted array
- number of multiplications needed to multiply two *n*-by-*n* matrices
- Lower bound can be
 - an exact count
 - an efficiency class (Ω)
- <u>*Tight*</u> lower bound: there exists an algorithm with the same efficiency as the lower bound

| Problem | Lower bound | | Tightness |
|---|--------------------|-----|-----------|
| sorting | $\Omega(n \log n)$ | yes | |
| searching in a sorted array | $\Omega(\log n)$ | | yes |
| element uniqueness | $\Omega(n \log n)$ | | yes |
| <i>n</i> -digit integer multiplication | $\Omega(n)$ | | unknown |
| multiplication of <i>n</i> -by- <i>n</i> matrices | $\Omega(n2)$ | | unknown |

Methods for Establishing Lower Bounds

- Trivial lower bounds
- Information-theoretic arguments (decision trees)
- Adversary arguments
- Problem reduction

Trivial Lower Bounds

<u>*Trivial lower bounds*</u>: based on counting the number of items that must be processed in input and generated as output

Examples

- finding max element
- polynomial evaluation
- sorting
- element uniqueness
- computing the product of two n-by-n matrices

Conclusions

- may and may not be useful
- be careful in deciding how many elements <u>must</u> be processed

6.2 DECISION TREES

<u>Decision tree</u> — a convenient model of algorithms involving Comparisons in which: (i) internal nodes represent comparisons (ii) leaves represent outcomes Decision tree for 3-element insertion sort

Decision tree for 3-element insertion sort



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- How does such a tree help us find lower bounds?
 - There must be at least one leaf for each correct output.
 - The tree must be tall enough to have that many leaves.
- In a binary tree with l leaves and height h,

 $h \geq \log_2 l$

Decision Tree and Sorting Algorithms



FIGURE 11.2 Decision tree for the three-element selection sort. A triple above a node indicates the state of the array being sorted. Note the two redundant comparisons b < a with a single possible outcome because of the results of some previously made comparisons.

Decision Tree and Sorting Algorithms

- Number of leaves (outcomes) $\geq n!$
- Height of binary tree with n! leaves $\geq \bigotimes \log_2 n! \diamondsuit$
- Minimum number of comparisons in the worst case $\geq \langle \mathbf{Q} \log_2 n | \mathbf{Q} \rangle$ for any comparison-based sorting algorithm
- $\partial \log_2 n! \partial \approx n \log_2 n$
- This lower bound is tight (e.g. mergesort)

Decision Tree and Searching a Sorted Array



FIGURE 11.4 Ternary decision tree for binary search in a four-element array

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Decision Tree and Searching a Sorted Array

- Number of leaves (outcomes) = n + n + 1 = 2n + 1
- Height of ternary tree with 2n+1 leaves $\geq \bigotimes \log_3 (2n+1) \bigotimes$
- This lower bound is NOT tight (the number of worst-case comparisons for binary search is \$log₂ (n+1)\$, and \$log₃ (2n+1)\$ ≤ \$log₂ (n+1)\$)
- Can we find a better lower bound or find an algorithm with better efficiency than binary search?

Decision Tree and Searching a Sorted Array



FIGURE 11.5 Binary decision tree for binary search in a four-element array

Decision Tree and Searching a Sorted Array

- Consider using a binary tree where internal nodes also serve as successful searches and leaves only represent unsuccessful searches
- Number of leaves (outcomes) = n + 1
- Height of binary tree with n+1 leaves $\geq \bigotimes \log_2(n+1) \bigotimes$
- This lower bound is tight

Decision-tree example





Each internal node is labeled i:j for $i, j \in \{1, 2, ..., n\}$.

- The left subtree shows subsequent comparisons if $a_i \le a_j$.
- The right subtree shows subsequent comparisons if $a_i \ge a_i$.

Each internal node is labeled i : j for $i, j \in \{1, 2, ..., n\}$.

• The left subtree shows subsequent comparisons if $a_i \leq a_j$.

• The right subtree shows subsequent comparisons if $a_i \ge a_j$.





Each internal node is labeled i j for $i, j \in \{1, 2, ..., n\}$.

• The left subtree shows subsequent comparisons if $a_i \leq a_j$.

• The right subtree shows subsequent comparisons if $a_i \ge a_j$.



If $a_i \ge a_j$. The right subtree shows subsequent comparis



Each leaf contains a permutation $\langle \pi(1), \pi(2), \dots, \pi(n) \rangle$ to indicate that the ordering $a_{\pi(1)} \leq a_{\pi(2)} \leq \dots \leq a_{\pi(n)}$ has been established.

Decision-tree model

A decision tree can model the execution of any comparison sort:

- One tree for each input size *n*.
- View the algorithm as splitting whenever it compares two elements.
- The tree contains the comparisons along all possible instruction traces.
- The running time of the algorithm = the length of the path taken.
- Worst-case running time = height of tree.

Any comparison sort can be turned into a Decision tree



Decision Tree Model : insertion sort

 For example, we trace the sequence of key comparisons of insertion sort on <a1, a2, a3> by a decision tree.





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Decision Tree Model

- In the insertion sort example, the decision tree reveals all possible key comparison sequences for 3 distinct numbers.
- There are exactly 3!=6 possible output sequences.
- Different comparison sorts should generate different decision trees.
- It should be clear that, in theory, we should be able to draw a decision tree for ANY comparison sort algorithm.
- Given a particular input sequence, the path from root to the leaf path traces a particular key comparison sequence performed by that comparison sort.

- The length of that path represented the number of key comparisons performed by the sorting algorithm.

- When we come to a leaf, the sorting algorithm has determined the sorted order.
- Notice that a correct sorting algorithm should be able to sort EVERY possible output sorted order.
- Since, there are n! possible sorted order, there are n! leaves in the decision tree.
- Given a decision tree, the height of the tree represent the longest length of a root to leaf path.
- It follows the height of the decision tree represents the largest number of key comparisons, which is the worst-case running time of the sorting algorithm.

"Any comparison based sorting algorithm takes $\Omega(n \log n)$ to sort a list of n distinct elements in the worst-case."

- any comparison sort \leftarrow model by a decision tree
- worst-case running time \leftarrow the height of decision tree

"Any comparison based sorting algorithm takes $\Omega(n \log n)$ to sort a list of n distinct elements in the worst-case."

- We want to find a lower bound (Ω) of the **height** of a binary tree that has n! Leaves.
 - What is the minimum height of a binary tree that has n! leaves?

