
A SYSTEMATIC APPROACH TO

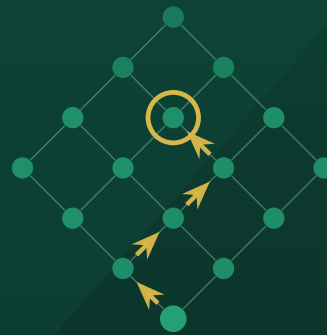
Sequential Algorithms

LLP-OddEvenSort

A: array of int

forbidden(j): $A[j] > A[j+1]$

advance(j): $\text{swap}(A[j], A[j+1])$



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To my family

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Preface

Approach

This book presents algorithms from two complementary perspectives. The *traditional* perspective develops each algorithm in its classical form: greedy choices, divide-and-conquer recursions, dynamic-programming table fills, network-flow augmenting paths, and so on. The *lattice-linear predicate* (LLP) perspective recasts each problem as a search for an extremal element—typically the least element—of an appropriate distributive lattice that satisfies a Boolean predicate B . An algorithm in the second form maintains a single state vector G , identifies any component that is *forbidden* under B , and *advances* it according to a rule derived from the problem’s structure.

The lattice-search perspective brings several advantages:

- **Simpler statements.** Almost every algorithm in this book uses a single state vector G . Any auxiliary state, when present, is syntactic sugar over the underlying lattice machinery.
- **Nondeterministic schedules.** The LLP loop fixes only that *some* forbidden index advances next; the choice of *which* is left open. A proof that holds for the nondeterministic schema applies uniformly to every concrete schedule, including the textbook sequential ones.
- **Vector-based reasoning.** Reasoning about G as a vector in a lattice replaces case analysis with monotonicity arguments.
- **Natural parallelism.** Because the schedule is open, every forbidden index is a candidate for advance in the same round. Many of the algorithms in this book are parallel by inheritance, requiring no separate derivation.
- **Easier correctness proofs.** The standard recipe—prove lattice-linearity of B , exhibit the advance rule, and apply the LLP fixed-point theorem—replaces the algorithm-specific invariant work found in classical presentations.

Guiding principles

The book is organized around three principles.

Derive, do not present. The goal of each chapter is to *derive* the algorithm from the feasibility and optimality structure of the underlying problem, rather than to present a finished algorithm and then verify that it works. The lattice formulation makes this derivation systematic: identify the lattice, write down B , check lattice-linearity, and read off the advance rule.

Prefer nondeterminism. Whenever possible, algorithms are stated nondeterministically. A nondeterministic statement is typically simpler than any of its deterministic refinements, and—crucially—a single proof applies to the entire class of refinements. The reader who wants a sequential schedule, a parallel schedule, or a distributed schedule can specialize the same algorithm to her needs.

Parallel by default. Many algorithms in this book admit a simple parallel form: start from a feasible point and improve it monotonically toward the optimum. The LLP framework exposes this structure explicitly—every forbidden index can advance in the same round—so parallelism need not be retrofitted onto a sequential description.

Companion material

Every algorithm in the book ships as runnable code on the companion website. Each algorithm is written once in a small lattice-linear DSL and compiled to Java, Python, C++, and JavaScript. The website also hosts interactive demos that step through the LLP loop on small concrete instances. Both the source code and the demos are intended to help the reader develop intuition for what an LLP algorithm *does*, beyond what the printed page alone can convey.

A solution manual covering the end-of-chapter problems is available to instructors on request.

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The list of known errors and supplementary material for the book is maintained at:

<http://www.ece.utexas.edu/~garg>

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Chapter 1

Introduction to Algorithms

1.1 Introduction

Computers are fast, but algorithms are what make them useful. The same problem can admit algorithms whose running times differ by factors of millions on inputs that already fit comfortably in memory: sorting a billion records takes minutes with a good algorithm but centuries with a bad one. Understanding what makes one algorithm faster than another, and being able to predict how an algorithm scales before running it, is the central skill of algorithm design.

This chapter collects the preliminaries that the rest of the book relies on. We define what an algorithm is, introduce the asymptotic notation used to compare running times, and review the basic data structures — linked lists, stacks, queues, binary search trees, and heaps — that the subsequent chapters take as building blocks. The chapter closes with a roadmap of the rest of the book, in which we preview each chapter and its connection to the lattice-linear-predicate (LLP) framework introduced in Chapter 2.

This chapter is organized as follows. Section 1.2 defines what an algorithm is, using Euclid’s GCD as a running example. Section 1.3 introduces asymptotic notation (Big-O, Big-Omega, Big-Theta) for describing growth rates. Section 1.4 analyses the running time of several familiar algorithms (linear search, binary search, the Tower of Hanoi) to make the asymptotic vocabulary concrete. Section 1.5 reviews the basic data structures (linked lists, stacks, queues, binary search trees) that later chapters take as primitives. Section 1.6 describes the heap data structure and the linear-time Build-Heap algorithm.

1.2 What is an Algorithm?

An algorithm is a finite, unambiguous sequence of instructions that solves a problem or computes a function, transforming an input into an output in a finite number of steps. It must exhibit *finiteness* (terminates), *definiteness* (each step is precisely specified), and *effectiveness* (each operation is basic and executable). Algorithms are the foundation of computer science, solving problems from simple arithmetic to complex optimization.

We start with one of the earliest algorithms due to Euclid. Suppose that we are required to find the greatest common divisor (GCD) of two natural numbers a and b (integers greater than or equal to 1). If they are both equal, then we have the answer: a . Suppose that a is greater than b , then we claim that it

is safe to reduce a to $a - b$ (prove it!). Similarly, if b is greater than a , then we reduce b to $b - a$. The algorithm terminates with both numbers identical and equal to the GCD.

Algorithm GCD: GCD(a , b)

Input: Two positive integers a , b greater than or equal to 1

Output: The greatest common divisor of a and b

```

1 while  $a \neq b$  do
2   | if  $(a > b)$  then  $a := a - b$ ;
3   | if  $(b > a)$  then  $b := b - a$ ;
4 end
5 return  $a$ 

```

Example 1.1 For $a = 48$, $b = 18$, in the first step, we get a equals 30 and b equals 18. Since a is still greater than b , a becomes 12 and b is still 18. Now b is greater than a . So, in the next step b becomes $b - a$ which equals 6. Finally, a becomes 6. Now, both a and b are equal to 6 and the algorithm terminates.

It is easy to see that the algorithm always terminates. All values are initially non-zero positive integers. They stay non-zero and integral and always decrease; therefore, the algorithm must terminate.

In our algorithm, we have used subtraction to decrease the numbers. We leave it as an exercise for the reader to use the *mod* function to speed up the above algorithm.

Suppose that we want to compute the factorial of a non-negative integer n . Algorithm [Factorial](#) shows a method that uses *recursion*. A recursive approach requires specification of the *base case*, when n equals 0. For a higher value of n , we use the value of $\text{Factorial}(n - 1)$ to compute $\text{Factorial}(n)$.

Algorithm Factorial: Factorial(n)

Input: A non-negative integer n

Output: $n!$

```

1 if  $n = 0$  then return 1 ;
2 return  $n \cdot \text{Factorial}(n - 1)$ 

```

As a final simple example of an algorithm, suppose that we are required to find the largest element in an array A . Algorithm [MaxElement](#) uses a *for* loop to find this element.

1.3 Asymptotic Notation (Big O, Big Omega, Big Theta)

Asymptotic notation quantifies runtime growth, abstracting constants and lower-order terms to focus on scalability. It is the cornerstone of algorithmic comparison.

- $T(n) = O(f(n))$ if $\exists c, n_0 > 0$ such that $T(n) \leq c \cdot f(n)$ for all $n \geq n_0$. For example, suppose that $T(n) = 2n + 5$. Since $2n + 5 \leq 3n$ for $n \geq 5$, we conclude that $T(n)$ equals $O(n)$.

Algorithm MaxElement: MaxElement(A)

Input: Array A of size n **Output:** Maximum element in A

```

1  $max := A[0]$  ;
2 for  $i = 1$  to  $n - 1$  do
3   | if  $A[i] > max$  then  $max := A[i]$  ;
4 end
5 return  $max$ 

```

- $T(n) = \Omega(f(n))$ if $\exists c, n_0 > 0$ such that $T(n) \geq c \cdot f(n)$ for all $n \geq n_0$. For example, suppose that $T(n) = n^2 - n$. Since $n^2 - n \geq 0.5n^2$ for $n \geq 2$, we conclude that $T(n)$ equals $\Omega(n^2)$.
- $T(n) = \Theta(f(n))$ if $T(n) = O(f(n))$ and $T(n) = \Omega(f(n))$. For example, suppose that $T(n) = 4n^2 + 3n$, $3n^2 \leq 4n^2 + 3n \leq 5n^2$ for $n \geq 3$. We conclude that $T(n)$ equals $\Theta(n^2)$.

Example 1.2 We first give an example, where $T(n)$ is a linearly growing function of n . Let $T(n) = 7n + 10$. Then, it is easy to see that $T(n)$ equals $O(n)$: $7n + 10 \leq 8n$ for $n \geq 10$. In addition, $T(n)$ equals $\Omega(n)$: $7n + 10 \geq 7n$. Thus, $T(n)$ equals $\Theta(n)$.

Example 1.3 As another example, suppose $T(n) = 2n^3 + 5n^2 + n$. Then $T(n)$ equals $O(n^3)$ because $2n^3 + 5n^2 + n \leq 3n^3$ for $n \geq 5$. Similarly, $T(n)$ equals $\Omega(n^3)$ because $2n^3 + 5n^2 + n \geq 2n^3$. Hence, $T(n)$ equals $\Theta(n^3)$.

1.4 Analyzing Algorithm Efficiency

Efficiency analysis evaluates the time and space complexity across the best, worst, and average cases, providing a holistic view of performance. We start with the Algorithm [LinearSearch](#).

Algorithm LinearSearch: LinearSearch(A, key)

Input: Array A of size n , key to find**Output:** Index of key or -1 if not found

```

1 for  $i = 0$  to  $n - 1$  do
2   | if  $A[i] = key$  then return  $i$  ;
3 end
4 return  $-1$ 

```

The time complexity of the algorithm is as follows.

- *Best:* $O(1)$, key at $A[0]$.

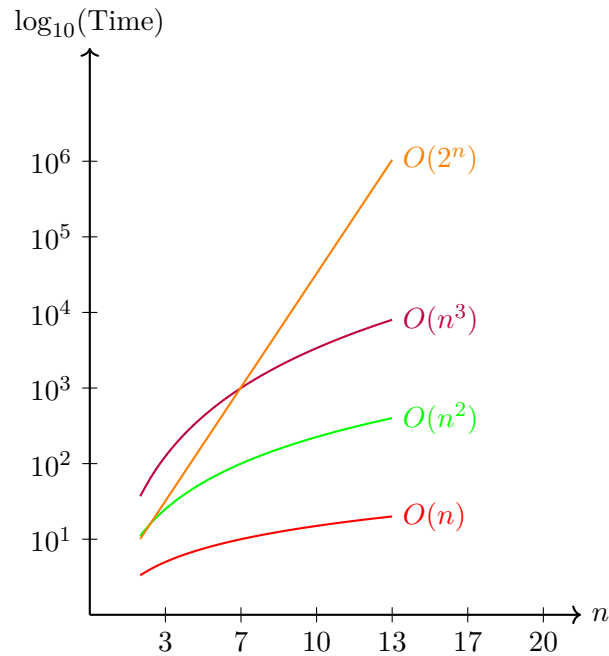


Figure 1.1: Comparison of Growth Rates for $n = 1$ to 20: Linear, Quadratic, Cubic, and Exponential (Log Scale)

- *Worst*: $O(n)$, key absent.
- *Average*: $\frac{n+1}{2} = O(n)$, assuming a uniform likelihood of the key.

We now consider the example of a *nested loop* shown in Algorithm [NestedSum](#). The time complexity of

Algorithm NestedSum: NestedSum(n)

Input: Integer n

Output: Sum of products

```

1 total := 0 ;
2 for i = 0 to n - 1 do
3   for j = 0 to n - 1 do
4     total := total + i · j ;
5   end
6 end
7 return total

```

the algorithm is $O(n^2)$. The space complexity of the algorithm is $O(1)$. From now on, the time and space complexity will always refer to the *worst case* unless otherwise specified.

Let us consider another example, in which our goal is to find the number of pairs in an array A that sum to the provided target sum. The reader should verify that the time complexity of the algorithm is $O(n^2)$.

Algorithm PairSum: PairSum(A , target)

Input: Array A of size n , target sum**Output:** Number of pairs summing to target

```

1 count := 0 ;
2 for i = 0 to n - 1 do
3   for j = i + 1 to n - 1 do
4     if A[i] + A[j] = target then count := count + 1 ;
5   end
6 end
7 return count

```

We now give an example of an algorithm [BinarySearch](#) that speeds up Algorithm [LinearSearch](#). Given a *key*, the algorithm compares it to the middle element of the array A . If *key* is equal to the middle element, then we have found the index in the array, *mid*, such that $A[\textit{mid}]$ equals *key*. If *key* is less than the middle element, then we know that it cannot be on the right side of the array A because the array is sorted. We can then restrict our attention to the left side of the array. Finally, if *key* is greater than the middle element, then we can restrict our attention to the right side of the array. If $T(n)$ equals the time complexity of searching in a sorted array of size n , then the recurrence $T(n) = T(n/2) + 1$ is satisfied. Later in this book, we show that $T(n) = O(\log n)$.

Algorithm BinarySearch: BinarySearch(A , key, low, high)

Input: Array A , key, range $[low, high]$ **Output:** Index of key or -1

```

1 if low > high then return -1 ;
2 mid := ⌊(low + high)/2⌋ ;
3 if A[mid] = key then return mid ;
4 else if A[mid] > key then return BinarySearch(A, key, low, mid - 1) ;
5 else return BinarySearch(A, key, mid + 1, high) ;

```

As another example, we describe the problem called Towers of Hanoi. The Towers of Hanoi problem involves three pegs and n disks of increasing size, initially stacked on one peg in decreasing size order. The goal is to move the entire stack to another peg, obeying two rules: only one disk can be moved at a time, and a larger disk cannot be placed on a smaller disk. The solution requires $2^n - 1$ moves, achieved recursively by moving $n - 1$ disks to a spare peg, transferring the largest disk, then moving the $n - 1$ disks onto the largest. Algorithm [Hanoi](#) shows a recursive solution to this problem. Here, the recurrence is: $T(n) = 2T(n - 1) + 1$, $T(1) = 1$. On solving this recurrence, we get $T(n) = 2^n - 1$.

Finally, we consider the problem of sorting an array A using a method called *MergeSort*. The algorithm uses recursion to sort an array using a method called divide-and-conquer. It sorts the left half of the array and the right half of the array, recursively. Now, given that the left side and the right side of the array are sorted, it merges these two sides to get a fully sorted array. In this example, we have the recurrence $T(n) = 2T(n/2) + n$. We later show that the recurrence can be solved to determine that $T(n) = O(n \log n)$.

Algorithm Hanoi: Hanoi(n , source, aux, target)

Input: Number of disks n , pegs

```

1 if  $n = 1$  then Move disk from source to target ;
2 else
3   | Hanoi( $n - 1$ , source, target, aux) ;
4   | Move disk from source to target ;
5   | Hanoi( $n - 1$ , aux, source, target) ;
6 end

```

Algorithm MergeSort: MergeSort(A , low, high)

Input: Array A , range $[low, high]$

```

1 if  $low < high$  then
2   |  $mid := \lfloor (low + high)/2 \rfloor$  ;
3   | MergeSort( $A$ , low, mid) ;
4   | MergeSort( $A$ , mid + 1, high) ;
5   | Merge( $A$ , low, mid, high) ;
6 end

```

1.5 Common Data Structures: Lists, Stacks, Queues, Binary Trees

This section briefly describes four fundamental data structures: linked lists, stacks, simple queues, and binary search trees, focusing on their main methods and time complexities.

Linked Lists

A linked list is a linear sequence of nodes, where each node contains data and a reference to the next node.

- **Insert** (at head/tail): $O(1)$
- **Delete** (a given node): $O(1)$ with pointer; $O(n)$ to find
- **Search**: $O(n)$

Stacks

A stack is a Last-In-First-Out (LIFO) structure, typically implemented with an array or linked list.

- **Push** (add to top): $O(1)$
- **Pop** (remove from top): $O(1)$
- **Top** (access top): $O(1)$

Simple Queues

A simple queue is a First-In-First-Out (FIFO) structure, often using an array or linked list.

- **Enqueue** (add to rear): $O(1)$
- **Dequeue** (remove from front): $O(1)$ with linked list; $O(n)$ with array
- **Front** (access front): $O(1)$

Binary Search Trees

A binary search tree (BST) is a binary tree in which the left subtree of each node has smaller values and the right subtree has larger values.

- **Insert**: $O(h)$ (height h ; $O(\log n)$ if balanced)
- **Delete**: $O(h)$ (height h ; $O(\log n)$ if balanced)
- **Search**: $O(h)$ (height h ; $O(\log n)$ if balanced)

1.6 Heaps

We now describe a data structure *heap* that is useful for many classical algorithms, such as Dijkstra's shortest path algorithm. A heap is a complete binary tree satisfying the heap property: In a minimum heap, the value of each parent is less than or equal to the values of its children. We focus on min-heaps here, implemented as arrays.

Structure and Representation

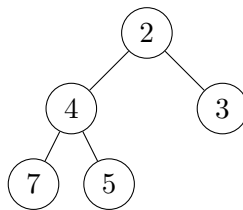


Figure 1.2: Min-heap as a binary tree

2	4	3	7	5
$A[0]$	$A[1]$	$A[2]$	$A[3]$	$A[4]$

Figure 1.3: Min-heap as an array

Figures 1.2 and 1.3 show a min-heap with values 2, 4, 3, 7, 5.

Heapify

The Heapify algorithm ensures the min-heap property at a given node by comparing it with its children and swapping with the smallest if necessary, then recursively applying the process downward. It takes an array A , an index i , and a heap size n , running in $O(\log n)$ time due to the height of the tree.

Algorithm Heapify: Heapify (Min-Heap)

Input: Array A , index i , heap size n

```

1 smallest := i ;
2 left := 2i + 1 ;
3 right := 2i + 2 ;
4 if left < n and  $A[\textit{left}] < A[\textit{smallest}]$  then smallest := left ;
5 if right < n and  $A[\textit{right}] < A[\textit{smallest}]$  then smallest := right ;
6 if smallest ≠ i then
7   | Swap  $A[i]$  and  $A[\textit{smallest}]$  ;
8   | Heapify( $A$ , smallest, n);
9 end
```

Insert

The Insert algorithm adds a new value to the heap by placing it at the end of the array and shifting it up to its correct position, comparing it with its parent until the min-heap property is restored. It runs in $O(\log n)$ time, as it traverses up the tree's height.

Algorithm HeapInsert: Heap-Insert

Input: Array A , value key , heap size n

```

1 n := n + 1 ;
2  $A[\textit{n} - 1]$  :=  $\infty$  ;
3 i := n - 1 ;
4 while i > 0 and  $A[\lfloor (i - 1)/2 \rfloor] > key$  do
5   |  $A[i]$  :=  $A[\lfloor (i - 1)/2 \rfloor]$  ;
6   | i :=  $\lfloor (i - 1)/2 \rfloor$ ;
7 end
8  $A[i]$  := key
```

Extract-Min

The Extract-Min algorithm removes the minimum element (root) by replacing it with the last element, reducing the heap size, and then calling Heapify to restore the min-heap property. It operates in $O(\log n)$ time due to the heapification step.

Algorithm HeapExtractMin: Heap-Extract-Min

Input: Array A , heap size n

```

1 if  $n < 1$  then return error ;
2  $min := A[0]$  ;
3  $A[0] := A[n - 1]$  ;
4  $n := n - 1$  ;
5 Heapify( $A, 0, n$ ) ;
6 return  $min$ 

```

Build-Heap

The Build-Heap algorithm constructs a min-heap from an unsorted array by iteratively applying Heapify from the last non-leaf node up to the root. It runs in $O(n)$ time, as the cumulative effect of heapifying all subtrees is linear in the number of nodes (Show this!).

Algorithm BuildHeap: Build-Heap (Min-Heap)

Input: Array A , size n

```

1 for  $i := \lfloor n/2 \rfloor - 1$  downto 0 do
2   | Heapify( $A, i, n$ )
3 end

```

The time complexity of various operations on a heap can be summarized as follows.

- **Heapify:** $O(\log n)$
- **Insert:** $O(\log n)$
- **Extract-Min:** $O(\log n)$
- **Build-Heap:** $O(n)$

1.7 Organization of the Book

Chapter 2 presents the general Lattice Linear Predicate (LLP) algorithm. It shows how many combinatorial optimization problems can be solved by a single framework and introduces the notions of forbidden indices and advance operations.

Chapter 3 presents the stable marriage problem. It gives the classical Gale–Shapley algorithm and shows that the LLP algorithm yields a more general version that handles additional constraints.

Chapter 4 discusses various sorting algorithms, including bubble sort, insertion sort, merge sort, quicksort, bitonic sort, and radix sort.

Chapter 5 discusses basic graph algorithms, including various traversal techniques (BFS, DFS), layering of directed acyclic graphs, and connected components.

Chapter 6 covers greedy algorithms: interval scheduling, interval partitioning, scheduling to minimize lateness, and Huffman coding.

Chapter 7 discusses algorithms for computing the shortest path in a graph, including Dijkstra’s algorithm, Bellman–Ford, Floyd–Warshall, Johnson’s algorithm, and delta-stepping. Both classical and LLP formulations are presented.

Chapter 8 describes algorithms for computing the minimum spanning tree in a weighted undirected graph. By considering different lattices on the set of edges, one can derive classical Kruskal’s, Prim’s, and Borůvka’s algorithms.