- The binary tree must be a complete tree (recall the definition of complete tree).
- Hence the minimum (lower bound) height is $\theta(\log_2(n!))$.
- $\log_2(n!)$ = $\log_2(n) + \log_2(n-1) + ... + \log_2(n/2) + ...$ $\ge n/2 \log_2(n/2) = n/2 \log_2(n) - n/2$ So, $\log_2(n!) = \Omega(n \log n)$.
- It follows the height of a binary tree which has n! leaves is at least $\Omega(n \log n) \diamond$ worstcase running time is at least $\Omega(n \log n)$
- Putting everything together, we have "Any comparison based sorting algorithm takes $\Omega(n \log n)$ to sort a list of n distinct elements in the worst-case."

Adversary Arguments

<u>Adversary argument</u>: a method of proving a lower bound by playing role of adversary that makes algorithm work the hardest by adjusting input

Example: "Guessing" a number between 1 and *n* with yes/no questions

Adversary: Puts the number in a larger of the two subsets generated by last question

Lower Bounds by Problem Reduction

Idea: If problem P is at least as hard as problem Q, then a lower bound for Q is also a lower bound for P.

Hence, find problem Q with a known lower bound that can be reduced to problem P in question.

Example: Euclidean MST problem

- Given a set of n points in the plane, construct a tree with minimum total length that connects the given points. (considered as problem P)
- To get a lower bound for this problem, reduce the *element uniqueness problem* to it. (considered as problem Q)
- If an algorithm faster than $n \log n$ exists for Euclidean MST, then one exists for element uniqueness also. Aha! A contradiction! Therefore, any algorithm for Euclidean MST must take $\Omega(n \log n)$ time.

Classifying Problem Complexity

Is the problem *tractable*, i.e., is there a polynomial-time (O(p(n)) algorithm that solves it? Possible answers:

- yes (give examples)
- no
 - -because it's been proved that no algorithm exists at all (e.g., Turing's *halting problem*)
 - -because it's been proved that any algorithm for the problem takes exponential time unknown

Problem Types: Optimization and Decision

- <u>Optimization problem</u>: find a solution that maximizes or minimizes some objective function
- <u>Decision problem</u>: answer yes/no to a question Many problems have decision and optimization versions. E.g.: traveling salesman problem
- optimization: find Hamiltonian cycle of minimum length
- *decision*: find Hamiltonian cycle of length $\leq m$

Decision problems are more convenient for formal investigation of their complexity.

6.3 CLASS P

<u>*P*</u>: the class of decision problems that are solvable in O(p(n)) time, where p(n) is a polynomial of problem's input size n

Examples:

- searching
- element uniqueness
- graph connectivity
- graph acyclicity
- primality testing (finally proved in 2002)

6.4 CLASS NP

<u>*NP*</u> (*nondeterministic polynomial*): class of decision problems whose proposed solutions can be verified in polynomial time = solvable by a *nondeterministic polynomial algorithm*

A *nondeterministic polynomial algorithm* is an abstract two-stage procedure that:

- generates a random string purported to solve the problem
- checks whether this solution is correct in polynomial time

By definition, it solves the problem if it's capable of generating and verifying a solution on one of its tries

Why this definition?

• led to development of the rich theory called "computational complexity"

Example: CNF satisfiability

Problem: Is a boolean expression in its conjunctive normal form (CNF) satisfiable, i.e., are there values of its variables that makes it true?

This problem is in NP. Nondeterministic algorithm:

- Guess truth assignment
- Substitute the values into the CNF formula to see if it evaluates to true

Example: $(A | \neg B | \neg C) \& (A | B) \& (\neg B | \neg D | E) \& (\neg D | \neg E)$ Truth assignments: <u>A B C D E</u> 0 0 0 0 0 . . . 1 1 1 1 1

Checking phase: O(n)

What problems are in NP?

- Hamiltonian circuit existence
- Partition problem: Is it possible to partition a set of *n* integers into two disjoint subsets with the same sum?
- Decision versions of TSP, knapsack problem, graph coloring, and many other combinatorial optimization problems. (Few exceptions include: MST, shortest paths)
- All the problems in *P* can also be solved in this manner (but no guessing is necessary), so we have:

 $P \diamondsuit NP$

• Big question: P = NP ?

P = NP?

- One of the most important unsolved problems is computer science is whether or not P=NP.
 - If P=NP, then a ridiculous number of problems currently believed to be very difficult will turn out have efficient algorithms.
 - If $P \neq NP$, then those problems definitely do not have polynomial time solutions.
- Most computer scientists suspect that $P \neq NP$. These suspicions are based partly on the idea of NP-completeness.

6.5 NP-COMPLETE PROBLEMS

A decision problem D is <u>NP-complete</u> if it's as hard as any problem in NP, i.e.,

- D is in NP
- every problem in NP is polynomial-time reducible to D



Other *NP*-complete problems obtained through polynomialtime reductions from a known *NP*-complete problem



Examples: TSP, knapsack, partition, graph-coloring and hundreds of other problems of combinatorial nature

General Definitions: P, NP, NP-hard, NP-easy, and NP-complete

- Problems
- Decision problems (yes/no)
- Optimization problems (solution with best score)
- P
- Decision problems (decision problems) that can be solved in polynomial time can be solved "efficiently"
- NP
- Decision problems whose "*YES*" answer can be verified in polynomial time, if we already have the *proof* (or *witness*)
- co-NP
- Decision problems whose "NO" answer can be verified in polynomial time, if we already have the *proof* (or *witness*)
- e.g. The satisfiability problem (SAT)
- Given a boolean formula

$$(x_1 \diamondsuit x_2 \diamondsuit x_3 \diamondsuit x_4) \diamondsuit (x_5 \diamondsuit x_6 \diamondsuit x_7) \diamondsuit x_8 \diamondsuit x_9^-$$

is it possible to assign the input x1...x9, so that the formula evaluates to TRUE?

- If the answer is YES with a proof (i.e. an assignment of input value), then we can check the proof in polynomial time (SAT is in NP)

- We may not be able to check the NO answer in polynomial time (Nobody really knows.)

- NP-hard
 - A problem is NP-hard iff an polynomial-time algorithm for it implies a polynomial-time algorithm for every problem in NP
 - NP-hard problems are at least as hard as NP problems
- NP-complete
 - A problem is NP-complete if it is NP-hard, and is an element of NP (NP-easy)

- Relationship between decision problems and optimization problems
 - every optimization problem has a corresponding decision problem
 - optimization: minimize x, subject to constraints
 - yes/no: is there a solution, such that x is less than c?
 - an optimization problem is NP-hard (NP-complete)

if its corresponding decision problem is NP-hard (NP-complete)

• How to know another problem, A, is NP-complete?

- To prove that A is NP-complete, reduce a known NP-complete problem to A



• Requirement for Reduction

- Polynomial time
- YES to A also implies YES to SAT, while
- NO to A also implies No to SAT

Examples of NP-complete problems

Vertex cover

- Vertex cover
 - given a graph G=(V,E), find the *smallest* number of vertexes that cover *each edge*
 - Decision problem: is the graph has a vertex cover of size K?
- Independent set
- independent set: a set of vertices in the graph with no edges between *each pair* of nodes.
- given a graph G=(V,E), find the *largest independent set*
- reduction from vertex cover:
 - Set cover
 - given a universal set U, and several subsets S1,...Sn
 - find the least number of subsets that contains each elements in the universal set

Polynomial (P) Problems

- Are solvable in polynomial time
- Are solvable in O(nk), where k is some constant.
- Most of the algorithms we have covered are in P

Nondeterministic Polynomial (NP) Problems

- This class of problems has solutions that are verifiable in polynomial time.
 - Thus any problem in P is also NP, since we would be able to solve it in polynomial time, we can also verify it in polynomial time

NP-Complete Problems

- Is an NP-Problem
- Is at least as difficult as an NP problem (is reducible to it)
- More formally, a decision problem C is NP-Complete if:
 - C is in NP
 - Any known NP-hard (or complete) problem $\leq p C$
 - Thus a proof must show these two being satisfied

Exponential Time Algorithms



Examples

- Longest path problem: (similar to Shortest path problem, which requires polynomial time) suspected to require exponential time, since there is no known polynomial algorithm.
- Hamiltonian Cycle problem: Traverses all vertices exactly once and form a cycle.

Reduction

- P1 : is an unknown problem (easy/hard ?)
- P2 : is known to be difficult

If we can easily solve P2 using P1 as a subroutine then P1 is difficult Must create the inputs for P1 in polynomial time.

* P1 is definitely difficult because you know you cannot solve P2 in polynomial time unless you use a component that is also difficult (it cannot be the mapping since the mapping is known to be polynomial)

Decision Problems

Represent problem as a decision with a boolean output

- Easier to solve when comparing to other problems
- Hence all problems are converted to decision problems.
- P = {all decision problems that can be solved in polynomial time}

NP = {all decision problems where a solution is proposed, can be verified in polynomial time} NP-complete: the subset of NP which are the "hardest problems"

Alternative Representation

- Every element p in P1 can map to an element q in P2 such that p is true (decision problem) if and only if q is also true.
- Must find a mapping for such true elements in P1 and P2, as well as for false elements.
- Ensure that mapping can be done in polynomial time.
- *Note: P1 is unknown, P2 is difficult

Cook's Theorem

• Stephen Cook (Turing award winner) found the first NP-Complete problem, 3SAT. Basically a problem from Logic.

Generally described using Boolean formula.

A Boolean formula involves AND, OR, NOT operators and some variables.

Ex: (x or y) and (x or z), where x, y, z are boolean variables.

Problem Definition – Given a boolean formula of m clauses, each containing 'n' boolean variables, can you assign some values to these variables so that the formula can be true?

Boolean formula: $(x v y v \hat{z}) \Lambda (x v y v \hat{z})$

Try all sets of solutions. Thus we have exponential set of possible solutions. So it is a NPC problem.

• Having one definite NP-Complete problem means others can also be proven NP-Complete, using reduction.

Unit 7

COPING WITH LIMITATIONS OF ALGORITHMIC POWER

- 7.1 Backtracking: n Queens problem,
- 7.2 Hamiltonian Circuit Problem,
- 7.3 Subset –Sum Problem.
- 7.4 Branch-and-Bound: Assignment Problem,
- 7.5 Knapsack Problem,
- 7.6 Traveling Salesperson Problem.
- 7.7 Approximation Algorithms for NP-Hard Problems Traveling Salesperson Problem, Knapsack Problem

Introduction

Tackling Difficult Combinatorial Problems

• There are two principal approaches to tackling difficult combinatorial problems (NP-hard problems):

• Use a strategy that guarantees solving the problem exactly but doesn't guarantee to find a solution in polynomial time

• Use an approximation algorithm that can find an approximate (sub-optimal) solution in polynomial time

Exact Solution Strategies

- *exhaustive search* (brute force)
 - useful only for small instances
- dynamic programming
 - applicable to some problems (e.g., the knapsack problem)
- backtracking
 - eliminates some unnecessary cases from consideration
 - yields solutions in reasonable time for many instances but worst case is still exponential
- branch-and-bound
 - further refines the backtracking idea for optimization problems

7.1 Backtracking

- Suppose you have to make a series of *decisions*, among various *choices*, where
 - You don't have enough information to know what to choose
 - Each decision leads to a new set of choices
 - Some sequence of choices (possibly more than one) may be a solution to your problem
- Backtracking is a methodical way of trying out various sequences of decisions, until you find one that "works"

Backtracking : A Scenario



A tree is composed of nodes

Backtracking can be thought of as searching a tree for a particular "goal" leaf node

- Each non-leaf node in a tree is a parent of one or more other nodes (its children)
- Each node in the tree, other than the root, has exactly one parent

The backtracking algorithm

- Backtracking is really quite simple--we "explore" each node, as follows:
- To "explore" node N:
 - 1. If N is a goal node, return "success"
 - 2. If N is a leaf node, return "failure"
 - 3. For each child C of N,
 - 3.1. Explore C
 - 3.1.1. If C was successful, return "success"
 - 4. Return "failure"
- Construct the <u>state-space tree</u>
 - nodes: partial solutions
 - edges: choices in extending partial solutions
- Explore the state space tree using depth-first search
- "Prune" nonpromising nodes
 - dfs stops exploring subtrees rooted at nodes that cannot lead to a solution and backtracks to such a node's parent to continue the search

Example:

n-Queens Problem Place *n* queens on an *n*-by-*n* chess board so that no two of them are in the same row, column, or diagonal

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State-Space Tree of the 4-Queens Problem

7.1.1N-Queens Problem:

- The object is to place queens on a chess board in such as way as no queen can capture another one in a single move
 - Recall that a queen can move horz, vert, or diagonally an infinite distance
 - This implies that no two queens can be on the same row, col, or diagonal
 - We usually want to know how many different placements there are



4-Queens

- Lets take a look at the simple problem of placing queens 4 queens on a 4x4 board
- The brute-force solution is to place the first queen, then the second, third, and forth
 After all are placed we determine if they are placed legally
- There are 16 spots for the first queen, 15 for the second, etc.
 - Leading to 16*15*14*13 = 43,680 different combinations
- Obviously this isn't a good way to solve the problem
- First lets use the fact that no two queens can be in the same col to help us
 - That means we get to place a queen in each col