Chapter 9 discusses the divide-and-conquer strategy, with applications to closest pair, counting inversions, convex hull, Karatsuba multiplication, Strassen matrix multiplication, and the FFT.

Chapter 10 discusses dynamic programming. Problems include the longest increasing subsequence, the optimal binary search tree, the knapsack problem, weighted interval scheduling, and segmented least squares. All of these are shown to be solvable using the LLP algorithm.

Chapter 11 discusses network flow, including the Ford–Fulkerson and Edmonds–Karp algorithms, the max-flow min-cut theorem, and the lattice of min-cuts.

Chapter 12 describes bipartite matching, including augmenting-path algorithms and maximum antichain computation via LLP.

Chapter 13 describes the theory of NP-completeness. This theory relates the difficulty of solving one problem to that of another and includes reductions among SAT, Clique, Vertex Cover, Hamiltonian Path, and Subset Sum. A standard course in algorithms may cover all the chapters up to this point. The remaining chapters may be covered depending on the interest of the instructor.

Chapter 15 discusses the housing allocation problem, a one-sided version of the stable marriage problem solved via the top-trading-cycles algorithm and its LLP formulation.

Chapter 16 discusses the assignment problem (weighted bipartite matching), with LLP algorithms for computing market-clearing prices.

Chapter 17 discusses subclasses of Boolean satisfiability — Horn formulas and 2-SAT — whose satisfiability can be evaluated efficiently, including an LLP formulation for Horn-SAT.

Chapter 18 discusses number-theoretic algorithms from the lens of LLP: GCD, extended GCD, least common multiple, the Chinese Remainder Theorem, and computing the multiplicative order.

Chapter 19 describes linear programming, another general algorithm design technique. We show that many problems described using the LLP paradigm can also be solved using linear programming.

1.8 Summary

This chapter provided an introduction to algorithms and their analysis. It covered the definition of an algorithm, asymptotic notation for characterizing growth rates, and efficiency analysis of several fundamental algorithms. The chapter also reviewed essential data structures—linked lists, stacks, queues, binary search trees, and heaps—that are used throughout the book.

Algorithm	Problem	Time Complexity
Euclid’s GCD	Greatest Common Divisor	$O(\log(\min(a, b)))$
Linear Search	Search in unsorted array	$O(n)$
Binary Search	Search in sorted array	$O(\log n)$
Tower of Hanoi	Move n disks	$O(2^n)$
Build-Heap	Build a min-heap from array	$O(n)$

1.9 Problems

- Prove or disprove each of the following:
 - $3^n = O(2^n)$
 - If $f(n) = O(g(n))$, then $2^{f(n)} = O(2^{g(n)})$
 - If $f(n) = O(g(n))$, then $f(n)^3 = O(g(n)^3)$
- Prove the following statements.
 - Prove that for any constant $d > 0$, $f(n) = O(d \cdot g(n))$ iff $f(n) = O(g(n))$.
 - Prove that if $f(n) = O(h_1(n))$ and $g(n) = O(h_2(n))$ then $f(n) + g(n) = O(\max\{h_1(n), h_2(n)\})$.
 - Prove that all logarithms are asymptotically equivalent. That is, prove that $\log_a(n) = O(\log_b(n))$ for any positive a and b .
- Show that the time complexity of `BuildHeap` is $O(n)$.
- Suppose that you have a min Heap of size n . Give the pseudo-code to find and delete the k^{th} smallest element in the heap. What is the time complexity of your algorithm in terms of n and k ? Assume that you have a method available to you for a Heap that can insert any element in the heap in $O(\log n)$ time.
- Suppose that we have an array A of n *distinct* integers in two forms: a min Heap H and a max heap K . We are given the following functions on the min-heap:


```
H.getMin() // return the least value in the heap H
H.deleteMin() // delete the least value from the heap H
```

 Similarly, we have analogous methods for the max-heap. Give pseudo-code of an efficient program to return a value $A[i]$ in the array A such that there exists j where $A[i] + A[j]$ equals m . If there are no two such distinct entries, then the program should return null. Give the tight asymptotic time complexity of your method.
- Rank the following functions by growth rate from slowest to fastest. For each consecutive pair in your ordering, state whether the relationship is O , Θ , or neither.

$$n^2, \quad 2^n, \quad n \log n, \quad n!, \quad \log n, \quad n^{1.5}, \quad n \log^2 n$$
- Solve the recurrence $T(n) = 2T(n/2) + n$ with $T(1) = 1$ and prove that $T(n) = \Theta(n \log n)$.
- Given an unsorted array A of n integers and an integer $k \leq n$, describe an algorithm that returns the k largest elements in A in $O(n \log k)$ time using a min-heap of size k . Prove the time bound.
- Prove the correctness of Algorithm `BinarySearch` (Algorithm `BinarySearch`) using a loop invariant. State the invariant precisely and show that it holds at initialisation, is maintained by each iteration, and implies correctness at termination.
- Let A be an array of n integers and let S be an initially empty stack. Consider the following algorithm: for $i = 1$ to n , push $A[i]$ onto S . After each push, while S is not empty and the top two elements of S are equal, pop both and push their sum. What does S contain when the algorithm finishes on input $A = [2, 2, 4, 8, 2, 2, 4]$? What is the worst-case time complexity of the algorithm? Justify your answer using an amortized argument.

11. Implement both versions of Euclid's GCD algorithm in Java: (a) the subtraction-based version (Algorithm GCD from this chapter), and (b) a version that uses the *mod* operator instead of repeated subtraction. Run both on the input pair $(a, b) = (10^9 + 7, 10^8 + 7)$ and compare the number of iterations each version requires. Report the ratio.
12. Implement a min-heap in Java with the following operations: `insert(int key)`, `extractMin()`, `buildHeap(int[] A)`. Use a 1-based array as the underlying storage. Test your implementation by inserting the values `[15, 10, 20, 17, 8, 25, 18, 5]` one at a time, then extracting all elements in order; verify that they come out sorted. As a second test, build a heap from the same array in $O(n)$ time using `buildHeap` and verify the output is the same.
13. Implement Merge Sort in Java. Instrument your code to count the exact number of element comparisons made during the sort. Run it on random arrays of sizes $n = 10^3, 10^4, 10^5$, and 10^6 . For each n , report the measured comparison count and the ratio $\text{count}/(n \log_2 n)$. Is the ratio approximately constant?
14. Given an array A of n integers, the *next greater element* of $A[i]$ is the first element to the right of $A[i]$ that is strictly larger than $A[i]$, or -1 if no such element exists. Implement a Java program that computes the next-greater-element array in $O(n)$ time using a stack. Test on $A = [4, 5, 2, 25, 7, 8, 6, 1]$ and verify the output is `[5, 25, 25, -1, 8, -1, -1, -1]`.

1.10 Bibliographic Remarks

For a discussion of the asymptotic notation of the algorithm time complexity, we refer the reader to Knuth [Knu97a]. Heaps were introduced by J.W.J. Williams in 1964 as part of the heapsort algorithm, which leverages the heap structure for efficient sorting in $O(n \log n)$ time. The linear-time Build-Heap construction was later analyzed by Floyd, showing its $O(n)$ complexity. Heaps have since become foundational in priority queue implementations, with applications in scheduling, graph algorithms (e.g., Dijkstra's), and data compression (e.g., Huffman coding).

Chapter 2

Lattice Linear Predicate Detection

2.1 Introduction

Many classical combinatorial optimization problems — stable marriage, shortest paths, minimum spanning trees, market-clearing prices, housing allocation — appear at first glance to require entirely different algorithmic techniques. The Gale–Shapley algorithm proposes and rejects; Dijkstra’s algorithm relaxes edges in priority order; Borůvka’s algorithm merges components. Yet a closer look reveals a shared structure: in every case, the set of feasible solutions is closed under a natural “meet” operation, and the optimal solution is the *least* element of a *distributive lattice* (see Appendix 21 for background on lattices) that satisfies a Boolean predicate expressing feasibility.

In this chapter, we formalize this observation as the Lattice-Linear Predicate (LLP) algorithm. The framework has three ingredients:

1. A *distributive lattice* L that models the search space. Each element of L is a candidate solution represented as a vector of component values.
2. A Boolean *predicate* B on L that captures the goal: $B(G)$ is true when G is a feasible (or optimal) solution.
3. A *lattice-linearity* condition on B : whenever $B(G)$ is false, at least one component j of G is *forbidden* — meaning that no solution above G with the same value of $G[j]$ can satisfy B . Advancing that component brings G closer to a feasible solution.

Once these ingredients are in place, a single generic algorithm repeatedly advances forbidden components until the predicate is satisfied (or the top of the lattice is reached, signaling that no solution exists). The algorithm is *correct by construction*: lattice-linearity guarantees that every advance makes irrevocable progress, so the algorithm terminates at the unique least fixpoint.

The practical payoff is substantial. The LLP framework unifies a wide range of problems that are traditionally taught with separate algorithms:

In each case, the classical algorithm emerges as a *specific schedule* of the generic LLP algorithm — a particular order in which forbidden components are advanced. Different schedules yield different classical

Problem	Classical algorithm	LLP lattice
Stable marriage	Gale–Shapley	proposal vectors
Single-source shortest path	Dijkstra, Bellman–Ford	distance vectors
Minimum spanning tree	Kruskal, Prim, Borůvka	edge-selection vectors
Market-clearing prices	Demange–Gale–Sotomayor	price vectors
Housing allocation	Top-trading cycles	allocation vectors
Horn satisfiability	Unit propagation	truth-assignment vectors
Job scheduling	Critical-path method	completion-time vectors

Table 2.1: Problems unified by the LLP framework. Each classical algorithm is a specific schedule of the generic LLP algorithm on the corresponding lattice.

algorithms for the same problem (for example, Dijkstra’s algorithm and Bellman–Ford are two schedules of the same LLP instance; delta-stepping interpolates between them).

A key advantage of the LLP viewpoint is *composability*: because the conjunction of two lattice-linear predicates is itself lattice-linear (Lemma 2.3(c)), one can add side constraints to any of the above problems — forced or forbidden pairs in stable marriage, fairness constraints in shortest paths, budget caps in market-clearing prices — and the same algorithm solves the constrained version with no additional machinery.

This chapter is organized as follows. Section 2.2 formally defines the lattice-linear predicates. These predicates are defined on a distributive lattice. The problem is framed as the search for an element in the lattice that satisfies the predicate. Generally, we are interested in the least element in the lattice that satisfies the predicate. Section 2.3 gives the notation that we use for programs based on lattice-linear predicates. It specifies the lattice that we are working on, the starting element in the lattice, the top element of the lattice, and the predicate *forbidden*, which allows us to advance in the lattice. Section 2.5 lists some desirable properties of the LLP algorithm. The algorithm is naturally *nondeterministic*. There are multiple ways in which the algorithm can advance in the lattice. The algorithm can advance on multiple components simultaneously. The algorithm is *online*. It does not inspect any element higher than the current element of the lattice that is under consideration. Finally, the algorithm returns an optimal answer for all components. Section 2.9 gives the dual of the lattice-linearity property. Many problems such as the stable marriage problem satisfy not only lattice-linearity but also its dual. For such predicates, one can also start from the top element of the lattice and traverse it downwards looking for an element that satisfies the given predicate. When applied to the stable marriage problem, one can find the woman-optimal stable marriage in this manner.

2.2 Lattice-Linear Predicates

Let L be the lattice of all n -dimensional vectors of reals greater than or equal to zero vector and less than or equal to a given vector T where the order on the set of vectors is defined by the natural componentwise \leq . The minimum element of this lattice is the zero vector. The lattice is used to model the search space of the combinatorial optimization problem. For simplicity, we are considering the lattice of vectors of nonnegative reals; later, we show that our results are applicable to any distributive lattice. The combinatorial optimization problem is modeled as finding the minimum element in L that satisfies a boolean *predicate* B , where B models *feasible solutions*. We are interested in algorithms to solve the combinatorial optimization problem with n processes. We will assume that the system maintains as its state the current candidate vector $G \in L$ in the search lattice. Even if we are using a sequential algorithm,

we say that $G[i]$ is maintained in the process i . We call G , the global state, and $G[i]$, the state of process i .

Finding an element in the lattice that satisfies the given predicate B is called the *predicate detection problem*. Finding the *minimum* element that satisfies B (whenever it exists) is the combinatorial optimization problem. We now define *lattice-linearity*, which enables efficient computation of this minimum element. Lattice-linearity was originally introduced in the context of detecting global conditions in a distributed system, where it is simply called linearity. We use the term *lattice-linearity* to avoid confusion with the standard usage of linearity. A key concept in the development of an efficient predicate detection algorithm is that of a *forbidden* state. Given a predicate B and a vector $G \in L$, a state $G[i]$ is *forbidden* (or equivalently, the index i is forbidden) if for any vector $H \in L$, where $G \leq H$, if $H[i]$ equals $G[i]$, then B is false for H . Formally,

Definition 2.1 (Forbidden State) *Given any distributive lattice L of n -dimensional vectors of $\mathbf{R}_{\geq 0}$, and a predicate B , we define $\text{forbidden}(G, i, B) \equiv \forall H \in L : G \leq H : (G[i] = H[i]) \Rightarrow \neg B(H)$.*

We define a predicate B as *lattice-linear* with respect to a lattice L if, for any global state G , B is false in G implies that G contains a *forbidden state*. Formally,

Definition 2.2 (Lattice-Linear Predicate) *A boolean predicate B is lattice-linear with the polytime algorithm \mathcal{A} with respect to a finite distributive lattice L generated from a poset P iff $\forall G \in L : \neg B(G) \Rightarrow (\exists i \in \mathcal{A}(G) : \text{forbidden}(G, i, B))$.*

The algorithm \mathcal{A} is efficient in the size of the poset P . It tells us which indices are forbidden. From now on, the algorithm \mathcal{A} is implicit. We now give some examples of lattice-linear predicates.

1. **Job Scheduling Problem:** Our first example relates to scheduling of n jobs. Each job j requires time t_j for completion and has a set of prerequisite jobs, denoted by $\text{pre}(j)$, such that it can be started only after all its prerequisite jobs have been completed. Our goal is to find the minimum completion time for each job. We let our lattice L be the set of all possible completion times. A completion vector $G \in L$ is feasible iff $B_{\text{jobs}}(G)$ holds where $B_{\text{jobs}}(G) \equiv \forall j : (G[j] \geq t_j) \wedge (\forall i \in \text{pre}(j) : G[j] \geq G[i] + t_j)$. B_{jobs} is lattice-linear because if it is false, then there exists j such that either $G[j] < t_j$ or $\exists i \in \text{pre}(j) : G[j] < G[i] + t_j$. We claim that $\text{forbidden}(G, j, B_{\text{jobs}})$. Indeed, any vector $H \geq G$ cannot be feasible with $H[j]$ equal to $G[j]$. The minimum of all vectors that satisfy feasibility corresponds to the minimum completion time.
2. **Prefix Sum Problem:** Our second example relates to (exclusive) prefix sum of an array A with non-negative reals. We are required to output an array G such that $G[j]$ equals sum of all entries in A from 0 to $j - 1$. We define G to be feasible iff B_{prefix} holds where $B_{\text{prefix}} \equiv (\forall j > 0) : (G[j] \geq G[j - 1] + A[j - 1])$. Again, it is easy to verify that B_{prefix} is lattice-linear. The minimum vector G that satisfies B_{prefix} corresponds to the exclusive prefix sum of the array A .
3. **Continuous Optimization Problem:** We are required to find minimum nonnegative x and y such that $B \equiv (x \geq y^2/4 + 5) \wedge (y \geq x - 4)$. We view this problem as finding the minimum (x, y) pair such that B holds. It is easy to verify that B is lattice-linear. If the first conjunct is false, then x is forbidden. Unless x is increased, the predicate cannot become true, even if other variables (y for

this example) increase. If the second conjunct is false, then y is forbidden. In this case, $x = 6$ and $y = 2$ is the pointwise minimum solution. For some predicates, there may not be any solution. For example, when $B \equiv (x \geq 2y^2 + 5) \wedge (y \geq x - 4)$, there is no nonnegative (x, y) pair such that B holds (verify this!). The predicate B is still lattice-linear, but by advancing along x and y we go beyond any bounded (x, y) .

4. **A Non Lattice-Linear Predicate:** As an example of a predicate that is not lattice-linear, consider the predicate $B \equiv \sum_j G[j] \geq 1$ defined on the space of two-dimensional vectors. Consider the vector G equal to $(0, 0)$. The vector G does not satisfy B . For B to be lattice-linear, either the first index or the second index should be forbidden. However, none of the indices are forbidden in $(0, 0)$. The index 0 is not forbidden because the vector $H = (0, 1)$ is greater than G , has $H[0]$ equal to $G[0]$ but it still satisfies B . The index 1 is also not forbidden because $H = (1, 0)$ is greater than G , has $H[1]$ equal to $G[1]$ but it satisfies B .

The following lemma is useful for proving lattice-linearity of predicates.

Lemma 2.3 *Let B be any Boolean predicate defined on a lattice L of vectors.*

(a) *Let $f : L \rightarrow \mathbf{R}_{\geq 0}$ be any monotone function defined on the lattice L of vectors of $\mathbf{R}_{\geq 0}$. Consider the predicate $B \equiv G[i] \geq f(G)$ for some fixed i . Then, B is lattice-linear.*

(b) *Let L_B be the subset of the lattice L of the elements that satisfy B . Then, B is lattice-linear implies that L_B is closed under meets.*

(c) *If B_1 and B_2 are lattice-linear then $B_1 \wedge B_2$ is also lattice-linear.*

Proof: (a) Suppose B is false for G . This implies that $G[i] < f(G)$. Consider any vector $H \geq G$ such that $H[i]$ is equal to $G[i]$. Since $G[i] < f(G)$, we get that $H[i] < f(G)$. The monotonicity of f implies that $H[i] < f(H)$ which shows that $\neg B(H)$.

(b) We show the contrapositive. Let $Y = \{H_1, H_2, \dots, H_k\}$ be any subset of L_B such that its infimum G does not belong to L_B . Since G is the infimum of Y , for any i , there exists $j \in \{1 \dots k\}$ such that $G[i] = H_j[i]$. Since $B(H_j)$ is true for all j , it follows that there exists a G for which lattice-linearity does not hold. None of the indices in G are forbidden. (c) Suppose that $\neg(B_1 \wedge B_2)$. This implies that one of the conjuncts is false and therefore, from the lattice-linearity of that conjunct, a forbidden state exists. ■

For the job scheduling example, we define B_j as $G[j] \geq \max(t_j, \max\{G[i] + t_j \mid i \in \text{pre}(j)\})$. Since $f_j(G) = \max(t_j, \max\{G[i] + t_j \mid i \in \text{pre}(j)\})$ is a monotone function, it follows from Lemma 2.3(a) that B_j is lattice-linear. The predicate $B_{j_{obs}} \equiv \forall j : B_j$ is lattice-linear due to Lemma 2.3(c). Also note that the problem of finding the minimum vector that satisfies $B_{j_{obs}}$ is well-defined due to Lemma 2.3(b).

We now discuss detection of lattice-linear predicates which requires an additional assumption called the *efficient advancement property* — there exists an efficient (polynomial time) algorithm to determine the forbidden state. This property holds for all the problems considered in this book. Once we determine j such that $\text{forbidden}(G, j, B)$, we also need to determine how to advance along the index j . To that end, we extend the definition of forbidden as follows.

Definition 2.4 (α -forbidden) Let B be any Boolean predicate on the lattice L of all assignment vectors. For any G, j and positive real $\alpha > G[j]$, we define $\text{forbidden}(G, j, B, \alpha)$ iff

$$\forall H \in L : H \geq G : (H[j] < \alpha) \Rightarrow \neg B(H).$$

Given any lattice-linear predicate B , suppose $\neg B(G)$. This means that G must be advanced on all indices j such that $\text{forbidden}(G, j, B)$. We use a function $\alpha(G, j, B)$ such that $\text{forbidden}(G, j, B, \alpha(G, j, B))$ holds whenever $\text{forbidden}(G, j, B)$ is true. With the notion of $\alpha(G, j, B)$, we have the algorithm *LLP* shown in Fig. [LLP](#). The algorithm *LLP* has two inputs, the predicate B and the top element of the lattice T . It returns the least vector G which is less than or equal to T and satisfies B (if it exists). Whenever B is not true in the current vector G , the algorithm advances on all forbidden indices j in parallel. This simple algorithm can be used to solve a wide variety of combinatorial optimization problems by instantiating different $\text{forbidden}(G, j, B)$ and $\alpha(G, j, B)$.

Algorithm LLP: Algorithm *LLP* to find the minimum vector at most T that satisfies B

Input: T : vector (top element), B : predicate

Output: the minimum vector $G \leq T$ that satisfies B , or null

```

1  $G$ : vector of reals initially  $\forall i : G[i] = 0$ ;
2 while  $\exists j : \text{forbidden}(G, j, B)$  do
3   forall  $j$  such that  $\text{forbidden}(G, j, B)$  in parallel do
4     if  $\alpha(G, j, B) > T[j]$  then return null;
5     else  $G[j] := \alpha(G, j, B)$ ;
6   end
7 end
8 return  $G$  ;
```

// the optimal solution

Theorem 2.5 Suppose that there exists a fixed constant $\delta > 0$ such that $\alpha(G, j, B) - G[j] \geq \delta$ whenever $\text{forbidden}(G, j, B)$. Then, the algorithm *LLP* finds the least vector $G \leq T$ that satisfies B , if one exists.

Proof: Since $G[j]$ increases by at least δ for at least one forbidden j in every iteration of the *while* loop, the algorithm terminates in at most $\sum_i \lceil T[i]/\delta \rceil$ number of steps.