- So we can place the first queen into the first col, the second into the second, etc.
- This cuts down on the amount of work
 - Now there are 4 spots for the first queen, 4 spots for the second, etc.
 - 4*4*4*4 = 256 different combinations
- However, we can still do better because as we place each queen we can look at the previous queens we have placed to make sure our new queen is not in the same row or diagonal as a previously place queen
- Then we could use a Greedy-like strategy to select the next valid position for each col



- So now what do we do?
- Well, this is very much like solving a maze
 - As you walk though the maze you have to make a series of choices
 - If one of your choices leads to a dead end, you need to back up to the last choice you made and take a different route
 - That is, you need to change one of your earlier selections
 - Eventually you will find your way out of the maze



• This type of problem is often viewed as a state-space tree

- A tree of all the states that the problem can be in
- We start with an empty board state at the root and try to work our way down to a leaf node
 - Leaf nodes are completed boar



Eight Queen Problem

- The solution is a vector of length 8 (a(1), a(2), a(3),, a(8)). a(i) corresponds to the column where we should place the i-th queen.
- The solution is to build a partial solution element by element until it is complete.
- We should backtrack in case we reach to a partial solution of length k, that we couldn't expand any more.

Eight Queen Problem: Algorithm

```
putQueen(row) {
 for every position col on the same row
   if position col is available
     place the next queen in position col
   if (row<8)
     putQueen(row+1);
   else success;
   remove the queen from position col
}
putQueen(row) {
 for every position col on the same row
   if position col is available
     place the next queen in position col
   if (row<8)
     putQueen(row+1);
   else success;
   remove the queen from position col
}
```

Eight Queen Problem: Implementation

- Define an 8 by 8 array of 1s and 0s to represent the chessboard
- The array is initialized to 1s, and when a queen is put in a position (c,r), board[r][c] is set to zero
- Note that the search space is very huge: 16,772, 216 possibilities.
- Is there a way to reduce search space? Yes Search Pruning.
- We know that for queens: each row will have exactly one queen each column will have exactly one queen each diagonal will have at most one queen
- This will help us to model the chessboard not as a 2-D array, but as a set of rows, columns and diagonals.

7.2 HAMILTONIAN CYCLE

- Hamiltonian Cycle: a cycle that contains every node exactly once
- Problem: Given a graph, does it have a Hamiltonian cycle?

Background

- **NP-complete** problem:
 - Most difficult problems in NP (non- deterministic polynomial time)
- A decision problem *D* is NP-complete if it is complete for NP, meaning that:
 - it is in NP
 - it is NP-hard (every other problem in NP is reducible to it.)
- As they grow large, we are not able to solve them in a reasonable time (polynomial time)

Alternative Definition

- . NP Problem such as Hamiltonian Cycle :
 - Cannot be solved in Poly-time
 - Given a solution, easy to verify in poly-time



7.3 SUBSET –SUM PROBLEM.

- Problem: Given *n* positive integers *w*1, ... *wn* and a positive integer S. Find all subsets of *w*1, ... *wn* that sum to S.
- Example: n=3, S=6, and w1=2, w2=4, w3=6
- Solutions: {2,4} and {6}
- We will assume a binary state space tree.
- The nodes at depth 1 are for including (yes, no) item 1, the nodes at depth 2 are for item 2, etc.
- The left branch includes wi, and the right branch excludes wi.
- The nodes contain the sum of the weights included so far

Sum of subset Problem: State SpaceTree for 3 items



The sum of the included integers is stored at the node.

A Depth First Search solution

- Problems can be solved using depth first search of the (implicit) state space tree.
- Each node will save its depth and its (possibly partial) current solution
- DFS can check whether node v is a leaf.
 - If it is a leaf then check if the current solution satisfies the constraints
 - Code can be added to find the optimal solution

A DFS solution

- Such a DFS algorithm will be very slow.
- It does not check for every solution state (node) whether a solution has been reached, or whether a *partial* solution can lead to a *feasible* solution
- Is there a more efficient solution?

Backtracking solution to Sum of Subsets

• Definition: We call a node *nonpromising* if it cannot lead to a feasible (or optimal)

solution, otherwise it is *promising*

- Main idea: Backtracking consists of doing a DFS of the state space tree, checking whether each node is promising and if the node is nonpromising backtracking to the node's parent
- The state space tree consisting of expanded nodes only is called the *pruned state space tree*
- The following slide shows the pruned state space tree for the sum of subsets example
- There are only 15 nodes in the pruned state space tree
- The full state space tree has 31 nodes



Backtracking algorithm

void checknode (node v) {
node u

Checknode

- Checknode uses the functions:
 - promising(v) which checks that the partial solution represented by v can lead to the required solution
 - aSolutionAt(v) which checks whether the partial solution represented by node v solves the problem.

Sum of subsets – when is a node "promising"?

- Consider a node at depth i
- *weightSoFar* = weight of node, i.e., sum of numbers included in partial solution node represents
- *totalPossibleLeft* = weight of the remaining items i+1 to n (for a node at depth i)
 - A node at depth i is non-promising if (*weightSoFar* + *totalPossibleLeft* < S) or (*weightSoFar* + w[i+1] > S)
- To be able to use this "promising function" the w_i must be sorted in non-decreasing order A Pruned State Space Tree



sumOfSubsets (i, weightSoFar, totalPossibleLeft) 1) if (promising (i)) //may lead to solution 2) then if (weightSoFar = S) **then** print *include*[1] to *include*[*i*] 3) //found solution 4) //expand the node when *weightSoFar* < S else 5) include [i + 1] = "yes" //try including sumOfSubsets (i + 1, 6) weightSoFar + w[i + 1], totalPossibleLeft - w[i + 1]) 7) include [*i* + 1] = "no" //try excluding 8) sumOfSubsets (i + 1, weightSoFar, totalPossibleLeft - w[i + 1]) boolean promising (i)1) return (*weightSoFar* + *totalPossibleLeft* \geq S) && (weightSoFar == S // weightSoFar + $w[i+1] \leq S$) Prints all solutions!

W_i

Initial call sum Of Subsets (0, 0,)

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Feasible Solution vs. Optimal Solution

- DFS, BFS, hill climbing and best-first search can be used to solve some searching problem for searching a feasible solution.
- However, they cannot be used to solve the optimization problems for searching an (the) optimal solution.

The Branch-and-bound strategy

- This strategy can be used to solve optimization problems without an exhaustive search in the average case.
- 2 mechanisms:
 - A mechanism to generate branches when searching the solution space
 - A mechanism to generate a bound so that many braches can be terminated
- It is efficient in the average case because many branches can be terminated very early.
- Although it is usually very efficient, a very large tree may be generated in the worst case.
- Many NP-hard problem can be solved by B&B efficiently in the average case; however, the worst case time complexity is still exponential.

Bounding

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- A bound on a node is a guarantee that any solution obtained from expanding the node will be:
 - Greater than some number (lower bound)
 - Or less than some number (upper bound)
- If we are looking for a minimal optimal, as we are in weighted graph coloring, then we need a lower bound
 - For example, if the best solution we have found so far has a cost of 12 and the lower bound on a node is 15 then there is no point in expanding the node
 - The node cannot lead to anything better than a 15
- We can compute a lower bound for weighted graph color in the following way:
 - The actual cost of getting to the node
 - Plus a bound on the future cost
 - Min weight color * number of nodes still to color
 - That is, the future cost cannot be any better than this
 - Recall that we could either perform a depth-first or a breadth-first search
 - Without bounding, it didn't matter which one we used because we had to expand the entire tree to find the optimal solution
 - Does it matter with bounding?
 - Hint: think about when you can prune via bounding
- We prune (via bounding) when:
 - (currentBestSolutionCost <= nodeBound)
- This tells us that we get more pruning if:
 - The currentBestSolution is low
 - And the nodeBound is high
- So we want to find a low solution quickly and we want the highest possible lower bound

 One has to factor in the extra computation cost of computing higher lower bounds vs. the expected pruning savings

The Assignment Problem

- In many business situations, management needs to assign personnel to jobs, jobs to machines, machines to job locations, or salespersons to territories.
- Consider the situation of assigning *n* jobs to *n* machines.
- When a job i (=1,2,...,n) is assigned to machine j (=1,2,n) that incurs a cost Cij.
- The objective is to assign the jobs to machines at the least possible total cost.
- This situation is a special case of the Transportation Model And it is known as the *assignment problem*.
- Here, jobs represent "sources" and machines represent "destinations."
- The supply available at each source is 1 unit And demand at each destination is 1 unit.

| | | 5-1 | A . | <i>l</i> achine | | a |
|-------------|-------|-----|--------|-----------------|-----|--------|
| 1 | | 1 | 2 | | n | Source |
| | 1 | C11 | C12 | · · · · · · · | C1n | 1 |
| 1 | 2 | C21 | C22 | | C2n | 1 |
| Job | 2 | j. | 8 | | | |
| | 4 | | j. (4. | | | |
| | | | | | 1.1 | 1 |
| 62 YE YE | n | Cnl | Cn2 | | Cnn | 1 |
| Destination | i (). | 1 | 1 | anan. | 1 | |

The assignment model can be expressed mathematically as follows:

0, if the job j is not assigned to machine i

1, if the job j is assigned to machine i

```
rn
Mir.ΣΣC:jXij
i=1i=:
```

(Sum of assignments from a source should be exactly equal to 1):

n Σ Xij = 1 For i=1,2,...,r. i=1

(Sum of assignments to a destination should be equal to the demanded quantity by that destination):

```
n
Σ Xij = 1 For j=1,2,...,n
i=1
```

(Quantizies to be assigned car. be either 0 or 1):

Xij=0 cr 1 Foraliandj.

Xij=

The Assignment Problem Example

- Ballston Electronics manufactures small electrical devices.
- Products are manufactured on five different assembly lines (1,2,3,4,5).
- When manufacturing is finished, products are transported from the assembly lines to one of the five different inspection areas (A,B,C,D,E).
- Transporting products from five assembly lines to five inspection areas requires different times (in minutes)

| | | Inspectio | on Area | | |
|---------------|----|-----------|---------|----|----|
| Assembly Line | A | В | С | D | E |
| 1 | 10 | 4 | 6 | 10 | 12 |
| 2 | 11 | 7 | 7 | 9 | 14 |
| 3 | 13 | 8 | 12 | 14 | 15 |
| 4 | 14 | 16 | 13 | 17 | 17 |
| 5 | 19 | 11 | 17 | 20 | 19 |

Under current arrangement, assignment of inspection areas to the assembly lines are 1 to A, 2 to B, 3 to C, 4 to D, and 5 to E. This arrangement requires 10+7+12+17+19 = 65 man minutes.

- Management would like to determine whether some other assignment of production lines to inspection areas may result in less cost.
- This is a typical assignment problem. n = 5 And each assembly line is assigned to each inspection area.
- It would be easy to solve such a problem when n is 5, but when n is large all possible alternative solutions are n!, this becomes a hard problem.
- Assignment problem can be either formulated as a linear programming model, or it can be formulated as a transportation model.
- However, An algorithm known as *Hungarian Method* has proven to be a quick and efficient way to solve such problems.

Assignment Problem Revisited:

Select one element in each row of the cost matrix *C* so that:

- no two selected elements are in the same column
- the sum is minimized

Example

| | Job 1 | Job 2 | Job 3 | Job 4 |
|------------|-------|-------|-------|-------|
| Person a | 9 | 2 | 7 | 8 |
| Person b | 6 | 4 | 3 | 7 |
| Person c | 5 | 8 | 1 | 8 |
| Person d | 7 | 6 | 9 | 4 |

Lower bound: Any solution to this problem will have total cost at least: 2 + 3 + 1 + 4 (or 5 + 2 + 1 + 4)

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lb = 2+3+1+4=10

a

 $\rightarrow 1$

lb = 9+3+1+4=17

Example: First two levels of the state-space tree $\begin{array}{c} 0\\ \hline Start\\ lb = 2+3+1+4=10\\ \hline 3\\ \hline 4\end{array}$

Example: First two levels of the state-space tree

а

→ 3

lb =7+4+5+4=20

Figure 11.5 Levels 0 and 1 of the state-space tree for the instance of the assignment problem being solved with the best-first branch-and-bound algorithm. The number above a node shows the order in which the node was generated. A node's fields indicate the job number assigned to person a and the lower bound value, lb, for this node.