We show that the algorithm maintains the invariant (I1) that for all indices j , any vector V such that $V[j]$ is less than $G[j]$ cannot satisfy B . Formally, the invariant (I1) is

$$\forall j : (\forall V \in L : (V[j] < G[j]) \Rightarrow \neg B(V)).$$

Initially, the invariant holds trivially because G is initialized to 0. Suppose $\text{forbidden}(G, j, B)$. Then, we increase $G[j]$ to $\alpha(G, j, B)$. We need to show that this change maintains the invariant. Pick any V such that $V[j] < \alpha(G, j, B)$. We now do a case analysis. If $V \geq G$, then $\neg B(V)$ holds from the definition of $\alpha(G, j, B)$. Otherwise, there exists some k such that $V[k] < G[k]$. In this case, $\neg B(V)$ holds due to (I1).

We can now show Theorem 2.5 using the invariant. First, suppose that the algorithm *LLP* terminates because $\alpha(G, j, B) > T[j]$. In this case, there is no feasible vector in L due to the invariant (because the

predicate B is false for all values of $G[j]$. Now suppose that the algorithm terminates because there does not exist any j such that $forbidden(G, j, B)$. This implies that G satisfies B due to lattice-linearity of B . It is also the least vector that satisfies B due to the invariant (I1). ■

For the job scheduling example, we get an algorithm to find the minimum completion time using $forbidden(G, j, B_{jobs}) \equiv (G[j] < t_j) \vee (\exists i \in pre(j) : G[j] < G[i] + t_j)$, and $\alpha(G, j, B_{jobs}) = \max\{t_j, \max\{G[i] + t_j \mid i \in pre(j)\}\}$.

For the prefix sum example, we get an algorithm using $forbidden(G, j, B_{prefix}) \equiv (G[j] < G[j - 1] + A[j - 1])$ and $\alpha(G, j, B_{prefix}) = G[j - 1] + A[j - 1]$ for all $j > 0$.

We now show, on account of Lemma 2.3(c), that if we have an algorithm for a problem, then we also have one for the constrained version of that problem.

Lemma 2.6 *Let LLP be the algorithm to find the least vector G that satisfies B_1 if one exists. Then, LLP can be adapted to find the least vector G that satisfies $B_1 \wedge B_2$ for any lattice-linear predicate B_2 .*

Proof: The algorithm LLP can be used with the following changes:
 $forbidden(G, j, B_1 \wedge B_2) \equiv forbidden(G, j, B_1) \vee forbidden(G, j, B_2)$, and
 $\alpha(G, j, B_1 \wedge B_2) = \max\{\alpha(G, j, B_1), \alpha(G, j, B_2)\}$. ■

For example, suppose that we want the minimum completion time of jobs with the additional lattice-linear constraint that $B_2(G) \equiv (G[1] = G[2])$. B_2 is lattice-linear with $forbidden(G, 1, B_2) \equiv (G[1] < G[2])$ and $forbidden(G, 2, B_2) \equiv (G[2] < G[1])$. Applying Lemma 2.6, we get an algorithm for the constrained version.

2.3 LLP Notation

We now describe the notation used in LLP algorithms. Fig. 2.1 shows LLP algorithms for the job scheduling problem. We have a single variable G in all the examples shown in Fig. 2.1.

All other variables are derived directly or indirectly from G . G is an array of objects such that $G[j]$ is the state of component j . There are three sections of the program.

The **var section** declares the state variables and their initial values. For example, the *var* section of the job scheduling program in Fig. 2.1 specifies that $G[j]$ is an integer initially equal to $t[j]$.

The **always section** defines additional variables that are derived from G . The actual implementation of these variables is left to the system. They can be viewed as macros.

The third section gives the desirable predicate, either by using the **forbidden** predicate or the **ensure** predicate. The *forbidden* predicate has an associated *advance clause* that specifies how $G[j]$ must be advanced whenever the forbidden predicate is true. For many problems, it is more convenient to use the complement of the forbidden predicate. The *ensure section* specifies the desirable predicates of the form $(G[j] \geq expr)$ or $(G[j] \leq expr)$. The statement *ensure* $G[j] \geq expr$ simply means that whenever component j finds $G[j]$ to be less than $expr$, it advances $G[j]$ to $expr$. Since $expr$ may refer to G , just by setting $G[j]$ equal to $expr$, there is no guarantee that $G[j]$ continues to be equal to $expr$ — the value of

$expr$ may change due to changes in other components. We use the *ensure* statement whenever $expr$ is a monotonic function of G and therefore the predicate is lattice-linear.

(a) Using **forbidden/advance**:

Algorithm LLP-JobScheduling-Forbidden: LLP Program for LLP-JobScheduling-Forbidden

Input: t : array of int; pre : array of sets of int

Output: G : array of int, initially $t[i]$

- 1 **forbidden**(j): $G[j] < \max\{G[i] + t[j] \mid i \in pre(j)\}$;
 - 2 **advance**: $G[j] := \max\{G[i] + t[j] \mid i \in pre(j)\}$;
-

(b) Using **ensure** clause:

Algorithm LLP-JobScheduling-Ensure: LLP Program for LLP-JobScheduling-Ensure

Input: t : array of int; pre : array of sets of int

Output: G : array of int, initially $t[i]$

- 1 **ensure**(j): $G[j] \geq \max\{G[i] + t[j] \mid i \in pre(j)\}$;
-

Figure 2.1: LLP Program for the job scheduling problem: (a) using the *forbidden/advance* notation, (b) using the compact *ensure* clause. Both are equivalent; the *ensure* form is used when the advance expression is a monotone function of G .

Consider four jobs with execution times $t = (3, 2, 4, 1)$ and prerequisites $pre(2) = \{1\}$, $pre(3) = \{1\}$, $pre(4) = \{2, 3\}$ (job 1 has no prerequisites). The LLP algorithm initializes $G = (3, 2, 4, 1)$. Depending on the evaluation order, a job may need to be advanced more than once.

Example 2.7 Suppose we evaluate job 4 before job 3:

1. Jobs 1 not forbidden. Advance job 2: $G = (3, 5, 4, 1)$.
2. Job 4: requires $\max(G[2] + 1, G[3] + 1) = \max(6, 5) = 6$, but $G[4] = 1$. Advance: $G[4] := 6$. Now $G = (3, 5, 4, 6)$.
3. Job 3: requires $G[3] \geq G[1] + 4 = 7$, but $G[3] = 4$. Advance: $G[3] := 7$. Now $G = (3, 5, 7, 6)$.
4. Job 4 is forbidden *again*: requires $\max(6, 8) = 8$, but $G[4] = 6$. Advance: $G[4] := 8$. Now $G = (3, 5, 7, 8)$.

Job 4 was advanced *twice* because it was processed before its prerequisite job 3 was settled.

An Efficient Reformulation Using Fixed Indices

To avoid such repeated advances, we reformulate the forbidden predicate using a *fixed* array. Let $fixed[j]$ be a boolean, initially false for all j . A job j is forbidden if $\neg fixed[j] \wedge (\forall i \in pre(j) : fixed[i])$. When j is forbidden, we advance $G[j] := \max\{G[i] + t[j] \mid i \in pre(j)\}$ and set $fixed[j] := true$. Since all predecessors

are already fixed (their values will not change), this advance is final — each job becomes forbidden at most once. Algorithm [LLP-JobScheduling-Fixed](#) gives the LLP program.

Algorithm LLP-JobScheduling-Fixed: LLP Program for LLP-JobScheduling-Fixed

Input: t : array of int; pre : array of sets of int

Output: G : array of int, initially $t[i]$; $fixed$: array of boolean, initially *false*

- 1 **forbidden**(j): $\neg fixed[j] \wedge \forall i \in pre(j) : fixed[i]$;
 - 2 **advance**: $G[j] := \max\{G[i] + t[j] \mid i \in pre(j)\}$; $fixed[j] := true$;
-

The efficiency of the algorithm may depend on the formulation of $\alpha(G, j, B)$. We will see such optimizations for many problems later in the book, including the shortest-path algorithm.

Example 2.8 Using the same four-job instance, we trace Algorithm [LLP-JobScheduling-Fixed](#). Initialize $G = (3, 2, 4, 1)$ and $fixed = (false, false, false, false)$.

1. Job 1: $pre(1) = \emptyset$, so all predecessors are vacuously fixed. Forbidden. Advance: $G[1] := 3$ (unchanged); $fixed[1] := true$. $G = (3, 2, 4, 1)$.
2. Job 2: $pre = \{1\}$ and $fixed[1]$ is true. Forbidden. Advance: $G[2] := G[1] + t[2] = 5$; $fixed[2] := true$. $G = (3, 5, 4, 1)$.
3. Job 3: $pre = \{1\}$ and $fixed[1]$ is true. Forbidden. Advance: $G[3] := G[1] + t[3] = 7$; $fixed[3] := true$. $G = (3, 5, 7, 1)$.
4. Job 4: $pre = \{2, 3\}$ and both are fixed. Forbidden. Advance: $G[4] := \max(6, 8) = 8$; $fixed[4] := true$. $G = (3, 5, 7, 8)$. All jobs fixed. Done.

Each job is advanced exactly once, giving $O(n + m)$ total work (where $m = |\bigcup_j pre(j)|$).

The *fixed* predicate ensures that a job is only processed after all its prerequisites are settled, eliminating repeated advances.

Fig. 2.2 illustrates the difference between the two formulations, projected onto the $(G[3], G[4])$ plane (since $G[1] = 3$ and $G[2] = 5$ are settled early in both traces). A state (g_3, g_4) is feasible when $g_3 \geq G[1] + t[3] = 7$ and $g_4 \geq \max(G[2] + t[4], g_3 + t[4]) = \max(6, g_3 + 1)$. The feasible states (shaded) are $(7, 8)$, $(7, 9)$, and $(8, 9)$; the least feasible state is $(7, 8)$.

2.4 A Sequential Implementation of the LLP Algorithm

The LLP algorithm can be implemented sequentially by maintaining a set S of forbidden indices. Algorithm [LLP-Sequential](#) gives LLP-SEQ, a general sequential implementation.

The algorithm maintains the invariant: **(I1)** S empty implies G is the optimal solution. The set S need not contain all forbidden indices at every step; it may also contain non-forbidden indices, which are simply skipped. The key implementation choices are:

- *Storage for S* : a linked list, queue, stack, or heap depending on the removal strategy. For the shortest path problem, a priority queue keyed by $G[j]$ is efficient.

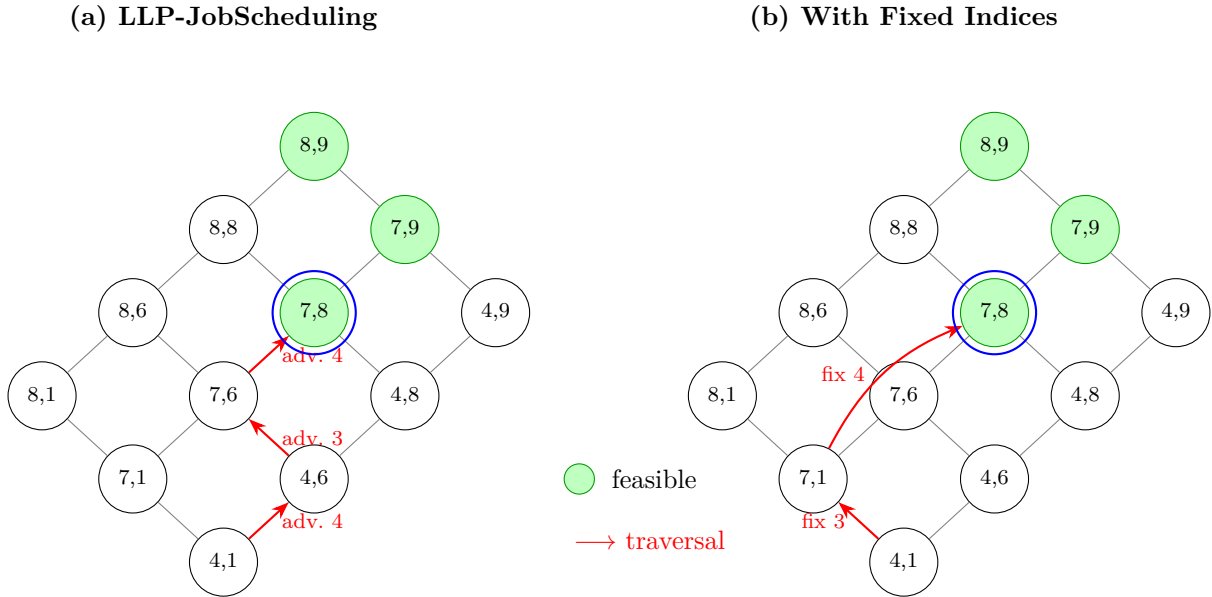


Figure 2.2: LLP traversal for the 4-job scheduling example, projected onto $(G[3], G[4])$ since $G[1] = 3$ and $G[2] = 5$ settle early. The feasible states $\{(7, 8), (7, 9), (8, 9)\}$ are shaded; the least is $(7, 8)$. (a) Basic formulation (bad evaluation order): advance job 4 to $(4, 6)$, then job 3 to $(7, 6)$, then job 4 *again* to $(7, 8)$. Job 4 is advanced twice. (b) With fixed indices: fix job 3 first (reaching $(7, 1)$), then fix job 4 which advances directly to $(7, 8)$. Each job advances once.

Algorithm LLP-Sequential: LLP-SEQ: A sequential algorithm to find the minimum vector $\leq T$ that satisfies B .

Input: T : vector (top element), B : predicate

Output: the minimum vector $G \leq T$ that satisfies B , or null

1 **var** G : vector initially $\forall i : G[i] := 0$;

2 **var** S : set of indices;

3 *initialize*(S, G) ;

// add initial forbidden indices to S

4 **while** $\neg S.empty()$ **do**

5 $j := S.remove(G)$;

6 **if** *forbidden*(G, j, B) **then**

7 $G[j] := \alpha(G, j, B)$;

8 **if** $G[j] > T[j]$ **then return** null;

9 *updateForbidden*(G, j);

10 **end**

11 **end**

12 **return** G

- *Initialization*: insert all indices into S , or only those initially forbidden. For job scheduling with acyclic prerequisites, initialize with jobs having no predecessors.
- *Removal order*: simple FIFO, or priority-based (e.g., smallest $G[j]$ first for shortest path, ensuring indices are fixed once processed).
- *Updating S* : use a dependency list $dependList(j)$ listing indices whose forbidden status may change when $G[j]$ changes. For graph problems, these are typically the neighbors of j .

The while loop executes at most $O(mn)$ times, since each of n components advances at most m times (where m is the height of a single chain in the lattice). If each iteration costs $O(n)$ for remove, advance, and update, the total is $O(mn^2)$; for problems where these operations are $O(1)$, the total is $O(mn)$.

2.5 Properties of the LLP Algorithm

The LLP algorithm has many useful properties. We list them here so that the reader can apply them to various problems studied in this book. These properties are applicable to all the problems for which the LLP algorithm is used. We illustrate them with the job scheduling problem; in Chapter 3, we reinforce them in the context of the stable marriage problem.

1. **Nondeterminism in Evaluation of Forbidden Predicate**: Given a global state G , there may be multiple indices j for which $G[j]$ is forbidden. The LLP algorithm is correct irrespective of the order in which these indices are updated. The efficiency may differ depending upon the order, but the correctness is independent of the order. For example, in the job scheduling problem, the minimum completion time vector is the same regardless of the order in which forbidden jobs are advanced.
2. **Parallel Evaluation of Forbidden Predicate**: Suppose that G is shared among different threads, such that thread j is responsible for evaluating $forbidden(G, j)$. While this thread is evaluating the predicate, other threads may have advanced on other indices, i.e., thread j may have old information of $G[i]$ for $i \neq j$. However, this would still keep the algorithm correct. For example, in job scheduling, multiple forbidden jobs can evaluate their prerequisites and advance their completion times simultaneously, even with stale values of G .
3. **No Lookahead Required for Evaluation of Forbidden Predicate**: The LLP algorithm determines whether an index j is forbidden depending upon only the current global state G . This means that these algorithms are applicable in online settings where the future part of the lattice is revealed only when a forbidden index needs to advance. For job scheduling, the forbidden predicate for job j depends only on the current completion times of its prerequisites — not on jobs that may become forbidden later. For some problems, such as the shortest path problem, the weights on the edges are required to evaluate the forbidden predicate, so the graph must be known and static.
4. **The Optimal Feasible Global State is Optimal for Individual Components**: Suppose that in the job scheduling problem, we are interested in the minimum completion time for a single job j , ignoring all other jobs. The LLP algorithm returns a vector G_{llp} such that $G_{llp}[j]$ equals this individually optimal value. More generally, let G be the global state that is feasible and optimal with respect to index i , i.e., for all feasible H , $G[i] \leq H[i]$. Let G_{llp} be the global state computed by the LLP algorithm, then $G[i] = G_{llp}[i]$. This follows from the meet-closure property of feasible states. If

$G[i] < G_{llp}[i]$, then the global state given by $G \sqcap G_{llp}$ is also feasible and strictly smaller than G_{llp} , contradicting that G_{llp} is the least global state that satisfies the feasibility predicate.

2.6 Splittable Predicates

Let L be a finite distributive lattice and B be any lattice-linear predicate. We call B splittable if there exist two lattices L_1 and L_2 and two lattice-linear predicates B_1 and B_2 such that given the least solution for B_1 and B_2 in L_1 and L_2 , one can determine the least solution for B in L . The effort to determine the least solution for B in L is reduced if we have the solutions for B_1 and B_2 . If B_1 and B_2 are also splittable, then we can apply this strategy recursively. Once the lattice is trivial, i.e., it is totally ordered, then finding the least solution is simple. Formally,

Definition 2.9 (Splittable Lattice-Linear Predicate) *A nonempty Lattice-Linear Predicate B is splittable for a finite distributive lattice L , if there exist two nonempty lattices L_1 and L_2 and two lattice-linear predicates B_1 and B_2 such that there exists a map $f : L_1 \times L_2 \rightarrow L$, where $f(G_1, G_2)$ is the least element satisfying B in L such that G_1 and G_2 are the least elements in L_1 and L_2 satisfying B_1 and B_2 .*

As an example of a predicate that is not splittable, consider the stable marriage problem. Given the distributive lattice of all assignments, the predicate B that the assignment is a stable marriage is not splittable. The predicate B that is true on all elements is trivially splittable into B_1 and B_2 also as true elements. The lattice L can be split into arbitrary $L_1 \times L_2$. Nontrivial splittable predicates underlie the divide-and-conquer approach to algorithm design; examples such as Quicksort and Mergesort expressed via splittable predicates are developed in Chapter 9.

2.7 Additional Examples of LLP Algorithms

To further illustrate the generality of the LLP framework, we present three additional examples: computing the greatest common divisor (GCD) of an array, finding the maximum element, and linear search. Fig. 2.3 shows the LLP programs for these problems.

(a) GCD of an array:

Algorithm LLP-GCD: LLP Program for LLP-GCD

Input: A : array of int

Output: G : array of int, initially $A[i]$

```

1 forbidden( $j$ ):  $\exists i \in [1..n] : G[j] > G[i]$  ; // pick such  $i$ 
2 advance: if  $G[j] \bmod G[i] = 0$  then  $G[j] := G[i]$  else  $G[j] := G[j] \bmod G[i]$ ;

```

Figure 2.3: LLP Program for GCD.

GCD. For computing the GCD of an array of positive integers A , the lattice is ordered by componentwise \geq (decreasing order). Component j is forbidden if $G[j] > G[i]$ for some i . In this case, we replace $G[j]$ by $G[i]$ if $G[i]$ divides $G[j]$; otherwise, we replace $G[j]$ by $G[j] \bmod G[i]$. When the algorithm terminates, all

components are equal to the GCD of A . The invariant is that every common divisor of A is also a common divisor of G . This holds initially because $G = A$. After each advance, if $G[i]$ divides $G[j]$, then every divisor of $G[i]$ also divides $G[j]$, so setting $G[j] := G[i]$ preserves the invariant. If $G[i]$ does not divide $G[j]$, the common divisors of $G[i]$ and $G[j]$ are exactly those of $G[i]$ and $G[j] \bmod G[i]$.

2.8 The priority Tag

The *forbidden* and *advance* clauses of an LLP program specify *which* indices can be advanced and *how* to advance them, but they deliberately leave the *order* in which forbidden indices are processed unspecified. Every fair schedule that keeps advancing forbidden indices reaches the same least fixed point, so correctness is independent of the order. For efficiency, however, the order often matters. The LLP framework provides an optional **priority** tag that lets the designer influence the schedule without changing the predicate.

An LLP program may carry a third clause alongside *forbidden* and *advance*:

priority(j): *expression*

where *expression* is a real-valued (or ordinal-valued) function of the current state G and the index j . The semantics is:

*Whenever more than one index is forbidden, remove (and advance) the forbidden index j whose **priority**(j) is smallest first, breaking ties arbitrarily.*

Because the least-fixed-point solution is independent of the schedule, adding a **priority** clause never changes the final answer of a lattice-linear program; it can only change the running time and, in some cases, the number of advance steps. A typical implementation maintains a priority queue keyed on **priority**(j) that contains all currently forbidden indices. After each advance, any indices whose forbidden status may have changed (either newly forbidden, or newly not forbidden) are inserted into or removed from the queue. The choice of data structure (binary heap, bucket queue, Fibonacci heap) depends on the range and update pattern of the priority.

A useful guideline is to pick **priority**(j) to be the value to which $G[j]$ will be advanced — or a monotone function thereof. This ensures that indices whose target value is small are fixed first, so later advances do not disturb them.

We now give some example where the tag **priority** will come handy.

BFS and Dijkstra's algorithm. In the LLP formulation of BFS (Chapter 5), an index j is forbidden when some predecessor i has $G[j] > G[i] + 1$. Setting

priority(j) := $G[j]$

processes vertices in increasing order of their current distance. The first time a vertex j is dequeued, $G[j]$ already equals its final shortest-path distance — so j is advanced at most once. The schedule matches the classical BFS-with-queue algorithm. The same priority turns the LLP program for the single-source shortest path (Chapter 7) into Dijkstra's algorithm.

Dynamic programming. Many dynamic-programming recurrences have the form

$$G[j] = \min_{i \in \text{pre}(j)} (G[i] + w(i, j)),$$

on a DAG (e.g., shortest path in a DAG, longest increasing subsequence ending at each index, optimal binary search tree). Taking

$$\text{priority}(j) := \text{topological-order rank of } j$$

forces predecessors to be fixed before their successors, so every $G[j]$ is advanced exactly once — recovering the familiar “fill the DP table in topological order” schedule. Chapter 10 exploits exactly this pattern.

Layering a DAG. In the LLP program for layering (Chapter 5), j becomes forbidden when all its predecessors are fixed. Choosing

$$\text{priority}(j) := \max_{i \in \text{pre}(j)} G[i] + 1$$

processes vertices in order of their final layer number, giving a single $O(n + m)$ pass.

In every case, correctness is inherited from lattice-linearity; the `priority` clause only changes the order of evaluation and thus the running time. This separation of concerns — correctness from the predicate, efficiency from the schedule — is a recurring theme throughout the book.

2.9 Dual of Lattice-Linearity

We briefly discuss detection of predicates when the search starts from the top of the lattice in addition to the bottom of the lattice. Many predicates, such as $B_{\text{stableMarriage}}$ satisfy not only the lattice-linearity but also its dual. Thus, the set of elements satisfying these predicates is closed not only for the meet operation but also for the join operation. In addition, whenever the predicate is false, we can efficiently determine the forbidden indices. If we are okay with returning either of the elements as our final answer, then searching for any of the elements in parallel may speed up the algorithm by a factor of the height of the lattice. The dual of the lattice-linear property can be formally defined as follows.

Definition 2.10 (dual of Lattice-Linearity Property) *A Boolean predicate B has the dual of lattice-linearity property with the polytime algorithm \mathcal{A} with respect to a finite distributive lattice L generated from a poset P if*

$$\forall G \in L : \neg B(G) \Rightarrow \exists i \in \mathcal{A}(G) : \text{dual-forbidden}(i, G, B)$$

where $\text{dual-forbidden}(i, G, B)$ is defined as

$$\forall H \leq G : (H[i] = G[i]) \Rightarrow \neg B(H).$$

The above definition states that whenever B is false in G , the algorithm $\mathcal{A}(G)$ returns an index such that any global state H less than G that matches G on that index also has B false.

We next define a predicate B as a regular predicate when it satisfies lattice-linearity as well as its dual.

Definition 2.11 (Regular Predicate) *A predicate B is regular in a lattice L iff B is lattice-linear and also dual lattice-linear.*

From the definition of regular predicates, it follows that the subset of elements that satisfy B is a sublattice of L . Since a sublattice of a distributive lattice is also distributive, we conclude that the subset of elements in L satisfying B is also distributive.

For a regular predicate, one can search for the satisfying element starting from both the bottom and the top of the lattice, as shown in Algorithm [Regular](#).

Algorithm Regular: An Algorithm for detecting predicates with efficient advancement property and its dual.

Input: B : a regular predicate on lattice L
Output: an element of L satisfying B , or false

```

1  $G$ : element of  $L$  initially  $\perp$  (the bottom element of  $L$ );
2  $Z$ : element of  $L$  initially  $\top$  (the top element of  $L$ );
3 while  $\neg B(G)$  and  $\neg B(Z)$  do
4   forall  $i$ : forbidden( $i, G$ ) do
5     if  $G$  cannot be advanced on  $i$  then return false;
6     else advance  $G$  on  $i$ ;
7   end
8   forall  $j$ : dual-forbidden( $j, Z$ ) do
9     if  $Z$  cannot be retreated on  $j$  then return false;
10    else retreat  $Z$  on  $j$ ;
11  end
12 end
13 if  $B(G)$  then return  $G$ ;
14 else return  $Z$  ; // an optimal solution;
```

Applying the idea to the stable marriage problem, one can search for the stable marriage starting from the top choices for all men (the \perp element) of the lattice (in parallel) with the bottom choices for all men (the \top) element of the lattice. This algorithm will traverse the distance in a lattice given by the minimum of the distance of a stable marriage from the top or the bottom. Hence, depending upon the distance of man-optimal and man-pessimal stable marriages, it will return the stable marriage that is closer to the bottom or the top of the lattice.

In the above analysis, we ignore the 2 penalty factor that we incur because we run the algorithm from both the bottom and top of the lattice.

2.10 Summary

This chapter presented the Lattice-Linear Predicate (LLP) algorithm, a general method for finding the least element in a distributive lattice satisfying a given predicate. The LLP algorithm is nondeterministic and online: it does not inspect any element of the lattice above the current state. We showed that lattice-linear predicates are closed under conjunction, which means that constrained versions of problems are solved by the same algorithm with no additional machinery.

We also discussed the dual of lattice-linearity, which enables traversal from the top of the lattice downward. For predicates that are both lattice-linear and dual lattice-linear (called *regular* predicates), one can search from both ends simultaneously.

Table [2.2](#) lists the algorithms presented in this chapter.

Here n is the number of components, m is the number of precedence edges (for job scheduling), and h is the height of the lattice. The LLP and Regular algorithms give the number of iterations of the outer

Algorithm	Problem/Topic	Time Complexity
LLP	General LLP (least fixpoint)	$O(h \cdot n)$ iterations
LLP-JobScheduling-Fixed	Job Scheduling with fixed	$O(n + m)$
LLP-SEQ	General LLP (sequential)	problem-dependent
Regular	Regular predicate (bidirectional)	$O(h \cdot n)$ iterations

Table 2.2: Algorithms for lattice-linear predicate detection

loop; the cost per iteration depends on the specific predicate and advance operation.

2.11 Problems

- Let G be an n -dimensional vector of positive numbers. Let $pred$ be a binary acyclic relation on $[n]$. Show that the predicate $B \equiv \forall(i, j) \in pred : G[j] \geq G[i] + 1$ is a lattice-linear predicate.
- Let (X, \leq) be a poset. A subset $Y \subseteq X$ is an order ideal if it satisfies

$$\forall u, v \in X : (v \in Y) \wedge (u \leq v) \Rightarrow (u \in Y)$$

Show that the predicate $B(Y) \equiv "Y \text{ is an order ideal of } (X, \leq)"$ is lattice-linear in the boolean lattice of all subsets of X .

- Show that lattice-linearity is not closed under disjunction.
- Show that lattice-linearity is not closed under negation.
- Let L be any finite distributive lattice of global states. Let B be any Boolean predicate defined on L such that B is lattice-linear as well as dual lattice-linear. Let $M = \{M_1, M_2, \dots, M_k\}$ be any subset of L of size k such that all elements of M satisfy B . Then, for each index m , we get a multiset $\{M_i[m] \mid 1 \leq i \leq k\}$. For any j , we construct the generalized j -median state, G^j as follows. For any m , $G^j[m]$ is given by the j^{th} element in the *sorted* multiset $\{M_i[m] \mid 1 \leq i \leq k\}$. Then, G^j also satisfies B .
- Consider the job scheduling instance with $n = 5$ jobs, processing times $t = [2, 3, 1, 4, 2]$, and precedence constraints $1 \rightarrow 3, 1 \rightarrow 4, 2 \rightarrow 4, 3 \rightarrow 5, 4 \rightarrow 5$. Trace the execution of Algorithm LLP-JobScheduling-Fixed. At each step, state which indices are forbidden, show the advance, and give the updated vector G . What is the final completion-time vector and the critical path?
- Determine whether each of the following predicates on vectors $G \in \mathbb{Z}_{\geq 0}^n$ is lattice-linear on the componentwise lattice. Prove your answer.
 - $B_1(G) \equiv \forall i < n : G[i] \leq G[i + 1]$ (the vector is non-decreasing).
 - $B_2(G) \equiv G[1] + G[2] + \dots + G[n] \geq k$ for a fixed constant k .
 - $B_3(G) \equiv \max_i G[i] - \min_i G[i] \leq d$ for a fixed constant d (bounded range).
- Given an array $A[1..n]$ of positive integers, the *least common multiple* (LCM) of A is the smallest positive integer divisible by every $A[i]$. Formulate the LCM computation as an LLP problem: define the lattice, the state vector, the predicate B , the forbidden condition, and the advance operation. Prove that B is lattice-linear.

9. Consider the job scheduling problem with the additional constraint that each job i has a *release time* $r[i]$: job i cannot start before time $r[i]$, so its completion time must satisfy $G[i] \geq r[i] + t[i]$.
- (a) Show that the release-time predicate $B_r(G) \equiv \forall i : G[i] \geq r[i] + t[i]$ is lattice-linear.
 - (b) Using Lemma 2.3(c), give the combined forbidden and advance operations for the constrained problem (precedence constraints \wedge release-time constraints).
10. Prove that the predicate $B(G) \equiv “G \text{ is a permutation of } (1, 2, \dots, n)”$ is *not* lattice-linear on the componentwise lattice of vectors in $\{1, \dots, n\}^n$.

2.12 Bibliographic Remarks

The Lattice-Linear Predicate (LLP) algorithm is a general technique for designing algorithms for combinatorial optimization problems. The lattice-linearity property was introduced by Chase and Garg [CG98] in the context of detecting global predicates in distributed computations. In that setting, a distributed system of n processes generates a lattice of global states, and the goal is to find the least global state satisfying a given predicate. The key observation — that if the predicate is lattice-linear, a single pass through the lattice suffices — was later applied to algorithm design.

The application of LLP to the stable marriage problem is shown in Garg [Gar17], and its systematic application to algorithm design — including the shortest path problem, the assignment problem, and many others — is developed in Garg [Gar20a]. The LLP framework has since been extended to dynamic programming problems [GS24], to the housing allocation problem [Gar22], and to the minimum spanning tree problem [GG19]. Gupta and Kulkarni extend LLP algorithms for deriving self-stabilizing algorithms [GK23]. Garg [Gar23] shows that many generalizations of the stable matching problem, including the constrained stable marriage problem with ties, can also be solved using the LLP framework.

When the feasibility predicate has the form $\forall i : G[i] \geq f_i(G)$ for monotone functions f_i , the LLP algorithm is closely related to computing the least fixed point of a monotone function on a lattice via the Knaster–Tarski fixed-point theorem [Tar55, LNS82]. However, the LLP framework is more general in three respects: (1) the predicate B need not have the form $G \geq f(G)$ — we only require B to be closed under meets; (2) the Knaster–Tarski theorem requires the function to map the lattice to itself, guaranteeing a fixed point exists, whereas in LLP the algorithm may return null (e.g., cyclic job scheduling with positive weights has no solution); (3) the LLP framework supports advancement on multiple components, which the classical fixed-point iteration does not address.

The technique of *chaotic iteration* (also called *asynchronous iteration*) in abstract interpretation and dataflow analysis [CC77] also computes least fixed points by iterating on individual components in arbitrary order. The LLP algorithm can be viewed as a generalization of chaotic iteration to predicates that are not necessarily of the fixed-point form.

The classical algorithms unified by the LLP framework include Dijkstra’s algorithm [Dij59] for shortest paths, the Gale–Shapley algorithm [GS62] for stable marriage, and Kruskal’s and Prim’s algorithms for minimum spanning trees. For lattice theory background, see Appendix 21 and the texts by Birkhoff [Bir67] and Davey and Priestley [DP90].

Chapter 3

The Stable Marriage Problem

3.1 Introduction

Matching markets are everywhere. The U.S. National Resident Matching Program (NRMP) places more than 40,000 medical-school graduates into hospital residency positions each year, using an algorithm that traces directly to the work of Gale and Shapley. Big-city public-school assignment systems — New York, Boston, Singapore — match hundreds of thousands of children to schools using essentially the same procedure. Kidney-exchange clearinghouses pair incompatible donor–recipient pairs into chains so that more transplants can take place. In each setting the underlying question is the same: given a set of agents on each side and a preference order from every agent, find a pairing in which no two unpaired agents would both prefer each other.

The Stable Matching Problem formalises this question. Gale and Shapley introduced it in their 1962 paper [GS62] and showed two surprising results: a stable matching always exists, and the deferred-acceptance algorithm they introduced constructs one in $O(n^2)$ time. The economic significance of this line of work was recognised by the 2012 Nobel Prize in Economics awarded to Lloyd Shapley and Alvin Roth for the theory of stable allocations and the practice of market design.

Beyond its applications, stable matching is also the first concrete instance of the lattice-linear-predicate (LLP) framework introduced in Chapter 2. The set of *proposal vectors* forms a finite distributive lattice; the stability requirement is a lattice-linear predicate; and the Gale–Shapley algorithm is exactly the LLP main loop applied to that predicate. The chapter therefore reads as the first concrete LLP application and provides a template that later chapters — shortest paths, minimum spanning trees, network flow, and several others — will reuse.

In the standard problem there are n men numbered $1, 2, \dots, n$ and n women numbered $1, 2, \dots, n$, each with a totally ordered preference list. The matrices $mpref$ and $wpref$ encode the lists: $mpref[i][k] = j$ iff woman j is man i 's k^{th} preference, and analogously for $wpref$. A matching has a *blocking pair* (m, w) if m and w are not married to each other yet each prefers the other to their current partner; the matching is *stable* if it has no blocking pair. Fig. 3.1 shows a small instance.

$mpref$		$wpref$	
m_1	$w_2 \quad w_3 \quad w_1$	w_1	$m_1 \quad m_3 \quad m_2$
m_2	$w_2 \quad w_3 \quad w_1$	w_2	$m_2 \quad m_1 \quad m_3$
m_3	$w_3 \quad w_1 \quad w_2$	w_3	$m_1 \quad m_2 \quad m_3$

Figure 3.1: Stable Matching Problem with men preference list ($mpref$) and women preference list ($wpref$).

This chapter is organized as follows. Section 3.2 introduces the proposal-vector lattice on which all algorithms in this chapter operate. Section 3.3 presents the Gale-Shapley algorithm with its $O(n^2)$ analysis. Section 3.4 generalises Gale-Shapley to Algorithm α , which takes any starting proposal vector to the nearest stable proposal vector that dominates it; we extend this to allow $n_m \neq n_w$ men and women, assuming $n_w \geq n_m$. Section 3.5 gives the LLP formulation of stable matching together with the constrained version, in which additional lattice-linear constraints — regret bounds (e.g., Peter’s regret must be less than Paul’s), *forced pairs*, and *forbidden pairs* — can be imposed without changing the algorithm.

3.2 Proposal Vector Lattice

We use the notion of a *proposal vector* for our algorithms. A vector of (man) proposals, G , is of dimension n , the number of men. We view any vector G as follows: ($G[i] = k$) if man i has proposed to his k^{th} preference, i.e., the woman given by $mpref[i][k]$. If $mpref[i][k]$ equals j , then $G[i]$ equals k corresponds to man i proposing to woman j . The woman that man i proposes to in G is $mpref[i][G[i]]$. The vector $(1, 1, \dots, 1)$ corresponds to the proposal vector in which every man has proposed his top choice. Similarly, (n, n, \dots, n) corresponds to the vector in which every man has proposed to his last choice. Our algorithms can also handle the case when the lists are incomplete, that is, a man prefers to stay alone to be matched to some women. However, for simplicity, we assume complete lists. It is clear that the set of all proposal vectors forms a distributive lattice under the natural less than order in which the meet and join are given by the component-wise minimum and the component-wise maximum, respectively. This lattice has n^n elements.

Given any proposal vector, G , there is a unique matching defined as follows: man i and $mpref[i][G[i]]$ are matched in G if the proposal of man i is the best proposal for that woman in G or any proposal before G . The man i is unmatched otherwise. A woman q is unmatched in G if she does not receive any proposal in G or earlier.

A proposal vector G represents a *man-saturating matching* iff no woman receives more than one proposal in G . Formally, G is a man-saturating matching if $\forall i, j : i \neq j : mpref[i][G[i]] \neq mpref[j][G[j]]$. When the number of men equals the number of women, a man-saturating matching is a perfect matching (all men and women are matched). When the number of men is less than the number of women, then G is a man-saturating matching if every man is matched (but some women are unmatched). We say that a matching M_1 (or a marriage) is less than another matching M_2 if the proposal vector for M_1 is less than that of M_2 .

A proposal vector G may have one or more blocking pairs. A pair of man and woman (p, q) is a *blocking pair* in G iff $mpref[p][G[p]]$ is not q , man p prefers q to $mpref[p][G[p]]$, and woman q prefers p to any proposal received in G . Observe that this definition works even when a woman q is unmatched, i.e. she has not received any proposals in G . In this case, woman q prefers p to staying alone, and p prefers q to $mpref[p][G[p]]$.

A proposal vector G is a stable marriage (or a stable proposal vector) iff it is a man-saturating matching and there are no blocking pairs in G . The man-optimal marriage is the *least* stable matching in the proposal

lattice, and the woman-optimal marriage is the *greatest* stable matching.

3.3 Gale-Shapley Algorithm

In this section, we first present an algorithm due to Gale and Shapley for this problem (also known as the deferred acceptance algorithm). In this algorithm, a free man proposes to women in his decreasing order of preferences, i.e., he first proposes to his top choice. Only when a man is rejected by his top choice would he move to his next top choice. We maintain the list of all men who are not engaged in the variable *mList*. Initially, all men are free (not engaged) and are on this list. For this section, recall that we assume that the number of men is equal to the number of women.

When any woman *z* receives a proposal from a man *i*, she always accepts it if she is not engaged. In this case, they both get engaged, and the variable *partner*[*z*] is set to *i*. If the woman *z* is engaged, then she compares this proposal with her existing partner. If she prefers this proposal to her existing partner, then she breaks the engagement with her existing partner and makes *i* her partner. The previous partner joins *mList*, the list of free men. If the woman *z* prefers her existing partner, then the proposal of the man *i* is rejected, and the man *i* goes back to *mList*. Algorithm [Gale-Shapley](#) shows the pseudocode for these steps.

Algorithm Gale-Shapley: Finding the man-optimal marriage

Input: *mpref*, *rank*
Output: man-optimal stable marriage

```

1 mList: list of 1..n // list of men that are free, initially includes all men
2 partner: array[1..n] of 0..n initially partner[i] = 0 for all i // Current fiance for woman i
3 G: array[1..n] of 0..n initially G[i] = 0 for all i // Number of proposals made by man i
4 mList := [1..n] // Initialize all men as part of this list
5 while mList is nonempty do
6   i := mList.removeFirst();
7   G[i] := G[i] + 1 // Move to the next available top choice for man i
8   z := mpref[i][G[i]] // Woman corresponding to that choice
9   if partner[z] = 0 then partner[z] := i;
10  else if rank[z][i] < rank[z][partner[z]] then
11    | mList.add(partner[z]);
12    | partner[z] := i;
13  end
14  else mList.add(i) ;
15 end
16 return G

```

It is easy to verify the following properties of the algorithm.

Lemma 3.1 *All the following statements are valid.*

1. *As the algorithm progresses, the partner for a man can only worsen and the partner for a woman can only improve.*
2. *Once a woman is engaged, she stays engaged.*

Observe that once all men are engaged, the list $mList$ is empty and the *while* loop in the algorithm terminates. Is it possible for all men to run out of all the choices and still not have a stable marriage? We claim that whenever a man i proposes to his last choice, the proposal is always accepted. The proposal is always accepted if the woman is not engaged. If the woman is engaged, we can deduce that all women are engaged because the man had previously proposed to all other women who are now engaged. However, this is a contradiction because all n women are engaged to different men and the man i is not engaged. Hence, we conclude that the last proposal of any man is always accepted.

How many times can the *while* loop iterate? Observe that in every iteration $G[i]$ increases by 1 for some i . Since $G[i]$ starts with 0 and cannot increase beyond n , the total number of iterations for the *while* loop is $O(n^2)$. Each iteration of the *while* loop takes $O(1)$ time. Hence, the time complexity of the Gale-Shapley Algorithm is $O(n^2)$.

We now show that the GS algorithm returns the man-optimal stable marriage. Suppose that a man m is rejected by his best valid partner woman w in the GS algorithm. Let this be the first time a valid pair is rejected. Suppose w rejected m for some man m' . Thus, we know that w prefers m' to m .

Let M be a marriage in which m is married to w because (m, w) is a valid pair. In the marriage, M , let m' be married to w' . When w rejected m for m' in the GS marriage, it was the first rejection of any valid pair. This means that m' must have proposed to w before proposing to w' . Thus, we also get that m' prefers w to w' . However, this results in (m', w) forming a blocking pair for M . The man m' prefers w to w' and the woman w prefers m' to m , a contradiction. Hence, we conclude that the GS algorithm returns the man-optimal stable marriage.

Example 3.2 We trace the Gale-Shapley algorithm on the instance in Fig. 3.1. Initially, $G = (0, 0, 0)$, $partner = (0, 0, 0)$, and $mList = [m_1, m_2, m_3]$.

1. m_1 proposes to $mpref[1][1] = w_2$. $G = (1, 0, 0)$. w_2 is free; $partner[w_2] := m_1$.
2. m_2 proposes to $mpref[2][1] = w_2$. $G = (1, 1, 0)$. w_2 prefers m_2 to m_1 ($rank[w_2][m_2] = 1 < rank[w_2][m_1] = 2$). $partner[w_2] := m_2$; m_1 returns to $mList$.
3. m_3 proposes to $mpref[3][1] = w_3$. $G = (1, 1, 1)$. w_3 is free; $partner[w_3] := m_3$.
4. m_1 proposes to $mpref[1][2] = w_3$. $G = (2, 1, 1)$. w_3 prefers m_1 to m_3 ($rank[w_3][m_1] = 1 < rank[w_3][m_3] = 3$). $partner[w_3] := m_1$; m_3 returns to $mList$.
5. m_3 proposes to $mpref[3][2] = w_1$. $G = (2, 1, 2)$. w_1 is free; $partner[w_1] := m_3$.