Figure 11.6 Levels 0, 1, and 2 of the state-space tree for the instance of the assignment problem being solved with the best-first branch-and-bound algorithm

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4

lb = 8+3+1+6=18

Example: Complete state-space tree



Figure 11.7 Complete state-space tree for the instance of the assignment problem solved with the best-first branch-and-bound algorithm

7.5 TRAVELING SALESPERSON PROBLEM

- This is a classic CS problem
- Given a graph (cities), and weights on the edges (distances) find a minimum weight tour of the cities
 - Start in a particular city
 - Visit all other cities (*exactly* once each)
 - Return to the starting city
- Cannot be done by brute-force as this is worst-case exponential or worse running time
 - So we will look to backtracking with pruning to make it run in a reasonable amount of time in most cases
- We will build our state space by:
 - Having our children be all the potential cities we can go to next
 - Having the depth of the tree be equal to the number of cities in the graph
 we need to visit each city exactly once
- So given a fully connected set of 5 nodes we have the following state space
 - only partially completed

Traveling Salesperson Problem



- Now we need to add bounding to this problem
 - It is a minimization problem so we need to find a lower bound
- We can use:
 - The current cost of getting to the node plus
 - An underestimate of the future cost of going through the rest of the cities
 - The obvious choice is to find the minimum weight edge in the graph and multiply that edge weight by the number of remaining nodes to travel through
- As an example assume we have the given adjacency matrix
- If we started at node A and have just traveled to node B then we need to compute the bound for node B
 - Cost 14 to get from A to B
 - Minimum weight in matrix is 2 times 4 more legs to go to get back to node A = 8
 - For a grand total of 14 + 8 = 22

| 0 | 14 | 4 | 10 | 20 | |
|----|----|----|----|----|--|
| 14 | 0 | 7 | 8 | 7 | |
| 4 | 5 | 0 | 7 | 16 | |
| 11 | 7 | 9 | 0 | 2 | |
| 18 | 7 | 17 | 4 | 0 | |

Recall that if we can make the lower bound higher then we will get more pruning Note that in order to complete the tour we need to leave node B, C, D, and E

- The min edge we can take leaving B is min(14, 7, 8, 7) = 7
- Similarly, C=4, D=2, E=4

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This implies that at best the future underestimate can be 7+4+2+4=17

- 17 + current cost of 14 = 31
 - This is much higher than 8 + 14 = 22

| 0 | 14 | 4 | 10 | 20 | |
|----|----|----|----|----|--|
| 14 | 0 | 7 | 8 | 7 | |
| 4 | 5 | 0 | 7 | 16 | |
| 11 | 7 | 9 | 0 | 2 | |
| 18 | 7 | 17 | 4 | 0 | |



7.6 THE 0/1 KNAPSACK PROBLEM



How to find the upper bound?

• Ans: by quickly finding a feasible solution in a greedy manner: starting from the smallest available i, scanning towards the largest i's until M is exceeded. The upper bound can be calculated.

The 0/1 knapsack problem

• E.g. n = 6, M = 34

| i | 1 | 2 | 3 | 4 | 5 | 6 |
|------------------|----|----|---|----|----|---|
| P _i | 6 | 10 | 4 | 5 | 6 | 4 |
| \mathbf{W}_{i} | 10 | 19 | 8 | 10 | 12 | 8 |

$$(P_i/W_i \geq \textbf{P}_{i+1}/W_{i+1})$$

• A feasible solution: X1 = 1, X2 = 1, X3 = 0, X4 = 0, X5 = 0, X6 = 0-(P1+P2) = -16 (upper bound) Any solution higher than -16 can not be an optimal solution.

How to find the lower bound?

• Ans: by relaxing our restriction from Xi = 0 or 1 to $0 \le Xi \le 1$ (knapsack problem)

Let $-\sum_{i=1}^{n} P_i X_i$ be an optimal solution for O/1 knapsack problem and $-\sum_{i=1}^{n} P_i X_i$ be an optimal solution for fractional knapsack problem. Let $Y = -\sum_{i=1}^{n} P_i X_i$, $Y' = -\sum_{i=1}^{n} P_i X_i$. \clubsuit $Y' \leq Y$

7.7 APPROXIMATION APPROACH

Apply a fast (i.e., a polynomial-time) approximation algorithm to get a solution that is not necessarily optimal but hopefully close to it

Accuracy measures:

accuracy ratio of an approximate solution sa

 $r(sa) = f(sa) / f(s^*)$ for minimization problems

 $r(sa) = f(s^*) / f(sa)$ for maximization problems

where f(sa) and $f(s^*)$ are values of the objective function f for the approximate solution sa and

actual optimal solution s^*

<u>performance ratio</u> of the algorithm A the lowest upper bound of r(sa) on all instances

Nearest-Neighbor Algorithm for TSP

Starting at some city, always go to the nearest unvisited city, and, after visiting all the cities, return to the starting one

Note: Nearest-neighbor tour may depend on the starting city



Accuracy: $RA = \infty$ (unbounded above) – make the length of AD arbitrarily large in the above example

Multifragment-Heuristic Algorithm

Stage 1: Sort the edges in nondecreasing order of weights. Initialize the set of tour edges to be constructed to empty set

Stage 2: Add next edge on the sorted list to the tour, skipping those whose addition would've created a vertex of degree 3 or a cycle of length less than n. Repeat this step until a tour of length n is obtained

Note: $RA = \infty$, but this algorithm tends to produce better tours than the nearest-neighbor algorithm

Twice-Around-the-Tree Algorithm

Stage 1: Construct a minimum spanning tree of the graph(e.g., by Prim's or Kruskal's algorithm) Stage 2: Starting at an arbitrary vertex, create a path that goes twice around the tree and returns to the same vertex

Stage 3: Create a tour from the circuit constructed in Stage 2 by making shortcuts to avoid visiting intermediate vertices more than once

Note: $RA = \infty$ for general instances, but this algorithm tends to produce better tours than the nearest-neighbor algorithm

Christofides Algorithm

Stage 1: Construct a minimum spanning tree of the graph

Stage 2: Add edges of a minimum-weight matching of all the odd vertices in the minimum spanning tree

Stage 3: Find an Eulerian circuit of the multigraph obtained in Stage 2

Stage 3: Create a tour from the path constructed in Stage 2 by making shortcuts to avoid visiting intermediate vertices more than once

 $RA = \infty$ for general instances, but it tends to produce better tours than the twice-around-theminimum-tree alg.

Euclidean Instances

<u>Theorem</u> If $P \neq NP$, there exists no approximation algorithm for TSP with a finite performance ratio.

<u>Definition</u> An instance of TSP is called *Euclidean*, if its distances satisfy two conditions: 1. *symmetry* d[i, j] = d[j, i] for any pair of cities *i* and *j*

2. *triangle inequality* $d[i, j] \le d[i, k] + d[k, j]$ for any cities *i*, *j*, *k*

For Euclidean instances:

approx. tour length / optimal tour length $\leq 0.5(\log 2n + 1)$ for nearest neighbor and multifragment heuristic; approx. tour length / optimal tour length ≤ 2 for twice-around-the-tree; approx. tour length / optimal tour length ≤ 1.5 for Christofides

Local Search Heuristics for TSP

Start with some initial tour (e.g., nearest neighbor). On each iteration, explore the current tour's neighborhood by exchanging a few edges in it. If the new tour is shorter, make it the current tour; otherwise consider another edge change. If no change yields a shorter tour, the current tour is returned as the output.