$mList$ is empty. The algorithm terminates with $G = (2, 1, 2)$: $m_1 \leftrightarrow w_3$, $m_2 \leftrightarrow w_2$, $m_3 \leftrightarrow w_1$. This is the man-optimal stable marriage.

3.4 Algorithm α : Upward Traversal

The algorithm α generalizes the Gale-Shapley algorithm in two fundamental ways.

- *Arbitrary Initial Proposal Vector*: Gale-Shapley algorithm always finds the man-optimal stable marriage. Suppose that we are interested in finding a stable marriage such that the proposal vector is at least I . For example, if $I = (3, 1, 2, 1)$, then the first man cannot propose to his two top choices, and the third man cannot propose to his top choice. The algorithm α works even when the initial proposal vector is arbitrary instead of the top choice for each man. Observe that the standard Gale-Shapley algorithm does not work as is when the starting proposal vector is arbitrary. The simple Gale-Shapley algorithm would require men to make proposals and women to accept the best proposals they have received so far. If the starting proposal vector is a perfect matching but not stable, then each woman gets a unique proposal. All women would accept the only proposal received, but the resulting marriage may not be stable.
- *Unequal number of men and women*: We consider the case where the number of women exceeds the number of men. If the number of men is greater than the number of women, then we can simply switch the roles in our algorithm.

If we started with the top choices of all men, then the Gale-Shapley algorithm would still return a man-optimal stable matching with the excess women unmatched. However, if we start from an arbitrary proposal vector, we can end up with all women getting unique proposals, but there may exist an unmatched woman who is preferred by some man over his current match. To address this problem, we first do a simple check on the initial proposal vector given by the following lemma. Let $numw(I)$ denote the total number of unique women that have been proposed in all vectors that are less than or equal to I .

Lemma 3.3 *Given any stable marriage instance with n_m men and the initial proposal vector I there is no stable marriage for any proposal vector $G \geq I$ whenever $numw(I)$ exceeds n_m .*

Proof: Consider any proposal vector $G \geq I$. Since the total number of men is n_m , there is at least one woman q who has been proposed in the past of G who does not have any proposal in G . Suppose that the proposal was made by man p . Then, man p prefers q to $mpref[p][G[p]]$ and q prefers p to staying alone. ■

Hence, in our algorithm, we only consider I such that the total number of women proposed up to I is at most n_m . Even when the number of men equals the number of women, the proposal vector may be a perfect matching but not stable. To address this problem, we first define $forbidden(G, i)$ as the predicate that there exists another man j such that (1) both i and j have proposed to the same woman in G and that woman prefers j , or (2) $(j, mpref[i][G[i]])$ is a blocking pair in G . We first show that

Lemma 3.4 *Let G be any proposal vector such that $numw(G) \leq n_m$. There exists a man i such that $forbidden(G, i)$ iff G is not a stable marriage.*

Proof: First, suppose that there exists i such that $forbidden(G, i)$. This means that there exists a man j such that j has proposed to the same woman and that woman prefers j or $(j, mpref[i][G[i]])$ is a blocking

pair in G . If both i and j have proposed to the same woman in G , then it is clearly not a matching. If $(j, mpref[i][G[i]])$ is a blocking pair, then G is not stable.

Conversely, assume that G is not a stable marriage. This means that either G is not a man-saturating matching or there is a blocking pair for G . If G is not a man-saturating matching, then there must be some woman who has been proposed by multiple men. Any man i who is not the most favored in the set of proposals satisfies $forbidden(G, i)$. If G is a perfect matching but not a stable marriage, then there must be a blocking pair (p, q) . If q has been proposed in G by man i , then $(p, mpref[i][G[i]])$ is a blocking pair. Hence, $forbidden(G, i)$ holds. If q has not been proposed in G then we know at least $n_m + 1$ women who belong to $numw(G)$ which violates our requirement on G . ■

Algorithm [UpwardTraversal](#) exploits the $forbidden(G, i)$ function to find a stable marriage in the proposal lattice. The basic idea is that if a man i is forbidden in the current proposal vector G , then he must go down his preference list until he finds a woman who is unmatched or prefers him to her current match. The *while* loop iterates until none of the men is forbidden in G . If the while loop terminates, then G is a stable marriage due to Lemma 3.4. The man i advances on his preference list until his proposal is the most preferred proposal to the woman among all proposals that are made to her in any proposal vector less than or equal to G . If there is no such proposal, then there does not exist any $G \geq I$ such that G is stable and the algorithm returns null. Otherwise, the man makes that proposal.

Algorithm UpwardTraversal: An algorithm that returns the man-optimal marriage greater than or equal to the given proposal vector I .

Input: A stable marriage instance, initial proposal vector I

Output: smallest stable marriage greater than or equal to I (if one exists)

```

1 forbidden( $G, i$ ) holds if there exists another man  $j$  such that either (1) both  $i$  and  $j$  have
   proposed to the same woman in  $G$  and that woman prefers  $j$ , or (2)  $(j, mpref[i][G[i]])$  is a
   blocking pair for  $G$ .;
2 if  $numw(I) > n_m$  then return null // no stable matching exists ;
3 else  $G := I$ ;
4 while there exists a man  $i$  such that  $forbidden(G, i)$  do
5   | find the next woman  $q$  in the list of man  $i$  s.t.  $i$  has the most preferred proposal to  $q$  until  $G$ ;
6   | if no such choice after  $G[i]$  or the number of women proposed including  $q$  exceeds  $n_m$  then
7     |   return null // no stable matching exists ;
8     |   else  $G[i] :=$  choice that corresponds to woman  $q$ ;
8 end
9 return  $G$ 

```

There are two main differences from the Gale-Shapley algorithm. The first difference is the simple check on the number of women who have been proposed up to G . We require $numw(G) \leq n_m$. Clearly, if the number of women is equal to the number of men, then $numw(G)$ cannot exceed n_m and this check can be dropped. Also, even if the number of women exceeds the number of men, but the initial proposal vector is $(1, 1, \dots, 1)$, the number of women proposed cannot exceed n_m . However, this check is required when the number of women exceeds n_m and the initial proposal vector is arbitrary.

The second difference is in the definition of $forbidden(G, i)$. In the standard Gale-Shapley algorithm, a man advances on his preference list only when the woman he has proposed to is either matched with someone more preferable or receives a proposal from a more preferable man. Whenever the Gale-Shapley algorithm reaches a perfect matching, it is a stable matching. For any arbitrary I (for example, a perfect matching that is not stable), it is important to take blocking pairs into consideration as part of the forbidden predicate.

The time complexity of Algorithm α is $O(n_m^2)$: each man advances at most n_m times on his preference list, and each advancement takes $O(1)$ time using appropriate data structures. When $n_m = n_w = n$, this reduces to the standard $O(n^2)$ complexity of the Gale-Shapley algorithm.

3.5 LLP Stable Matching Algorithm

We now derive the algorithm for the stable matching problem using Lattice-Linear Predicates. We assume that the number of men equals the number of women in this section. We let $G[i]$ be the choice number that man i has proposed to. Initially, $G[i]$ is 1 for all men. The woman that man i proposes to in G is $mpref[i][G[i]]$.

Definition 3.5 *An assignment G is feasible for the stable marriage problem if (1) it corresponds to a perfect matching (all men are paired with different women) and (2) it has no blocking pairs.*

We show that the predicate “ G is a stable marriage” is a lattice-linear predicate.

Lemma 3.6 *The predicate that a vector G corresponds to a stable marriage is lattice-linear.*

Proof: Let $z = mpre[f][j][G[j]]$, the woman that corresponds to choice $G[j]$ for man j . We define j to be forbidden in G if there exists a man i such that z prefers man i to man j and either man i has also been assigned z in G or he prefers z to his current choice, i.e., man i and woman z would form a blocking pair in G . Formally, $forbidden(G, j)$ is defined as $(\exists i : \exists k \leq G[i] : (z = mpre[f][i][k]) \wedge (rank[z][i] < rank[z][j]))$.

It is easy to see that G is not a stable marriage iff $\exists j : forbidden(G, j)$. If G is not a perfect matching then there must be at least one woman who is assigned to two men. In that case, the less preferred man is forbidden. If G is a perfect matching but has a blocking pair, then the partner of the woman in the blocking pair is forbidden. Conversely, $forbidden(G, j)$ implies that either G is not a perfect matching or has a blocking pair.

We only need to show that if $forbidden(G, j)$ holds, then there is no proposal vector H such that $(H \geq G)$ and $(G[j] = H[j])$ and H is a stable marriage.

Consider any H such that $(H \geq G)$ and $(G[j] = H[j])$. We show that H is not a stable marriage. Since $G[j]$ is equal to $H[j]$, $mpref[j][G[j]]$ is equal to $mpref[j][H[j]]$. Let i be such that $\exists k \leq G[i] : (z = mpre[f][i][k]) \wedge (rank[z][i] < rank[z][j])$. Since $G \leq H$, $G[i] \leq H[i]$, we get that $\exists k \leq H[i] : (z = mpre[f][i][k]) \wedge (rank[z][i] < rank[z][j])$. Hence, $forbidden(H, j)$ also holds. ■

Lemma 3.6 immediately gives us the Algorithm [LLP-StableMarriage](#). The **always** section defines variables which are derived from G . These variables can be viewed as macros. For example, for any thread $z = mpre[f][j][G[j]]$. This means that whenever $G[j]$ changes, so does z .

If man j is forbidden, it is clear that any vector in which man j is matched with z and man i is matched with his current or a worse choice can never be a stable marriage. Thus, it is safe for man j to advance to the next choice.

Algorithm LLP-StableMarriage: LLP Program for LLP-StableMarriage

Input: $mpref$: 2D array of int; $rank$: 2D array of int; I : array of int

Output: G : array of int, initially $I[i]$

- 1 **forbidden**(j): $\exists i \in [1..n] : \exists k \in [1..G[i]] : (mpref[j][G[j]] = (mpref[i][k]) \wedge (rank[mpref[j][G[j]]][i]) < (rank[mpref[j][G[j]]][j]));$
 - 2 **advance**: $G[j] := G[j] + 1;$
-

The general properties of the LLP algorithm (Section 2.5) are well illustrated by the stable marriage problem:

- *Nondeterminism*: The man-optimal stable marriage is the same regardless of the order in which forbidden men advance on their preference lists.
- *No lookahead*: A man need not reveal his full preference list in advance. Only when he is rejected (forbidden) does he reveal his next choice. This makes the algorithm applicable in online settings.
- *Individual optimality*: The LLP solution is optimal for each man individually. If man m seeks the stable marriage that gives him his best possible partner (among all stable marriages), the LLP algorithm returns exactly that partner for m .

Algorithm **LLP-StableMarriage** works for any initialization I of G vector. If we know that G is initialized to $[1, 1, \dots, 1]$, then we can simplify the forbidden condition to

$$(\exists i : (z = mpref[i][G[i]]) \wedge (rank[z][i] < rank[z][j])).$$

Observe that if we assume sequential implementation and if we implement a variable $partner[z]$ for any woman z , we can further simplify the condition to

$$(partner[z] \neq null) \wedge (rank[z][partner[z]] < rank[z][j]).$$

This is precisely the condition used in the Gale-Shapley algorithm which is sequential and starts with the initial vector $[1, 1, \dots, 1]$.

We now present an algorithm to find stable marriages that satisfy additional constraints. The following lemma proves lattice-linearity of many such constraints.

Lemma 3.7 *The following constraints are lattice-linear.*

1. *The regret of man i is at most that of man j .*
2. *Man i cannot be married to woman j .*
3. *The regret of man i is equal to that of man j .*

Proof: Let B be the predicate that G is a stable marriage and it satisfies the corresponding additional constraint.

1. Suppose that G is a stable marriage but it does not satisfy B . This means that the regret of man i is more than the regret of man j . In this case, we have $forbidden(G, j)$, because unless $G[j]$ is advanced, the predicate cannot become true.
2. If $mpref[i][G[i]] = j$, then $forbidden(G, i)$ holds.
3. This condition is a conjunction of two lattice-linear conditions of type in part (1).

■

Observe that we have not given the argument that the LLP predicate eventually becomes true. In general, this may not happen for all LLP predicates. For the standard stable marriage, however, the LLP predicate becomes true. We can define the predicate $D(k)$ on the set of assignments as “By the assignment G , at least k different women have been proposed.” It is clear that before all the proposals are done $D(n)$ holds. It is easy to show that $D(k)$ implies that there are k valid pairs in the assignment. Therefore, $D(n)$ implies that B holds on G .

We now enumerate advantages of the LLP algorithm.

1. It gives us a more general algorithm. Instead of starting from the initial vector, we can start from any vector and find a stable marriage greater than or equal to that vector, if one exists.
2. We can use the LLP algorithm for finding a stable marriage that satisfies additional constraints such as the regret of man i is at most that of man j .
3. We can automatically deduce that the intersection of two stable marriages is also a stable marriage.

Example 3.8 As an example of a constrained stable marriage, consider the instance in Fig. 3.1 with the additional constraint that the regret of m_1 must be at most the regret of m_2 (i.e., $G[1] \leq G[2]$). This constraint is lattice-linear: $forbidden(1) \equiv (G[1] > G[2])$ with $advance(1) : G[1] := G[2]$. By Lemma 2.3(c), the conjunction of the stability predicate and this constraint is also lattice-linear, so the LLP algorithm finds the least stable marriage satisfying the constraint. In the unconstrained man-optimal marriage $G = (2, 1, 2)$, m_1 has regret 2 while m_2 has regret 1, violating the constraint. The LLP algorithm with the combined predicate would advance further to find a stable marriage where $G[1] \leq G[2]$.

3.6 Extending Algorithms

The Gale–Shapley algorithm and its LLP variants presented in this chapter all assume a clean, static, centralized setting: a fixed set of n men and n women, complete and truthful preference lists, and a single process that has access to all of them. Real applications rarely come packaged so neatly. Readers are encouraged to study how the algorithms of this chapter adapt, generalize, or fail under the following conditions:

1. **Incremental setting:** Men and/or women may join or leave the system dynamically after an initial matching has been computed. Does the existing matching need to be recomputed from scratch, or can it be repaired locally while preserving stability?

2. **Dynamic setting:** The preference lists change over time (for example, a participant revises their ranking after learning more about the candidates). How should the algorithm track a moving stable matching?
3. **Distributed setting:** Each participant has access only to their own preference list and can communicate only with a subset of the other side. Design a stable matching algorithm in which no single node sees the entire instance.
4. **Privacy setting:** Each participant is willing to reveal only partial information about their preferences (e.g., their top k choices, or a coarse ranking). Can a stable matching still be computed, possibly with a degradation in the quality of the result?
5. **Constrained setting:** The matching must satisfy additional side constraints—for example, forced pairs that must appear in the matching, forbidden pairs that must not, regional or diversity caps, or couples who must be matched to nearby partners. Some such constraint families are already treated in the book; others make for good exercises.
6. **Ties and indifference:** Participants are allowed ties in their preference lists. Stable matchings may no longer be unique, and several notions of stability (weakly, strongly, super-) diverge. Which of them can be computed with LLP-style algorithms?
7. **Parallel setting:** The algorithm is parallelized to reduce the wall-clock time of computing a stable matching. How large can the parallel speedup be in the worst case, and how does it compare with the critical-path analysis of Algorithms α and β ?
8. **Approximation setting:** When no stable matching exists (e.g., in the stable roommate problem with odd-length preference cycles), the goal is to find a near-stable solution that minimizes the number or severity of blocking pairs.
9. **Fairness setting:** Gale–Shapley is known to be men-optimal and women-pessimal (or vice versa). The objective is to produce a stable matching that is fair across the two sides, for example by minimizing the sum or the maximum of regrets.
10. **Enumeration setting:** The task is to enumerate all stable matchings of a given instance, or the top- k stable matchings according to a given objective. The lattice of stable matchings makes this a natural LLP target.
11. **Online setting:** Participants arrive one at a time in an adversarial or random order, and the algorithm must make irrevocable decisions as each participant is revealed. How close to an offline stable matching can one stay?
12. **Byzantine setting:** Some participants behave strategically or maliciously—misreporting preferences, refusing to execute the protocol, or colluding to disrupt the matching for truthful participants. Which of the algorithms in this chapter are robust to such behaviour, and what defenses are possible?

3.7 Summary

This chapter studied the stable marriage problem through the lens of lattice-linear predicate detection. We modeled the problem using the proposal vector lattice, where each vector G assigns a choice number

to every man. The key insight is that the predicate “ G is a stable marriage” is lattice-linear: if man j is forbidden in G (another man is preferred by $\text{mpref}[j][G[j]]$), then j remains forbidden in every $H \geq G$ with $H[j] = G[j]$.

We presented the Gale–Shapley algorithm as a special case of the LLP framework. Algorithm α generalizes Gale–Shapley to find the man-optimal stable marriage from any initial proposal vector I , and LLP-StableMarriage expresses the same algorithm in the LLP forbidden/advance notation.

Table 3.1 lists the algorithms discussed in this chapter.

Algorithm	Problem	Time Complexity
Gale–Shapley	Stable Marriage	$O(n^2)$
α (Upward Traversal)	Stable Marriage $\geq I$	$O(n^2)$
LLP-StableMarriage	Stable Marriage	$O(n^2)$

Table 3.1: Algorithms for the stable marriage problem

3.8 Problems

1. Show that Gale–Shapley algorithm maintains the invariant that every woman is partnered with the most preferred man of all the proposals made so far to her. Use this claim to show that there is no blocking pair for all the engaged couples during the execution of Gale–Shapley algorithm.
2. Show that the man-oriented Gale–Shapley algorithm returns the stable marriage, which is the worst stable marriage from the perspective of women.
3. Show that the set of stable marriages is closed under the meet operation and the join operation of the proposal lattice.
4. Given an assignment vector, how will you check whether it forms a stable marriage?
5. Consider $n = 4$ with the following preferences. Men: $m_1: w_1, w_2, w_3, w_4$; $m_2: w_2, w_1, w_4, w_3$; $m_3: w_1, w_4, w_3, w_2$; $m_4: w_4, w_1, w_2, w_3$. Women: $w_1: m_2, m_1, m_4, m_3$; $w_2: m_4, m_3, m_1, m_2$; $w_3: m_1, m_3, m_2, m_4$; $w_4: m_3, m_4, m_2, m_1$. Trace the man-oriented Gale–Shapley algorithm. Show the proposal vector G after each round and give the final stable matching.
6. Consider $n = 3$ with men’s preferences: $m_1: w_1, w_2, w_3$; $m_2: w_2, w_1, w_3$; $m_3: w_1, w_3, w_2$. Women’s preferences: $w_1: m_2, m_1, m_3$; $w_2: m_1, m_3, m_2$; $w_3: m_3, m_2, m_1$. The matching $M = \{(m_1, w_2), (m_2, w_3), (m_3, w_1)\}$ is given. Find all blocking pairs, or prove that M is stable.
7. Using the instance from the Gale–Shapley trace problem above (the $n = 4$ instance), find the man-optimal stable matching subject to the constraint that m_1 must be matched to either w_1 or w_2 (i.e., $G[1] \leq 2$). Does a constrained stable matching exist?
8. The *stable roommate problem* is the non-bipartite version of stable matching: given $2n$ people (no men/women distinction), each with a preference ranking over all others, find a matching into n pairs with no blocking pair. Show that stable matchings need not exist in the roommate setting by constructing a 4-person instance with no stable matching. Explain why the bipartite structure of the stable marriage problem guarantees existence.

9. Let S_1 and S_2 be two distinct stable matchings for the same instance. A man m is said to *prefer* S_1 if he prefers his partner in S_1 to his partner in S_2 , and similarly for women. Prove: if man m prefers S_1 , then his partner in S_1 prefers S_2 .
10. Construct an instance with n men and n women in which the man-oriented Gale–Shapley algorithm makes $\Omega(n^2)$ proposals.
11. Give an $O(n^2)$ implementation of Algorithm α (Upward Traversal). Your implementation should use:
 - An array $curBest[1..n]$ where $curBest[q]$ is the most preferred man among all men who have proposed to woman q in any proposal vector less than or equal to G .
 - A list $mList$ of men i such that $forbidden(G, i)$ holds.

Show that $forbidden(G, i)$ holds iff $curBest[mpref[i][G[i]]] \neq i$. Use this to argue that the while loop terminates with a stable marriage or correctly reports that none exists.

3.9 Bibliographic Remarks

The stable matching problem was introduced by Gale and Shapley [GS62] and has been studied extensively, with several books and survey articles devoted to the topic [GI89, Knu97c, RS92, IM08, Man13]. The notion of *regret* in a matching is discussed in [GI89]. Stable marriage with restricted pairs (forced and forbidden pairs) is studied by Dias et al. [DdFDFS03]. The lattice theory background on order ideals and distributive lattices used in this chapter can be found in Davey and Priestley [DP90]. The existence of a stable matching via Tarski’s fixed-point theorem is shown by Adachi [Ada00].

Chapter 4

Sorting Algorithms

4.1 Introduction

In this chapter, we consider some basic ideas in the design of algorithms that are crucially based on the notion of *order* in a set of elements of size n . We will assume that the order is total, that is, for every two elements x and y , either x is less than or equal to y , or y is less than or equal to x in that order.

Sorting is among the most fundamental problems in computer science and one of the most studied. A large fraction of the running time of real systems is spent arranging data into some order, and many algorithms that appear unrelated to sorting reduce to it as a subroutine. Once the data is sorted, many tasks become dramatically cheaper: searching for a value takes $O(\log n)$ time by binary search instead of $O(n)$; computing the median, quantiles, or any order statistic becomes $O(1)$ after an $O(n \log n)$ preprocessing; and detecting duplicates or taking the union, intersection, or difference of two sets takes linear time. Sorting also enables efficient algorithms for geometric problems such as finding the closest pair of points and constructing convex hulls, and it is a building block for data compression, indexing, and scheduling.

The ubiquity of sorting makes even small constant-factor improvements worthwhile.

Before we discuss sorting algorithms, let us consider the problem of searching for an element in a sorted array. Suppose that we are given a sorted array A of size n and an element x . We are interested in finding out if there exists i such that $A[i]$ equals x . On a single processor, we can accomplish this using binary search in $O(\log n)$ time. The idea is to compare x with the middle element of the array. If the middle element is equal to x , we are done; otherwise, the range of indices of A that can possibly have x is divided by a factor of 2.

This chapter is organized as follows. Section 4.2 discusses sorting algorithms based on the swapping of consecutive entries that are out of order. This is a general class of algorithms that are very natural. However, these algorithms take $O(n^2)$ sequential time in the worst case. Section 4.4 discusses *MergeSort*. Section 4.3 discusses *QuickSort*. Since MergeSort and QuickSort are based on the divide-and-conquer paradigm, they are revisited in Chapter 9. All these sorting algorithms are based on the comparison of two entries in the array. Finally, Section 4.6 describes a sorting algorithm that is not based on the comparison of two entries.

4.2 Algorithms Based on Swapping Consecutive Entries

Let A be an array of distinct integers. Our goal is to sort the array.

Algorithm BubbleSort: Bubble Sort

Input: $A[1..n]$: array of int
Output: $A[1..n]$: sorted array

```

1 repeat
2    $found := false$ ;
3   for  $j = 1$  to  $n - 1$  do
4     if  $A[j] > A[j + 1]$  then
5        $found := true$ ;
6        $swap(A[j], A[j + 1])$ ;
7     end
8   end
9 until  $\neg found$ ;
```

Algorithm `BubbleSort` swaps entries that are out of order in one iteration of the *for* loop.

Example 4.1 Consider sorting the array $A = [5, 3, 8, 1]$ using Bubble Sort.

Pass 1 ($j = 1, 2, 3$): Compare (5, 3), swap $\rightarrow [3, 5, 8, 1]$. Compare (5, 8), no swap. Compare (8, 1), swap $\rightarrow [3, 5, 1, 8]$. The largest element 8 has “bubbled” to the end.

Pass 2: Compare (3, 5), no swap. Compare (5, 1), swap $\rightarrow [3, 1, 5, 8]$. Compare (5, 8), no swap.

Pass 3: Compare (3, 1), swap $\rightarrow [1, 3, 5, 8]$. No more swaps needed; *found* remains false in the next pass, and the algorithm terminates.

Each pass moves at least one element to its correct position. Hence, the repeat loop is executed at most $n - 1$ times, giving a sequential time complexity of $O(n^2)$.

Another schedule is to increase the size of the sorted array one at a time.

Algorithm InsertionSort: Insertion Sort

Input: $A[1..n]$: array of int
Output: $A[1..n]$: sorted array

```

1 for  $i = 2$  to  $n$  do
2   for  $j = i - 1$  downto 1 do
3     if  $A[j] \leq A[j + 1]$  then break;
4      $swap(A[j], A[j + 1])$ ;
5   end
6 end
```

The time complexity is clearly $O(n^2)$ due to nested *for* loops.

Example 4.2 Consider sorting $A = [5, 3, 8, 1]$ using Insertion Sort.

$i = 2$: Insert $A[2] = 3$ into the sorted subarray $[5]$. Compare $(5, 3)$, swap $\rightarrow [3, 5, 8, 1]$.

$i = 3$: Insert $A[3] = 8$ into $[3, 5]$. Compare $(5, 8)$, no swap. Compare $(3, 5)$, no swap. Array: $[3, 5, 8, 1]$.

$i = 4$: Insert $A[4] = 1$ into $[3, 5, 8]$. Compare $(8, 1)$, swap $\rightarrow [3, 5, 1, 8]$. Compare $(5, 1)$, swap $\rightarrow [3, 1, 5, 8]$. Compare $(3, 1)$, swap $\rightarrow [1, 3, 5, 8]$. Sorted.

We now prove that any comparison-based sorting algorithm that restricts itself to comparing consecutive entries (adjacent elements) in an array of size n requires at least $\Omega(n^2)$ comparisons in the worst case. This applies to algorithms such as Bubble Sort or Insertion Sort, where comparisons are limited to pairs of elements at positions i and $i + 1$.

Theorem 4.3 Consider an array $A = [a_1, a_2, \dots, a_n]$ of n distinct elements. A sorting algorithm that uses only consecutive comparisons requires at least $\Omega(n^2)$ comparisons.

Proof:

The maximum number of inversions in a permutation of n elements is achieved by the reverse order $[n, n - 1, \dots, 2, 1]$, with:

$$\text{Number of inversions} = \binom{n}{2} = \frac{n(n-1)}{2}.$$

Each adjacent comparison can resolve at most one inversion (e.g., swapping a_i and a_{i+1}), so a permutation with $\frac{n(n-1)}{2}$ inversions requires at least that many comparisons to sort fully. ■

4.3 Quicksort

Quicksort (due to C.A.R. Hoare) is one of the fastest sorting algorithms on sequential computers. It has $O(n^2)$ worst case time complexity but requires $O(n \log n)$ time on average. The algorithm is based on partitioning the array into two parts using a *pivot*. Once we have the property that the lower side of the array is less than or equal to the pivot, and the upper side of the array has elements greater than the pivot, then we can recurse on each side. Since sorting each of the sides is independent, they can be sorted in parallel.

There are multiple methods to choose a pivot to partition the array A . Choosing an element at random to be the pivot is an easy method that will result in approximately equal sized partitions on average. This gives us the average case sequential time complexity of $O(n \log n)$. In the worst case, the recursion may reduce the range by only 1, resulting in the sequential time complexity of $O(n^2)$. For example, if the array is already sorted and the pivot is always the first element, then each partition produces one empty part and one part of size $n - 1$, giving the recurrence $T(n) = T(n - 1) + O(n)$, which yields $T(n) = O(n^2)$.

We describe a slight variant of the Quicksort algorithm in which we partition the array into three parts. Algorithm [ThreeWayPartition](#) takes array A , *low*, and *high* as input parameters. The part of the array this method partitions is given by

$$\{A[i] \mid \text{low} \leq i < \text{high}\}$$

Algorithm QuickSort: Sequential QuickSort

```

1 QuickSort( $A, low, high$ );
2 if  $low < high$  then
3   |  $pivot := choosePivot()$ ;
4   |  $(p, q) := Partition(A, pivot, low, high)$ ;
5   | QuickSort( $A, low, p$ );
6   | QuickSort( $A, q, high$ );
7 end

```

Note that when low equals $high$, the range is empty.

The method *partition* returns two indices p and q . All elements in the range $[low \dots p)$ are strictly less than the pivot, in the range $[p \dots q)$ are equal to the pivot, and in the range $[q \dots high)$ are greater than the pivot. We only need to recurse on the first and the third parts. Depending upon the pivot, any (or both!) of the first and third parts may be empty.

Once we have a pivot, how do we partition the array into three parts: the first partition with all elements less than the pivot, the second one with all elements equal to the pivot, and the third partition with elements strictly greater than the pivot?

Algorithm [ThreeWayPartition](#) is due to Dijkstra, who called this problem the Dutch National Flag problem. The *while* loop maintains the invariant that the entries $[low \dots p)$ are less than the pivot, $[p \dots q)$ are equal to the pivot and $[k \dots high)$ are greater than the pivot. The algorithm initializes p and q to low , therefore the first two ranges are empty initially and trivially satisfy the invariants. It also initializes k to $high$, making the range $[k \dots high)$ empty and thereby ensuring that the invariant holds initially. The range $[q..k)$ corresponds to the initial input and initially contains entries that may be less than, equal to, or greater than the pivot. In each iteration of the *while* loop, this range is reduced by one by increasing q or decreasing k . Depending on the comparison between $A[q]$ and the pivot, the entry $A[q]$ is placed in the appropriate range keeping the invariants.

It is easy to verify that the algorithm takes $O(n)$ time when the range has n elements.

Example 4.4 Consider partitioning $A = [3, 7, 2, 5, 1, 8, 4]$ with $pivot = 4$, $low = 1$, $high = 8$. Initially $p = q = 1$, $k = 8$.

p	q	k	Array	Action
1	1	8	[3, 7, 2, 5, 1, 8, 4]	$A[1]=3 < 4$: swap $A[1], A[1]$; $p=2, q=2$
2	2	8	[3, 7, 2, 5, 1, 8, 4]	$A[2]=7 > 4$: $k=7$; swap $A[2], A[7]$
2	2	7	[3, 4, 2, 5, 1, 8, 7]	$A[2]=4 = 4$: $q=3$
2	3	7	[3, 4, 2, 5, 1, 8, 7]	$A[3]=2 < 4$: swap $A[2], A[3]$; $p=3, q=4$
3	4	7	[3, 2, 4, 5, 1, 8, 7]	$A[4]=5 > 4$: $k=6$; swap $A[4], A[6]$
3	4	6	[3, 2, 4, 8, 1, 5, 7]	$A[4]=8 > 4$: $k=5$; swap $A[4], A[5]$
3	4	5	[3, 2, 4, 1, 8, 5, 7]	$A[4]=1 < 4$: swap $A[3], A[4]$; $p=4, q=5$

Now $q = k = 5$, so the loop ends. Return $(p, q) = (4, 5)$. The array is $[3, 2, 1, 4, 8, 5, 7]$: positions $[1..3]$ are less than 4, position $[4]$ equals 4, and positions $[5..7]$ are greater than 4.

Algorithm ThreeWayPartition: Three-Way Partition (Dutch National Flag)

```

1 Partition( $A, pivot, low, high$ ) returns (int, int);
2  $p, q := low$ ;
3  $k := high$ ;
4 while  $q < k$  do
5   if  $A[q] < pivot$  then
6     |  $swap(A[p], A[q])$ ;
7     |  $p := p + 1$ ;
8     |  $q := q + 1$ ;
9   end
10  else if  $A[q] > pivot$  then
11    |  $k := k - 1$ ;
12    |  $swap(A[q], A[k])$ ;
13  end
14  else  $q := q + 1$ ;
15 end
16 return ( $p, q$ )

```

4.4 Merge Sort

Merge Sort is the classic example of the divide-and-conquer method in algorithm design. To sort an array A , we divide it into two halves, recursively sort each half, and then merge the two sorted halves. The base case is when a half has only a single element and is already sorted. The key step is the merge: given two sorted arrays B and C , we would like to merge them into another array D so that D is sorted.

Example 4.5 Sort $A = [5, 3, 8, 1]$ using Merge Sort.

Split: $[5, 3]$ and $[8, 1]$

Recursively sort left: split $[5]$ and $[3]$, merge $\rightarrow [3, 5]$

Recursively sort right: split $[8]$ and $[1]$, merge $\rightarrow [1, 8]$

Merge $[3, 5]$ and $[1, 8]$: compare 3 and 1 $\rightarrow 1$; compare 3 and 8 $\rightarrow 3$;
compare 5 and 8 $\rightarrow 5$; copy 8. Result: $[1, 3, 5, 8]$.

So, it is sufficient to consider the following problem: we have two sorted arrays B and C , each of size n . We would like to merge them into another array D so that D is sorted.

It is easy to design a sequential algorithm that merges two arrays B and C into D . We simply keep two indices i and j in the arrays B and C , respectively. In any step of the algorithm, we compare $B[i]$ and $C[j]$. If $B[i]$ is smaller than $C[j]$, then we copy $B[i]$ into the next available slot in D and advance the index i . If $C[j]$ is smaller than $B[i]$, then we copy $C[j]$ into the next available slot in D and advance the index j . This algorithm takes $O(n)$ time.

Sequentially, *MergeTwo* takes $O(m + n)$ time.

Algorithm MergeSort: Sequential Merge Sort

```

1 MergeSort( $A, low, high$ );
2 if  $low < high$  then
3   |  $mid := \lfloor (low + high)/2 \rfloor$ ;
4   |  $B := \text{MergeSort}(A, low, mid)$ ;
5   |  $C := \text{MergeSort}(A, mid + 1, high)$ ;
6   | return MergeTwo( $B, C$ );
7 end
8 else return  $A[low]$  ;

```

Algorithm MergeTwo: Merging Two Sorted Arrays

Input: $B[1..m], C[1..n]$: sorted arrays of int

Output: $D[1..m+n]$: merged sorted array

```

1  $i, j, k := 1, 1, 1$ ;
2 while  $i \leq m$  and  $j \leq n$  do
3   | if  $B[i] < C[j]$  then
4     |  $D[k] := B[i]$ ;
5     |  $i := i + 1$ ;
6   | end
7   | else
8     |  $D[k] := C[j]$ ;
9     |  $j := j + 1$ ;
10  | end
11  |  $k := k + 1$ ;
12 end
13 while  $i \leq m$  do
14 |  $D[k] := B[i]; i := i + 1; k := k + 1$ 
15 end
16 while  $j \leq n$  do
17 |  $D[k] := C[j]; j := j + 1; k := k + 1$ 
18 end

```

Example 4.6 Merge $B = [2, 5, 8]$ and $C = [1, 4, 9]$.

Step 1: Compare $B[1]=2$ and $C[1]=1$. Since $1 < 2$, copy 1 to D . $D = [1]$.

Step 2: Compare $B[1]=2$ and $C[2]=4$. Since $2 < 4$, copy 2. $D = [1, 2]$.

Step 3: Compare $B[2]=5$ and $C[2]=4$. Copy 4. $D = [1, 2, 4]$.

Step 4: Compare $B[2]=5$ and $C[3]=9$. Copy 5. $D = [1, 2, 4, 5]$.

Step 5: Compare $B[3]=8$ and $C[3]=9$. Copy 8. $D = [1, 2, 4, 5, 8]$.

B is exhausted; copy remaining $C[3]=9$. $D = [1, 2, 4, 5, 8, 9]$.

For n total elements, the recurrence of the merge sort is $T(n) = 2T(n/2) + O(n)$, resulting in $O(n \log n)$ time.

4.5 Lower Bound for Comparison-Based Sorting

We show that any comparison-based sorting algorithm for an array of n distinct elements requires at least $\Omega(n \log n)$ comparisons in the worst case. This result applies to algorithms using unrestricted pairwise comparisons (e.g., Merge Sort), contrasting with the $\Omega(n^2)$ bound for consecutive-only comparisons. We use a decision tree model.

Theorem 4.7 *Given an array $A = [a_1, a_2, \dots, a_n]$ of n distinct elements, a comparison-based sorting algorithm uses binary comparisons ($a_i < a_j$) to sort A into $a_{\pi(1)} < a_{\pi(2)} < \dots < a_{\pi(n)}$, where π is a permutation. The minimum number of comparisons needed to sort any input permutation in the worst case is $\Omega(n \log n)$.*

Proof: We model the sorting algorithm as a decision tree, where internal nodes represent comparisons, and leaves correspond to permutations. The worst-case number of comparisons is the tree's height.

For n distinct elements, there are $n!$ possible permutations. The algorithm must distinguish all $n!$ permutations, so the decision tree requires at least $n!$ leaves.

In a comparison-based algorithm, each node compares two elements (a_i vs. a_j), branching on the less-than operator ($<$) or the greater-than operator ($>$). With distinct elements, each node has two children (binary tree). After k comparisons, the maximum number of leaves is 2^k .

To cover all permutations:

$$2^k \geq n!.$$

Taking the base-2 logarithm:

$$k \geq \log_2(n!).$$

To show that $\log_2(n!) = \Omega(n \log n)$, observe that

$$n! \geq \left(\frac{n}{2}\right)^{n/2},$$

since the product $n!$ contains at least $n/2$ factors each at least $n/2$. Taking logarithms:

$$\log_2(n!) \geq \frac{n}{2} \log_2\left(\frac{n}{2}\right) = \Omega(n \log n).$$

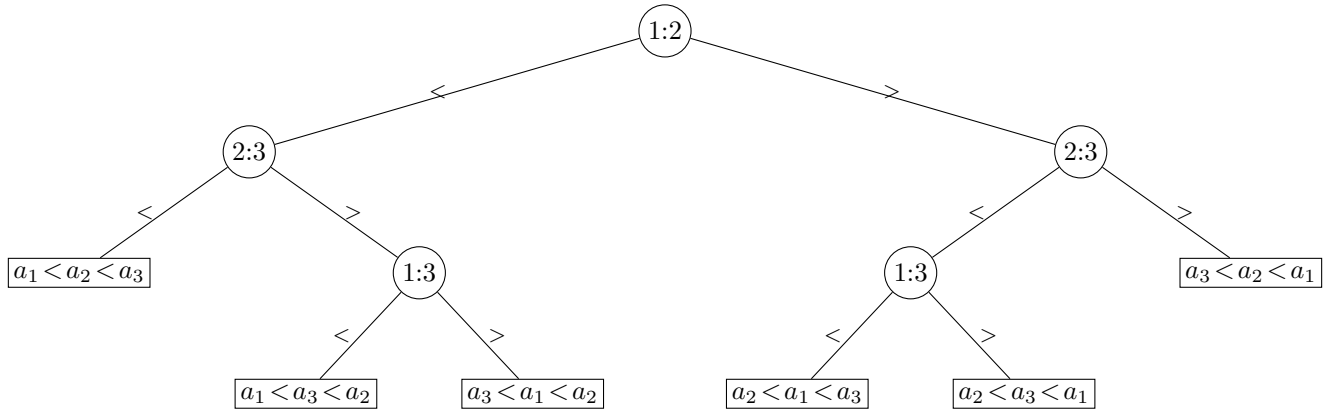


Figure 4.1: A decision tree for sorting three elements. Internal nodes are comparisons $i:j$ (is $a_i < a_j$?); left branches mean “yes,” right branches mean “no.” The six leaves correspond to the six permutations.

For $n = 3$, there are $3! = 6$ permutations, so the decision tree needs at least $\lceil \log_2 6 \rceil = 3$ levels. Fig. 4.1 shows a decision tree that sorts three elements using at most 3 comparisons. At each internal node, the label $i:j$ denotes the comparison $a_i < a_j$; the left child corresponds to “yes” and the right child to “no” (i.e., $a_i > a_j$). Each leaf is labeled with the sorted order of the indices.

Since $k \geq \log_2(n!)$ and $\log_2(n!) = \Omega(n \log n)$, we conclude:

$$k = \Omega(n \log n).$$

■

4.6 Radix Sort

All our sorting algorithms, so far, have been based on comparison. Any comparison-based sorting algorithm must do at least $\Omega(n \log n)$ work. We now show an algorithm that works when the sorting keys have a fixed number of digits (in any radix r , generally a power of 2). In our following examples, we simply use digits at the base 10, as they are easy for us humans. Suppose that we need to sort the following list. [85, 72, 94, 45, 13, 12, 61, 81]. Here, our keys have two digits. The idea of radix sorting is to sort numbers one digit at a time. When sorting on a single digit, we use a simple linear time sorting, exploiting the fact that there are only r possibilities for each digit. Upon getting any number, we can simply add it to the pile associated with that value. The number of passes that we will make on the array is equal to the number of digits.

It may appear that it is easier to sort starting from the most significant digit (msd) first. If we used this strategy, we would get [13, 12, 45, 61, 72, 85, 81, 94] after the first pass. The second pass would consist of sorting all subarrays with the same msd, and we would get the array [12, 13, 45, 61, 72, 81, 85, 94]. The problem with this approach is that we are forced to maintain different subarrays, one for each digit after the first pass. With every subsequent pass, it becomes even more cumbersome. Hence, somewhat counterintuitively, we will employ the least significant digit first strategy. After the first pass, we get [61, 81, 72, 12, 13, 94, 85, 45]. We do not need to remember any sublists that are created during the first pass. Now, we sort by the second least significant digit. We need to ensure that if two numbers have the same digit, then their relative order from the previous pass is preserved. In other words, we require our

sorting algorithm at each pass to be *stable*: a sorting algorithm is stable if elements with equal keys appear in the output in the same order as they appear in the input. After the second pass, we get the sorted array [12, 13, 45, 61, 72, 81, 85, 94]. Since 12 appeared before 13 after the first pass, the order is preserved after the second pass.

Example 4.8 Sort [85, 72, 94, 45, 13, 12, 61, 81] by LSD radix sort (base 10).

Pass 1 (sort by ones digit): distribute into buckets:

Digit	Numbers
1	61, 81
2	72, 12
3	13
4	94
5	85, 45

Concatenating: [61, 81, 72, 12, 13, 94, 85, 45].

Pass 2 (sort by tens digit, stably):

Digit	Numbers
1	12, 13
4	45
6	61
7	72
8	81, 85
9	94

Concatenating: [12, 13, 45, 61, 72, 81, 85, 94]. Sorted!

Note that stability is essential: in Pass 2, both 81 and 85 have tens digit 8, and their relative order from Pass 1 (81 before 85) is preserved, ensuring correctness.

Thus, the sequential algorithm is simply stated as Algorithm [RadixSort](#).

Algorithm RadixSort: Radix Sort for integers with k digits

```

1 RadixSort( $A[1..n]$ );
2 for  $i = 1$  to  $k$  do
3   | StableSort( $A$ , digit  $i$ )
4 end
```

The time complexity of this algorithm is $O(kn)$ assuming that the stable sort is achieved in $O(n)$ time. For $r = 10$, sorting [85, 72, 94, 45, 13, 12, 61, 81] yields $O(kn)$ time (here, $k = 2$, $O(n)$ per pass).

4.7 LLP Perspective

At face value, the sorting problem does not appear to be searching for a satisfying element in a lattice. However, it can be viewed from that angle. Sorting an array A is the same as finding a permutation π

such that applying π to A yields the sorted array. For example, if $A = [45, 12, 15]$, then the permutation $[3, 1, 2]$ sorts A once the entry at position i moves to position $\pi(i)$.