Greedy Algorithm for Knapsack Problem

Step 1: Order the items in decreasing order of relative values:

 $v_1/w_1 \ge \ldots \ge v_n/w_n$

Step 2: Select the items in this order skipping those that don't fit into the knapsack

| Examp | le: The | knapsack's | capacity | is 16 |
|-------|----------|------------|----------|-------|
| Lamp | 10. 1110 | mupbuen b | cupacity | 10 10 |

| item | weight | value | v/w |
|------|--------|-------|-----|
| 1 | 2 | \$40 | 20 |
| 2 | 5 | \$30 | 6 |
| 3 | 10 | \$50 | 5 |
| 4 | 5 | \$10 | 2 |

Accuracy

- R_A is unbounded (e.g., n = 2, C = m, $w_1=1$, $v_1=2$, $w_2=m$, $v_2=m$)
- yields exact solutions for the continuous version

Approximation Scheme for Knapsack Problem

Step 1: Order the items in decreasing order of relative values:

$$v_1/w_1 \ge \ldots \ge v_n/w_n$$

Step 2: For a given integer parameter k, $0 \le k \le n$, generate all subsets of k items or less and for each of those that fit the knapsack, add the remaining items in decreasing order of their value to weight ratios

Step 3: Find the most valuable subset among the subsets generated in Step 2 and return it as the

algorithm's output

Bin Packing Problem: First-Fit Algorithm

<u>*First-Fit (FF)* Algorithm</u>: Consider the items in the order given and place each item in the first available bin with enough room for it; if there are no such bins, start a new one Example: n = 4, $s_1 = 0.4$, $s_2 = 0.2$, $s_3 = 0.6$, $s_4 = 0.7$

Accuracy

- Number of extra bins never exceeds optimal by more than 70% (i.e., $R_A \le 1.7$)
- Empirical average-case behavior is much better. (In one experiment with 128,000 bins, the relative error was found to be no more than 2%.)

Bin Packing: First-Fit Decreasing Algorithm

First-Fit Decreasing (FFD) Algorithm: Sort the items in decreasing order (i.e., from the largest to the smallest). Then proceed as above by placing an item in the first bin in which it fits and starting a new bin if there are no such bins

Example: n = 4, $s_1 = 0.4$, $s_2 = 0.2$, $s_3 = 0.6$, $s_4 = 0.7$

Accuracy

- Number of extra bins never exceeds optimal by more than 50% (i.e., $R_A \le 1.5$)
- Empirical average-case behavior is much better, too

UNIT-8

PRAM ALGORITHMS

8.1 Introduction,

8.2 Computational Model,

8.3 Parallel Algorithms for Prefix Computation,

8.4 List Ranking, and Graph Problems,

8.1 INTRODUCTION

In this section a few basic facts about parallel processing in general. One very basic fact that applies to parallel computation, regardless of how it is implemented, is the following:

Suppose the fastest sequential algorithm for doing a computation with parameter n has execution time of T(n). Then the fastest parallel algorithm with m processors (each comparable to that of the sequential computer) has execution time T(n)/m. The idea here is: If you could find a faster parallel algorithm, you could execute it sequentially by having a sequential computer *simulate* parallelism and get a faster sequential algorithm. This would contradict the fact that the given sequential algorithm is the fastest possible. We are making the assumption that the *cost* of simulating parallel algorithms by sequential ones is negligible. This claim is called the "Principle of Unitary Speedup". As usual, the parameter *n* represents the relative size of the instance of the problem being considered. For instance, if the problem was that of sorting, *n* might be the number of items to be sorted and T(*n*) would be $O(n \lg n)$ for a sorting algorithm based upon comparisons.

As simple as this claim is, it is a bit controversial. It makes the tacit assumption that the algorithm in question is *deterministic*. In other words, the algorithm is like the usual idea of a computer program — it performs calculations and makes decisions based on the results of these calculations.

There is an interesting area of the theory of algorithms in which statement is *not necessarily true* — this is the theory of *randomized* algorithms. Here, a solution to a problem may involve making random "guesses" at some stage of the calculation. In this case, the parallel algorithm using *m* processors can run faster than $m \times$ the speed of the sequential algorithm ("Super-unitary speedup"). This phenomenon occurs in certain problems in which random search is used, and

most guesses at a solution quickly lead to a valid solution, but there are a few guesses that execute for a long time without producing any concrete results.

The expected execution-time of a *single (sequential) attempt* to find a solution is the *average* of all of these times, or 10.99 time-units.

8.2. MODELS OF PARALLEL COMPUTATION

If we attack this problem with a parallel computer that has 2 processors that try *distinct* possibilities, the expected time (and even the *worst-case* time) is 1 unit, since at least one of these two distinct possibilities will be a *fast solution*, consequently, see a *super-unitary* speedup in the parallel algorithm. In other words the *expected*1 running-time of the algorithm is divided by > 10, which is much greater than the ratio of processors.

The opponents of the concept of super-unitary speedups (including the author) would argue that the original sequential algorithm was not *optimal* — and that the optimal sequential algorithm would have attempted two possible solutions with two distinct processes, run *concurrently*). A sequential algorithm that created two processes would have a running time of 2 units. The speedup that results by going to the sample parallel algorithm is 2, which is exactly equal to the ratio of processors. Thus, by modifying the sequential algorithm used, the validity is restored.

In this section, consider some simple algorithms that can be implemented when a SIMD computer in which every processor can access common RAM. In general, a computer in which many processors can access common RAM in a single program-step is called the **PRAM** model of computer. This is one of the oldest models of parallel processing to be considered, although there have never been any large parallel computers that implement it. The PRAM model is a kind of mathematical idealization of a parallel computer that eliminates many of the low-level details and allows a researcher to concentrate on the "purely parallel" aspects of a problem.

Traditionally, the PRAM model of computation has been regarded as more of theoretical than practical interest. This is due to the fact that it requires large numbers of processors to be physically connected to the same memory location. The two examples cited above (the Sequent and the Encore Multimax) aren't an exception to this statement: they only have small numbers of processors. Several researchers are exploring the possibility of physically realizing of PRAM computers using optical interfaces.

There exist several different models of program control. Flynn listed several basic schemes:

SIMD: Single Instruction Multiple Data. In this model the processors are controlled by a program whose instructions are applied to all of them simultaneously (with certain qualifications). Assume that each of the processors has a unique number that is "known" to the processor in the sense that instructions to the parallel computer can refer to processor numbers. An example of this type of machine is: 1Recall that the expected running-time of an algorithm like the one in the example is the average of actual running times, weighted by probabilities that these running times occur.