There are many ways to represent a permutation. We represent it by its *inversion vector*. The *inversion number* of entry i is the number of entries less than i that appear to the right of i in the permutation. Thus, the permutation $[3, 1, 2]$ is equivalently written as the inversion vector $[2, 0, 0]$: there are two numbers less than 3 that appear to the right of 3, zero numbers less than 1 to the right of 1, and zero to the right of 2. There is a one-to-one correspondence between permutations and inversion vectors.

Consider the set of all inversion vectors. The last entry is always zero (nothing can appear to its right). The first entry can have $0 \dots n - 1$ inversions, the second entry $0 \dots n - 2$, and so on; the total number of inversion vectors is $n!$. We order inversion vectors by componentwise comparison. Under this order, the set of inversion vectors forms a finite distributive lattice. The bottom element is the zero vector, corresponding to the identity permutation. Applying the zero inversion vector to any array leaves it unchanged. *Our goal is to find the inversion vector whose application produces the sorted array.* Figure 4.2 illustrates this lattice for arrays of size three: each node is labeled with an inversion vector $(a, b, 0)$ and the array obtained by applying the corresponding permutation to $A = [45, 12, 15]$. The sorted array $[12, 15, 45]$ sits at $(2, 0, 0)$, so sorting A is equivalent to advancing from the bottom of the lattice to that node.

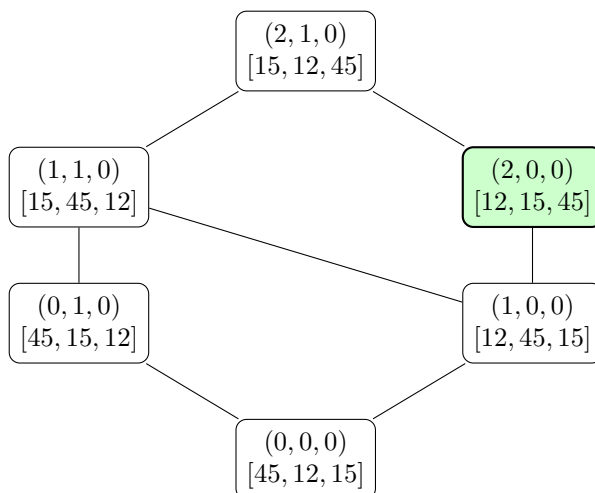


Figure 4.2: Lattice of inversion vectors for arrays of size three, applied to $A = [45, 12, 15]$. Each node shows an inversion vector $(a, b, 0)$ and the array obtained by applying the corresponding permutation to A . The sorted array $[12, 15, 45]$ sits at $(2, 0, 0)$ (shaded); sorting reduces to advancing upward in the lattice to that node.

We formalize when an inversion vector is *feasible*. Suppose G is less than the unique inversion vector that sorts A . Then, when G is applied to A , the array is not sorted, so there exists an index i with a missing inversion: some $j > i$ with $A[j] < A[i]$. Any inversion vector in which the count for index i is not increased cannot reach the sorted array. Rather than maintain G explicitly, we keep only the effect of applying G to A — that is, we mutate A in place.

The simplest choice of forbidden predicate checks only *immediate* inversions: $A[i] > A[i + 1]$. Swapping

$A[i]$ and $A[i + 1]$ advances $G[i]$ by one. This yields Algorithm LLP-Sort-Adjacent.

Algorithm LLP-Sort-Adjacent: Sorting via Adjacent Swaps

Input: $A[1..n]$: array of int

Output: $A[1..n]$: sorted array

```

1 forbidden( $j$ ):  $A[j] > A[j + 1]$  ; //  $G[j]$  misses an inversion
2 advance( $j$ ):  $swap(A[j], A[j + 1])$  ; // Increment  $G[j]$ 

```

Both Bubble Sort and Insertion Sort are deterministic schedules of LLP-Sort-Adjacent: they fix an order in which the forbidden indices are evaluated (round-robin left-to-right for Bubble Sort; incremental for Insertion Sort). The generic LLP framework, by contrast, is nondeterministic — multiple indices may be forbidden at any point, and any of them may advance.

A slight generalization widens the forbidden predicate to witness *any* right-hand index, not just the immediate neighbor. Algorithm LLP-Sort-Pairwise declares j forbidden whenever any $k > j$ has $A[j] > A[k]$, and advances by swapping $A[j]$ with $A[k]$.

Algorithm LLP-Sort-Pairwise: Sorting via Non-Adjacent Swaps

Input: $A[1..n]$: array of int

Output: $A[1..n]$: sorted array

```

1 forbidden( $j$ ):  $\exists k > j : A[j] > A[k]$  ; // pick such  $k$ 
2 advance( $j$ ):  $swap(A[j], A[k])$  ; // advance  $G[j]$ 

```

The forbidden predicate of LLP-Sort-Pairwise implies that of LLP-Sort-Adjacent, so any execution of LLP-Sort-Adjacent is also a valid execution of LLP-Sort-Pairwise. The extra freedom in choosing k lets a single advance step resolve multiple inversions at once; Quicksort can be viewed as an implementation of LLP-Sort-Pairwise where the pivot induces a batched advance: every j on the less-than-pivot side and every k on the greater-than-pivot side are resolved in a single partition pass.

4.8 Summary

Table 4.1 lists the sorting algorithms discussed in this chapter, with their sequential time complexities.

Problem	Algorithm	Time
Sorting	Bubble Sort	$O(n^2)$
Sorting	Insertion Sort	$O(n^2)$
Sorting	Merge Sort	$O(n \log n)$
Sorting	QuickSort	$O(n^2)$ worst, $O(n \log n)$ avg
Sorting	Radix Sort	$O(kn)$

Table 4.1: Sequential Sorting Algorithms

4.9 Problems

1. Implement the binary search algorithm discussed in Section 4.1.
2. Show that any algorithm that is based on comparison of consecutive entries must take $\Omega(n^2)$ comparisons in the worst case.

3. Give an algorithm to merge k sorted arrays of size n into a single sorted array.
4. We have partitioned the array into three parts in the QuickSort algorithm. Give a version of the QuickSort algorithm in which the array is partitioned only in two parts: entries that are less than *pivot* and the entries that are greater than or equal to the pivot.
5. Give a randomized version of Quicksort in which the pivot is chosen at random. Give the expected running time of your algorithm.
6. (Stability.) A sorting algorithm is *stable* if it preserves the relative order of records with equal keys. State which of the following are stable, and justify each: Bubble Sort, Insertion Sort, Merge Sort, QuickSort (Lomuto and Hoare partitions), and Radix Sort.
7. (Counting Sort.) Given an array $A[1..n]$ of integers in the range $[0, k]$, give an algorithm that sorts A in $O(n + k)$ time. Discuss when Counting Sort is preferable to Radix Sort.
8. (External Merge Sort.) Suppose the array $A[1..n]$ does not fit in memory. RAM holds $M \ll n$ elements, and reads and writes to disk occur in blocks of B elements. Describe an external-memory Merge Sort and count the number of block I/Os it performs.

4.10 Bibliographic Remarks

Quicksort was invented by Hoare in 1961 [Hoa61]; the original paper already contained a two-way in-place partition. The three-way “Dutch National Flag” partition (Algorithm [ThreeWayPartition](#)) is due to Dijkstra and appears in *A Discipline of Programming* [Dij76]. Merge sort is usually attributed to von Neumann in 1945, and its recurrence-based analysis is given in Knuth’s *The Art of Computer Programming, Vol. 3: Sorting and Searching* [Knu98], which also catalogues Bubble Sort, Insertion Sort, and numerous refinements. Radix sort predates digital computers: it was the basis of Herman Hollerith’s electromechanical tabulator used for the 1890 U.S. Census, and a modern analysis (including stability via Counting Sort as the inner pass) appears in [Knu98]. The $\Omega(n \log n)$ lower bound for comparison-based sorting via the decision-tree argument is also treated in [Knu98].

Chapter 5

Graphs

5.1 Introduction

Graph theory, a cornerstone of algorithmic design, is the foundation of numerous modern applications in diverse domains. In *social network analysis*, graphs model relationships where vertices represent individuals and edges denote interactions, enabling the detection of communities using algorithms such as clustering. In *transportation and logistics*, directed graphs optimize routing: companies like Amazon use shortest-path algorithms (for example, Dijkstra’s algorithm) in road networks to minimize delivery times, while airlines use bipartite matching to schedule crews efficiently. *Computer networks* rely on graphs to represent topologies, with spanning tree protocols ensuring loop-free communication and centrality measures identifying critical nodes for cybersecurity. In *bioinformatics*, graphs map protein interactions or gene regulatory networks, where traversal algorithms identify functional pathways, helping drug discovery.

Graphs come in numerous varieties. They may be *directed* or *undirected*. They may be *simple* or with loops and parallel edges. They may be *weighted* or *unweighted*. They may be *acyclic* or not. For example, when modeling transportation networks, the nodes may represent important milestones in a city. A directed edge may represent a one-way street, and an undirected edge may represent a street that can be traversed in either direction. The weights may represent the distance between the nodes.

In this chapter, we cover some basic traversal algorithms in a simple directed graph. This chapter is organized as follows. Section 5.2 describes various commonly used representations of graphs. Section 5.3 gives the breadth-first-search (BFS) method of traversing a graph. Section 5.4 gives the depth-first-search (DFS) method of traversing a graph. Section 5.5 gives an algorithm to “layer” a directed acyclic graph. Section 5.6 gives an algorithm to compute the strongly connected components of a directed graph. Section 5.7 revisits these problems from the Lattice-Linear Predicate (LLP) perspective. Many other algorithms on graphs, such as the shortest path algorithms and minimum spanning tree algorithms, are covered in later chapters.

5.2 Graph Representations

We present an undirected graph and its directed counterpart, each with vertices v_0, v_1, v_2, v_3 , shown with adjacency matrix and adjacency list representations (as linked lists).

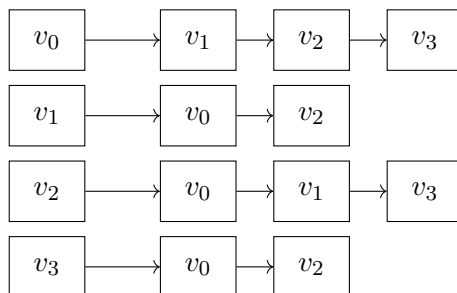


Figure 5.2: Adjacency list representation of an undirected graph

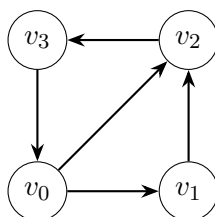


Figure 5.3: A directed graph.

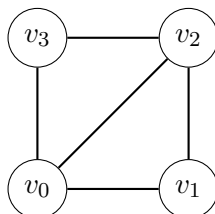


Figure 5.1: An undirected graph

The graph in Fig. 5.1 can be represented using an adjacency matrix as follows.

$$\begin{bmatrix}
 & v_0 & v_1 & v_2 & v_3 \\
 v_0 & 0 & 1 & 1 & 1 \\
 v_1 & 1 & 0 & 1 & 0 \\
 v_2 & 1 & 1 & 0 & 1 \\
 v_3 & 1 & 0 & 1 & 0
 \end{bmatrix}$$

An entry of 1 indicates an edge; the matrix is symmetric. Such a representation allows one to answer any question of the form: is there an edge between v_i and v_j in constant time. However, it takes $O(n^2)$ space irrespective of the number of edges in the graph. When the graph is sparse, the adjacency list representation is less wasteful. Fig. 5.2 shows the adjacency list representation of the same graph.

We now consider the directed version of the graph in Fig. 5.3.

As before, we can have the adjacency matrix and adjacency list representations shown below.

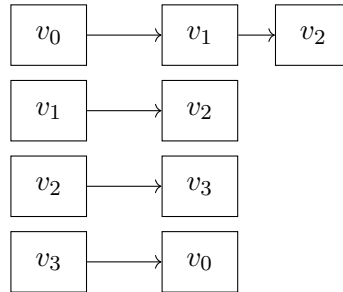


Figure 5.4: Adjacency representation of a directed graph

$$\begin{bmatrix}
 & v_0 & v_1 & v_2 & v_3 \\
 v_0 & 0 & 1 & 1 & 0 \\
 v_1 & 0 & 0 & 1 & 0 \\
 v_2 & 0 & 0 & 0 & 1 \\
 v_3 & 1 & 0 & 0 & 0
 \end{bmatrix}$$

5.3 Breadth-First Search (BFS)

In this section, we describe the breadth-first-search (BFS), one of the two most common ways of traversing a directed graph (the other being depth-first search, covered in the next section). Traversal is required for many reasons. We may be interested in searching for a node, finding the number of components in a graph, or checking for some user-specified property. We first present a general (nondeterministic) traversal algorithm; different strategies for choosing which vertex to explore next yield BFS or DFS as special cases.

General Traversal of a Directed Graph

In Algorithm [Traversal](#), we keep G as the set of vertices that can be reached from the vertex v_0 . We also keep S as a set of vertices that have been visited but not explored. Using different strategies to remove an index from S , we can traverse the graph in a different order. If S is maintained as a queue, the removal corresponds to removing from the head of the queue, and the addition corresponds to appending at the end of the queue, then we traverse the graph in a *breadth-first* manner. If S is maintained as a stack such that removal corresponds to the *pop()* and addition corresponds to the *push()* method on the stack, then we traverse the graph in a depth-first search manner.

Each vertex is added to and removed from S at most once, and each edge (j, k) is examined exactly once (when j is being explored). Hence Algorithm [Traversal](#) runs in $O(n + m)$ time.

Suppose that we are given one of the nodes v_0 in the graph. Our task is to find all the nodes that are reachable from v_0 . Algorithm [BFS-Traversal](#) can be used for this purpose. It will generate a tree rooted at v_0 such that every node is at the minimum distance from v_0 . We maintain a variable *parent* that gives the parent of any node x in the graph. If x is reachable from v_0 , then following the *parent* pointer from x to v_0 , we will get a shortest path from v_0 to x . A node v_i has its parent set to v_j if v_j is along a path from v_0 to v_i and has its distance one less than that of v_i .

Algorithm Traversal: Finding the reachable set of vertices from v_0 in a Directed Graph

Input: $dep(j)$: adjacency list for each vertex j
Output: $G[0..n-1]$: vector of int initially $\forall i : G[i] = 0$

```

1  $S$ : set of indices;
2  $S.add(0)$  // add 0 as the initial vertex to start
3  $G[0] := 1$ ;
4 while  $\neg S.empty()$  do
5    $j := S.removeAny(G)$  // remove any index from  $S$ 
6   forall ( $k \in dep(j)$ ) do
7     if ( $G[k] = 0$ ) then
8        $G[k] := 1$  // advance on  $k$ 
9        $S.add(k)$  // update frontier
10    end
11  end
12 end
13 return  $G$ ; // the reachable set

```

Algorithm BFS-Traversal: Algorithm BFS to find the Breadth-First-Search Tree in a Directed Graph

Input: $dep(j)$: adjacency list for each vertex j ; source vertex s
Output: $G[0..n-1]$: vector of int initially $\forall i : G[i] = maxint$; $parent[0..n-1]$: vector of int initially $\forall i : parent[i] = -1$

```

1  $S$ : queue of indices;
2  $S.add(s)$ ;
3  $G[s] := 0$ ;
4 while  $\neg S.empty()$  do
5    $j := S.removeFirst()$ ;
6   // remove an index from  $S$  forall ( $k \in dep(j)$ ) do
7     if ( $G[k] > G[j] + 1$ ) then
8        $G[k] := G[j] + 1$ ;
9        $parent[k] := j$ ;
10    append  $k$  to  $S$ ;
11  end
12 end
13 end
14 return  $G$ ;
15 // the BFS tree

```

Each vertex is enqueued at most once and dequeued at most once; each edge (j, k) is examined exactly once when j is dequeued. Hence Algorithm [BFS-Traversal](#) runs in $O(n + m)$ time.

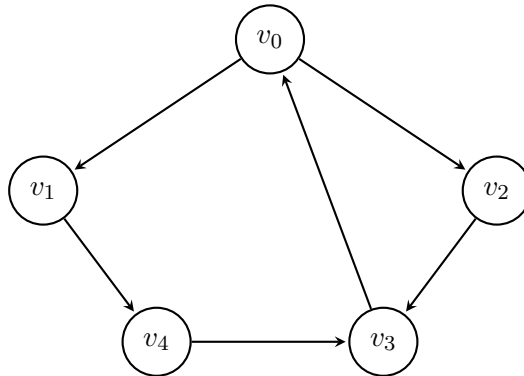


Figure 5.5: A directed graph used to illustrate BFS and DFS traversals.

Example 5.1 [BFS on a directed graph] Run BFS from source v_0 on the graph in Figure 5.5. Initially $G[v_0] = 0$ and the queue is $[v_0]$.

- Dequeue v_0 . Neighbours v_1, v_2 have $G = \infty$; set $G[v_1] = 1, G[v_2] = 1$, $parent[v_1] = parent[v_2] = v_0$, enqueue both.
- Dequeue v_1 . Neighbour v_4 : set $G[v_4] = 2$, $parent[v_4] = v_1$, enqueue v_4 .
- Dequeue v_2 . Neighbour v_3 : set $G[v_3] = 2$, $parent[v_3] = v_2$, enqueue v_3 .
- Dequeue v_4 . Neighbour v_3 already has $G[v_3] = 2$ (not improved).
- Dequeue v_3 . Neighbour v_0 already has $G[v_0] = 0$ (not improved).

Final distances from v_0 :

Node	$G[\cdot]$ (distance)	$parent[\cdot]$
v_0	0	—
v_1	1	v_0
v_2	1	v_0
v_4	2	v_1
v_3	2	v_2

The BFS tree is shown in Figure 5.6.

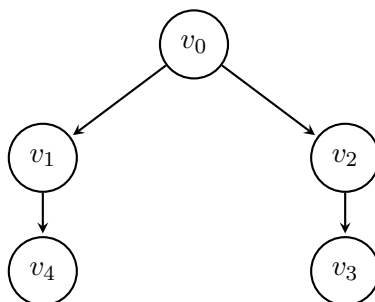


Figure 5.6: BFS tree rooted at v_0 for Example 5.1. Every vertex appears at its shortest distance from v_0 .

5.4 Depth-First Search (DFS)

Another way to traverse a graph is the depth-first search method. While the breadth-first search traverses the graph one layer at a time, the depth-first search continues to take a path as far as it can go, and then it backtracks to traverse the remaining graph (in the same fashion!).

Algorithm DFS-Traversal: Sequential Algorithm DFS to find the Depth-First-Search Tree in a Directed Graph

Input: $dep(j)$: adjacency list for each vertex j

Output: $G[0..n-1]$: int, initially 0; $parent[0..n-1]$: int, initially -1 ; $discovered[0..n-1]$: int; $finished[0..n-1]$: int

```

1 tick: int, initially 1;
2 Function DFS( $j$ ):
3    $G[j] := 1$ ;
4    $discovered[j] := tick$ ;
5    $tick := tick + 1$ ;
6   for  $k \in dep(j)$  do
7     if  $G[k] = 0$  then
8        $parent[k] := j$ ;
9       DFS( $k$ );
10    end
11  end
12   $finished[j] := tick$ ;
13   $tick := tick + 1$ ;
14 DFS(0);
  
```

The recursive call $Dfs(k)$ is made for each vertex k at most once, and each edge (j, k) is examined exactly once when j 's adjacency list is scanned. Hence Algorithm [DFS-Traversal](#) runs in $O(n + m)$ time.

Algorithm [DFS-Traversal](#) visits all vertices reachable from vertex v_0 in a depth-first search manner. The variable G is used to mark all the vertices that can be reached from v_0 . The variable S is used as a stack to store the vertices while traversing the graph. The variable $discovered$ is used to store the time (tick) when each vertex in the graph is explored. The variable $finished$ is used to store the tick when the algorithm is finished exploring a vertex. The variable $parent$ is used to store the depth-first tree. Anytime

a vertex k is visited for the first time from a vertex j , the $parent[k]$ is recorded as j .

The variables *discovered* and *finished* are used only to record the *tick* when a vertex is starting to be explored and finished for exploration.

Example 5.2 [DFS on a directed graph] Run DFS from source v_0 on the graph in Figure 5.5. From v_0 we visit v_1 ; from v_1 we visit v_4 ; from v_4 we visit v_3 . When v_3 is explored, its only outgoing edge points to v_0 , which is already visited, so v_3 finishes. We backtrack to v_4 (no other outgoing edges), then to v_1 , and finally to v_0 . From v_0 we now visit v_2 , whose outgoing edge leads to v_3 (already visited); v_2 finishes, and DFS terminates.

The resulting *discovered* and *finished* ticks are:

Node	<i>discovered</i>	<i>finished</i>
v_0	1	10
v_1	2	7
v_4	3	6
v_3	4	5
v_2	8	9

The DFS tree is shown in Figure 5.7.

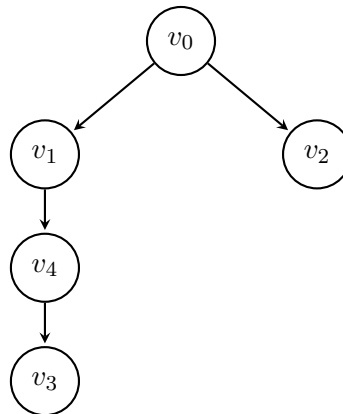


Figure 5.7: DFS tree rooted at v_0 for Example 5.2. The edge $v_2 \rightarrow v_3$ of the original graph is a *cross edge* (not part of the DFS tree, since v_3 is discovered earlier via v_4).