MIMD: Multiple Instruction Multiple Data. In this model processors can each have independent programs that are read from the common RAM. This model is widely used in several settings: The data-movement and communications problems that occur in all parallel computation are more significant here because the instructions to the processors as well as the data must be passed between the common memory and the processors. Due to these data-movement problems, commercial MIMD computers tend to have a relatively small number of processors (1/4 20). In general, it is easier to program a MIMD machine if one is only interested in a very limited form of parallelism—namely the formation of processes. Conventional operating systems like UNIX form separate processes to carry out many functions, and these processes *really* execute in parallel on commercially-available MIMD machines. It follows that, with one of these MIMD machines, one can reap some of the benefits of parallelism without explicitly doing any parallel programming. For this reason, most of the parallel computers in commercial use today tend to be MIMD machines, run as general-purpose computers. On the surface, it would appear that MIMD machine are strictly more powerful than SIMD machines with the same number of processors. Interestingly enough, this is not the case—it turns out that SIMD machines are more suited to performing computations with a very *regular* structure. MIMD machines are not as suited to solving such problems because their processors must be *precisely synchronized* to implement certain algorithms. Pure MIMD machines have no hardware features to guarantee synchronization of processors. In general, it is not enough to simply load multiple copies of a program into all of the processors and to start all of these copies at the same time. In fact many such computers have hardware features that tend to *destroy* synchronization, once it

has been achieved. For instance, the manner in which memory is accessed in the Sequent Symmetry series, generally causes processes to run at different rates even if they are synchronized at some time. Many Sequent's even have processors that run at different clockrates.

8.2.1 MODELS OF PARALLEL COMPUTATION

Three other terms that fill out this list are:

SISD Single Instruction, Single Data. This is nothing but conventional sequential computing.

MISD This case is often compared to computation that uses *Systolic Arrays*. These are arrays of processors that are developed to solve specific problems — usually on a single VLSI chip. A clock coordinates the data movement operations of all of the processors, and output from some processors are pipelined into other processors. The term "Systolic" comes from an analogy with an animal's circulatory system — the data in the systolic array playing the part of the blood in the circulatory system. In a manner of speaking, one can think of the different processors in a systolic array as constituting "multiple processors" that work on one set of (pipelined) data

(pipelined) data.

SIMD-MIMD Hybrids This is a new category of parallel computer that is becoming very significant. These machines are also called SAMD machines (Synchronous-Asynchronous Multiple Data). The first announced commercial SAMD computer is the new Connection Machine, the CM-5.

This is essentially a MIMD computer with hardware features to allow:

• Precise synchronization of processes to be easily achieved.

• Synchronization of processors to be maintained with little or no overhead,

Once it has been achieved (assuming that the processors are all executing the same instructions in corresponding program steps). It differs from pure MIMD machines in that the hardware maintains a uniform "heartbeat" throughout the machine, so that when the same program is run on all processors, and all copies of this program are started at the same time, it is possible to the execution of all copies to be kept in lock-step with essentially no overhead. Such computers allow efficient execution of MIMD and SIMD programs.

8.3 PARALLEL ALGORITHMS FOR PREFIX COMPUTATION

PRAM Recursive Prefix Sum Algorithm

Input: Array of (x1, x2, ..., xn) elements, n = 2k**Output**: Prefix sums si, 1 < i < n**begin if** n = 1 **then** s1 = x1; **exit for** 1 < i < n/2 **pardo** yi := x2i-1 + x2iRecursively compute prefix sums of y and store in z**for** 1 < i < n **pardo if** i is even **then** si := zi/2**if** i > 1 is odd **then** si := z(i-1)/2 + xi**if** i = 1 **then** s1 := x1

Theorem: An algorithm that runs in T time on the p-processor priority CRCW PRAM can be simulated by EREW PRAM to run in $O(T \log$

p) time. A concurrent read or write of an *p*-processor CRCW PRAM can be implemented on a *p* processor EREW PRAM to execute in $O(\log p)$ time

Q1,...,Qp CRCW processors, such that Qi has to read (write) M[ji]

P1,...,Pp EREW processors

M1,...,Mp denote shared memory locations for special use Pi stores $\langle ji,i \rangle$ in MiSort pairs in lexicographically non-decreasing order in O(log p) time using EREW merge sort algorithm

Pick representative from each block of pairs that have same firs component in O(1) time Representative Pi reads (writes) from M[k] with $\langle k, \rangle$ in Mi and copies data to each M in the block in O(log p) time using EREW segmented parallel prefix algorithm Pi reads data from Mi

For n x n matrix M with non-negative integer coefficients, define M and give an algorithm for computing m. prove that M cnan be computed from n x n matrix M in O(log n)time using CRCW PRAM processors for any fixed $\epsilon > 0$

Consider $n \times n$ matrix multiplication C = A B using n3 processors.

Each element of C

 $cij = \Sigma k = 1..n \ aik \ bkj$

can be computed on the CREW PRAM in parallel using *n* processors in $O(\log n)$ time All *cij* can be computed using *n*3 processors in $O(\log n)$

Matrix multiply with *n*3 processors (i,j,l)Each processor (i,j,l) executes: **Input**: $n \times n$ matrices *A* and *B*, n = 2k**Output**: C = A B**begin** C'[i,j,l] := A[i,l]B[l,j]**for** h = 1 **to** log *n* **do**

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if *i* < *n*/2*h* **then** *C*'[*i*,*j*,*l*] := *C*'[*i*,*j*,2*l*-1] + *C*'[*i*,*j*,2*l*] **if** *l* = 1 **then** *C*[*i*,*j*] := *C*'[*i*,*j*,1] **End**

8.4 LIST RANKING AND GRAPH PROBLEMS. Convex Hull Problem

The *planar convex hull* of a set of points $S = \{p1, p2, ..., pn\}$ of pi = (x,y) coordinates is the smallest convex polygon that encompasses all points *S* on the *x*-*y* plane



Preprocess the points by sorting in O(log *n*) parallel time (pipelined merge sort), such that x(p1) < x(p2) < ... < x(pn)Remove duplicates x(pi) = x(pj) (for UCH and LCH)

Divide-and-conquer:

1. Split *S* into *S*1 *S*2 and recursively compute the UCH of *S*1 and *S*2 2. Combine UCH(*S*1) and UCH(*S*2) by computing the upper common tangent in O(1) time to form UCH(*S*) Repeat to compute the LCH Parallel time (assuming p = O(n) processors) T(n) = T(n/2) + O(1) gives $T(n) = O(\log n)$