5.5 Layering of a Directed Acyclic Graph

Suppose that we are given a directed graph with no cycles. This graph can be “layered” as follows. Every vertex i in the graph is assigned a number $G[i]$ such that if there is a directed edge from i to j , then the number assigned to i is strictly less than that of j , i.e., for all edges $(i, j) \in E$, $G[i]$ is less than $G[j]$. An application of this concept is the prerequisite structure of courses in a university, and the integer assigned to the course corresponds to the earliest semester in which the course can be taken by a student.

Our goal is to compute, for each vertex j , the integer $G[j]$ equal to the length of the longest path from any source vertex (a vertex with no incoming edges) to j . We present a classical algorithm that maintains the in-degree of every vertex and processes vertices in the order in which all of their predecessors have been assigned layers.

The idea is simple. Initially, every vertex with in-degree 0 lies at layer 0. We place all such vertices in a queue. We then repeatedly remove a vertex j from the queue, and for every successor k of j (i.e., $(j, k) \in E$), we decrement k 's in-degree. Whenever k 's in-degree becomes 0, we have seen all of k 's predecessors and know their layers; we then set $G[k]$ to one more than the maximum of $G[i]$ over all predecessors i of k , and enqueue k . Algorithm [Layering](#) gives the details.

Algorithm Layering: Layering of a Directed Acyclic Graph.

Input: Directed acyclic graph (V, E) with $|V| = n$, $|E| = m$; $pre(j)$ the list of predecessors of j ;
 $succ(j)$ the list of successors of j

Output: $G[0..n - 1]$: layer of each vertex

```

1   $indeg[j] := |pre(j)|$  for every  $j$ ;
2   $Q$ : queue of indices initially empty;
3  forall  $j \in V$  do
4      if  $indeg[j] = 0$  then
5           $G[j] := 0$ ;
6          enqueue  $j$  into  $Q$ ;
7      end
8  end
9  while  $\neg Q.empty()$  do
10      $j := Q.removeFirst()$ ;
11     forall  $k \in succ(j)$  do
12          $indeg[k] := indeg[k] - 1$ ;
13         if  $indeg[k] = 0$  then
14              $G[k] := 1 + \max_{i \in pre(k)} G[i]$ ;
15             enqueue  $k$  into  $Q$ ;
16         end
17     end
18 end
19 return  $G$ ;

```

Each vertex is enqueued and dequeued at most once, and each edge (j, k) is processed exactly once when j is dequeued. Hence the time complexity of the algorithm is $O(n + m)$.

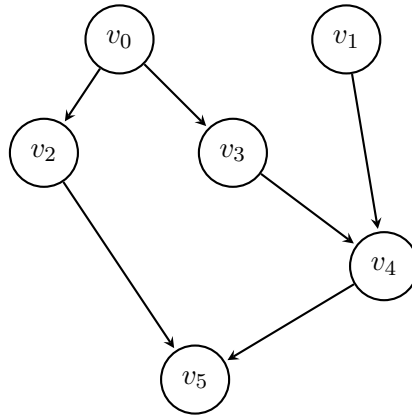


Figure 5.8: A directed acyclic graph used to illustrate layering.

Example 5.3 [Layering a DAG] Run the layering algorithm on the DAG of Figure 5.8. In-degrees are $\text{indeg}[v_0] = \text{indeg}[v_1] = 0$, $\text{indeg}[v_2] = \text{indeg}[v_3] = 1$, $\text{indeg}[v_4] = \text{indeg}[v_5] = 2$.

- Initially v_0 and v_1 have in-degree 0; set $G[v_0] = G[v_1] = 0$ and enqueue both. Queue: $[v_0, v_1]$.
- Dequeue v_0 . Successors v_2, v_3 : their in-degrees drop to 0; set $G[v_2] = G[v_3] = 1$ and enqueue them.
- Dequeue v_1 . Successor v_4 : in-degree drops to 1 (not yet ready).
- Dequeue v_2 . Successor v_5 : in-degree drops to 1 (not yet ready).
- Dequeue v_3 . Successor v_4 : in-degree drops to 0; set $G[v_4] = 1 + \max(G[v_1], G[v_3]) = 1 + \max(0, 1) = 2$ and enqueue v_4 .
- Dequeue v_4 . Successor v_5 : in-degree drops to 0; set $G[v_5] = 1 + \max(G[v_2], G[v_4]) = 1 + \max(1, 2) = 3$ and enqueue v_5 .
- Dequeue v_5 (no successors).

The computed layers are

Vertex	$G[\cdot]$ (layer)
v_0, v_1	0
v_2, v_3	1
v_4	2
v_5	3

The layered view of the DAG is shown in Figure 5.9. Each $G[j]$ equals the length of the longest path from any source to j ; the longest path to v_5 is $v_0 \rightarrow v_3 \rightarrow v_4 \rightarrow v_5$ of length 3.

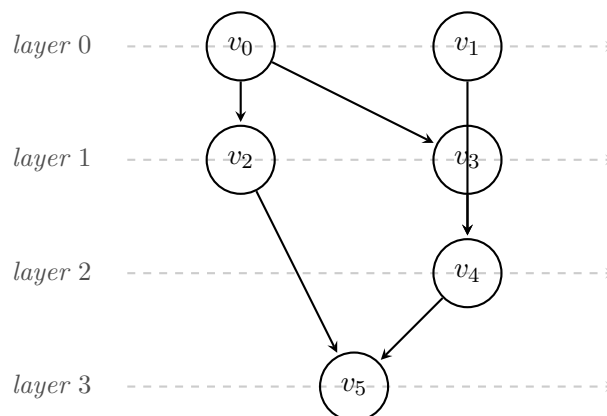


Figure 5.9: The DAG of Figure 5.8 redrawn with each vertex placed on its computed layer. Every directed edge goes from a strictly smaller layer to a strictly greater one.

In many applications, we are interested in coming up with a *total order* on all vertices such that for all edges (i, j) , $G[i]$ is strictly less than $G[j]$. This operation is called *topological sort* of a directed acyclic graph. The reader is invited to develop an algorithm for such applications.

5.6 Strongly Connected Components of a Directed Graph

Given a directed graph $G = (V, E)$, two vertices u and v are said to be in the same *strongly connected component* (SCC) if there is a directed path from u to v and a directed path from v to u . The relation “ u and v are mutually reachable” is an equivalence relation, so the vertex set V partitions into maximal strongly connected components. Collapsing each SCC into a single super-vertex yields the *component graph* G^{scc} , which is always a directed acyclic graph.

Layering handles the acyclic case; SCC computation is the natural counterpart for graphs that may contain cycles. SCC decomposition is used, for example, in compiler dataflow analysis (to identify strongly connected regions of a control-flow graph), in dependency resolution (to detect cyclic package or module dependencies), and in 2-SAT (where satisfiability reduces to computing SCCs of the implication graph).

We present Kosaraju’s algorithm, a simple and efficient $O(n + m)$ -time algorithm that uses two depth-first searches (DFS). Its correctness rests on the following lemma.

Lemma 5.4 *Let u and v be vertices in distinct strongly connected components C_u and C_v of G . Let $f(C) = \max_{w \in C} f[w]$ denote the maximum DFS finishing time among vertices of component C . If there is an edge in G^{scc} from C_u to C_v , then $f(C_u) > f(C_v)$.*

Proof: Consider the DFS executed on G and the vertex that is visited first among $C_u \cup C_v$. If it is in C_u , then by white-path reachability the DFS explores all of C_u and, via the cross edge, all of C_v before finishing the start vertex, so $f(C_u) > f(C_v)$. If instead the first visited vertex lies in C_v , then DFS finishes C_v first (since no edge returns from C_v to C_u in G^{scc}); the earliest vertex visited in C_u is explored later, and its finishing time — hence $f(C_u)$ — is strictly greater than $f(C_v)$. ■

As a consequence, if we process vertices in decreasing order of finishing time from the first DFS but on the *transpose* graph G^T (the graph obtained by reversing every edge), then each DFS tree from a new root is exactly one SCC. Edges in G^T go from lower- f components to higher- f components, so a DFS from the vertex of maximum f in G^T cannot leave its SCC; after that SCC is exhausted, we pick the next unvisited vertex with the largest remaining f , and so on.

Algorithm Kosaraju-SCC: Kosaraju's Algorithm for Strongly Connected Components.

Input: Directed graph $G = (V, E)$ with $|V| = n$, $|E| = m$

Output: $C[1..n]$: $C[v]$ is the identifier of the SCC containing v

// Pass 1: finishing-time DFS on G

- 1 Perform DFS on G and record the finishing time $f[v]$ for every vertex v ;
- 2 Let $S = [v_1, v_2, \dots, v_n]$ be the list of vertices sorted in decreasing order of f ;

// Pass 2: DFS on the transpose in reverse finishing order

- 3 Construct $G^T = (V, E^T)$ where $E^T = \{(v, u) \mid (u, v) \in E\}$;
 - 4 Mark all vertices in G^T as unvisited;
 - 5 $k := 0$;
 - 6 **for** $i = 1$ to n **do**
 - 7 **if** v_i is unvisited in G^T **then**
 - 8 $k := k + 1$;
 - 9 Run DFS in G^T starting at v_i ; for every vertex w reached, set $C[w] := k$ and mark w visited;
 - 10 **end**
 - 11 **end**
 - 12 **return** C
-

Theorem 5.5 *Algorithm Kosaraju-SCC computes the strongly connected components of G in $O(n+m)$ time.*

Proof: Correctness. By Lemma 5.4, for every edge $(C_u, C_v) \in G^{\text{scc}}$ we have $f(C_u) > f(C_v)$. Reversing all edges gives G^T with $(C_v, C_u) \in G^{\text{scc}, T}$; consequently, when we launch a DFS in G^T from the vertex with the largest f -value in its component, no transpose edge can leave the component, so the reached set is exactly that SCC. Subsequent iterations remove fully visited components and repeat on the remaining sub-DAG.

Complexity. Each of the two DFS passes runs in $O(n+m)$ time. Constructing G^T also takes $O(n+m)$ time using adjacency lists. Sorting vertices by finishing time can be done implicitly by pushing each vertex onto a stack when it finishes during the first DFS, so no additional sorting cost is incurred.

■

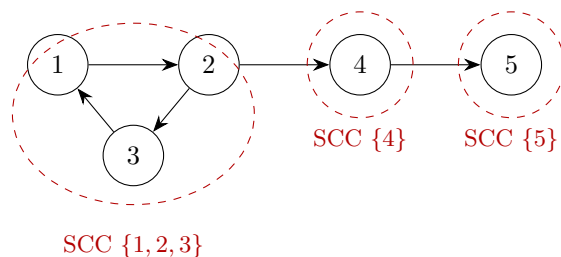


Figure 5.10: Example directed graph for Kosaraju’s algorithm. The three strongly connected components are $\{1, 2, 3\}$, $\{4\}$, and $\{5\}$.

Example 5.6 Consider the directed graph on $\{1, \dots, 5\}$ with edges E shown in Figure 5.10, where

$$E = \{(1, 2), (2, 3), (3, 1), (2, 4), (4, 5)\}.$$

DFS starting at vertex 1 finishes the vertices in order 3, 5, 4, 2, 1, giving finishing times $f[3]=1$, $f[5]=2$, $f[4]=3$, $f[2]=4$, $f[1]=5$. The transpose graph has edges

$$E^T = \{(2, 1), (3, 2), (1, 3), (4, 2), (5, 4)\}.$$

Processing vertices in decreasing order of f on the transpose yields three DFS trees: $\{1, 2, 3\}$ from vertex 1, $\{4\}$ from vertex 4, and $\{5\}$ from vertex 5, matching the three SCCs circled in Figure 5.10.

5.7 LLP Perspective

The graph algorithms of this chapter can also be expressed in the Lattice-Linear Predicate (LLP) framework of Chapter 2. In the LLP view, the algorithm designer specifies (a) a lattice of candidate solutions, (b) a predicate B describing feasibility, (c) a *forbidden* condition identifying indices whose current value prevents B from holding, and (d) an *advance* rule that moves a forbidden index towards feasibility. Each iteration of an LLP algorithm picks any forbidden index and advances it; the specification does not fix a schedule, leaving the designer free to choose an evaluation order that yields an efficient implementation.

Graph traversal as an LLP algorithm

Before considering BFS, which tracks distances, we revisit the plain reachability problem of Section 5.3: given a source vertex v_0 , compute the set of vertices reachable from v_0 . We maintain a Boolean vector G with $G[0] = 1$ and $G[j] = 0$ for $j \neq 0$ and search for the least vector satisfying

$$B_{reach}(G) \equiv G[0] = 1 \wedge \forall (i, j) \in E : (G[i] = 1 \Rightarrow G[j] = 1).$$

An index j is forbidden when some predecessor i has $G[i] = 1$ but $G[j] = 0$; the advance rule sets $G[j] := 1$. Algorithm [LLP-Traversal](#) captures this.

Algorithm LLP-Traversal: Reachable set of vertices via LLP.

- 1 Process P_j ;
Input: $edges(j)$: list of adjacent vertices of j
Output: $G[j]$: int, initially 1 if $j = 0$, else 0
 - 2 **forbidden**(j): $\exists i : j \in edges(i) \wedge G[i] = 1 \wedge G[j] = 0$;
 - 3 **advance**(j): $G[j] := 1$;
-

Each $G[j]$ advances at most once (from 0 to 1), and evaluating the forbidden predicate amortises to $O(n + m)$ work across all indices. The sequential running time is therefore $O(n + m)$.

BFS as an LLP algorithm

For breadth-first search from a source vertex v_0 we maintain a distance vector G with $G[0] = 0$ and $G[j] = \infty$ for $j \neq 0$. We want the least vector satisfying

$$B_{bfs}(G) \equiv \forall j \neq 0 : \forall (i, j) \in E : G[j] \leq G[i] + 1.$$

An index j is forbidden if it has a predecessor i with $G[j] > G[i] + 1$; the advance rule sets $G[j] := G[i] + 1$. Algorithm [LLP-BFS](#) gives the resulting LLP algorithm.

Algorithm LLP-BFS: LLP Program for LLP-BFS

- Input:** pre : array of sets of int; G : array of int
- 1 **forbidden**(j): $G[j] > \min\{G[i] + 1 \mid i \in pre(j)\}$;
 - 2 **advance**: $G[j] := \min\{G[i] + 1 \mid i \in pre(j)\}$;
-

A sequential implementation that evaluates forbidden indices in breadth-first order runs in $O(n + m)$ time and matches the classical BFS bound.

The LLP specification above does not fix an order in which forbidden indices are evaluated. In sequential or resource-constrained settings, it can be advantageous to prioritise which forbidden index is processed first. We write the prioritisation with a **priority** clause alongside **forbidden** and **advance**. In Algorithm [LLP-Traversal-BFS-priority](#), priority is given by the current value of $G[j]$, so the search explores vertices in order of distance from v_0 — exactly as in classical BFS with a queue.

Algorithm LLP-Traversal-BFS-priority: Prioritised BFS traversal via LLP.

- 1 Process P_j ;
Input: $pre(j)$: list of predecessors of vertex j
Output: $G[j]$: int, initially 0 if $j = 0$, else ∞
 - 2 **ensure**(j): $G[j] \leq \min\{G[i] + 1 \mid i \in pre(j)\}$;
 - 3 **priority**: $G[j]$
-

Processing indices in increasing $G[j]$ order using a priority queue keyed on $G[j]$ matches the classical BFS schedule and gives a sequential running time of $O(n + m)$.

Layering as an LLP algorithm

For layering a directed acyclic graph we want the least vector G of natural numbers satisfying

$$B_{layer}(G) \equiv \forall (i, j) \in E : G[i] < G[j].$$

Define $fixed[j]$ to be true once $G[j]$ has been finalised. An index j is forbidden when it is not fixed but all of its predecessors are fixed; the advance rule then sets $G[j]$ to one more than the maximum layer of its predecessors. Algorithm [LLP-Layering](#) captures this.

Algorithm LLP-Layering: Layering of a directed acyclic graph via LLP.

1 Process P_j ;
Input: $pre(j)$: list of predecessors of vertex j
Output: $G[j]$: int, initially 0; $fixed[j]$: boolean, initially *true* if $pre(j) = \{\}$, else *false*
2 **forbidden**(j): $\neg fixed[j] \wedge (\forall i \in pre(j) : fixed[i])$;
3 **advance**(j): $G[j] := \max_{i \in pre(j)} G[i] + 1$; $fixed[j] := true$;

A sequential implementation that evaluates forbidden indices in any topological order runs in $O(n+m)$ time.

5.8 Summary

This chapter introduced fundamental graph traversals (BFS and DFS) and showed that several derived problems — BFS layer assignment, layering of a DAG, connected components, and strongly connected components — admit clean LLP formulations on top of the same primitives.

Table [5.1](#) summarizes the time complexities of the sequential algorithms discussed in this chapter.

Problem	Algorithm	Time Complexity
Breadth-First Search	BFS-Traversal	$O(m+n)$
Depth-First Search	DFS-Traversal	$O(m+n)$
Breadth-First Search	LLP-BFS-Traversal	$O(m+n)$
Layering	LLP-Layering	$O(m+n)$
Strongly Connected Components	Kosaraju-SCC	$O(m+n)$

Table 5.1: Algorithms discussed in this chapter.

5.9 Problems

1. Given an undirected graph with n vertices and m edges, find the size (number of vertices) of each connected component using BFS. The graph is represented as an adjacency list. Return a list of sizes for all components.
2. Given an undirected graph with n vertices and m edges, determine if the graph contains a cycle using DFS traversal. The graph is represented as an adjacency list.
3. Give an efficient algorithm to check if the given undirected graph $G = (V, E)$ is bipartite.
4. Give an algorithm to find the number of connected components in an undirected graph using DFS.
5. Give an efficient algorithm to find all bridges in an undirected graph. A bridge is an edge whose removal increases the number of connected components.

6. Given an undirected graph with n vertices and m edges, find all articulation points using DFS traversal. An articulation point is a vertex whose removal increases the number of connected components in the graph.
7. Algorithm [Layering](#) gives a level number $G[i]$ for each vertex i such that for any edge (i, j) , we have that $G[i]$ is strictly less than $G[j]$. Modify the algorithm to generate a total order on all vertices.
8. Algorithm [Layering](#) gives us the least integral labels satisfying that, for all edges (i, j) , $G[i]$ is strictly less than $G[j]$. Such a property is possible only for acyclic graphs. Modify the algorithm to output “error” whenever there is a cycle in the input graph.
9. Algorithm [Layering](#) works by assigning level numbers to the minimal vertices in the graph and then to the next layer of vertices. Give the dual algorithm that starts with assigning level number to the maximal vertices and then to the layer of vertices that are below the maximal level.
10. For the connected components problem, we had marked each vertex with a label from $1..n$ such that all vertices in the same component had the same label. Now suppose that our goal is to find out the total number of connected components. Give an $O(\log n)$ parallel algorithm to do so. Assume that you are allowed to have additional arrays of $O(n)$.

5.10 Bibliographic Remarks

Breadth-first and depth-first search are classical graph-traversal techniques; both are treated in detail in Cormen, Leiserson, Rivest, and Stein [[CLRS01](#)]. The layering of a directed acyclic graph (equivalently, topological sort by longest-path length) is covered in most algorithms texts, including [[CLRS01](#)]. The in-degree-based approach used in Algorithm [Layering](#) is due to A. B. Kahn (1962). Algorithm Kosaraju-SCC for strongly connected components (two DFS passes on G and G^T) is due to S. Rao Kosaraju (unpublished, 1978) and was independently described by M. Sharir in 1981. An alternative $O(n + m)$ -time one-pass algorithm based on DFS low-link values is due to R. E. Tarjan (1972). A full treatment of both algorithms is given in [[CLRS01](#)]. 2-SAT, which reduces in linear time to SCC of the implication graph, is due to Aspvall, Plass, and Tarjan (1979) and is discussed in Chapter [17](#).

Chapter 6

Greedy Algorithms

6.1 Introduction

Many problems can be solved by making multiple choices such that at the end of these choices, we have the solution to the problem. The greedy algorithm proceeds as follows: at every step, commit irrevocably to the locally-best choice under a fixed selection rule, and never revisit past choices. The algorithm typically starts with the trivial solution of size 0 or 1 and increases the size of the partial solution one step at a time until the original problem of size n is solved. For many applications, this strategy results in a global optimum or a provably near-optimal solution.

A greedy algorithm is cheaper than exhaustive search or dynamic programming precisely because it never backtracks, but this restriction is also its weakness: committing too early can miss the global optimum. Part of the task of this chapter is to identify problems and greedy rules that *provably* yield an optimum, and to prove correctness using the **exchange argument**: take any optimal solution O that disagrees with the greedy solution G , identify the earliest point of disagreement, and show that swapping G 's choice into O produces a solution at least as good; iterating transforms O into G without loss.

This chapter is organized as follows. Section 6.2 describes a greedy algorithm for the interval scheduling problem. In this problem, we schedule a maximum number of jobs on a processor. Section 6.3 describes a greedy algorithm for the problem of scheduling intervals using as few rooms as possible so that no two overlapping intervals are assigned to the same room. Section 6.4 discusses the problem of scheduling n jobs so that the maximum lateness of jobs is minimized. Section 6.5 discusses the construction of a Huffman tree that minimizes the expected length of the binary code for a symbol. Section 6.7 describes the LLP versions of these algorithms.

6.2 Interval Scheduling Problem

We start with a simple greedy algorithm. Suppose that we have n intervals with start times s_i and finish times f_i for jobs $i = 1..n$, where $s_i < f_i$. We assume that activity i occurs during the half-open interval $[s_i, f_i)$. Two intervals i and j are *compatible* if $f_i \leq s_j$ or $f_j \leq s_i$. Our goal is to select a maximum-sized subset of mutually compatible intervals. We assume that the intervals are sorted by their finish times. We

assume that jobs with the same finish times are sorted according to their start times. Furthermore, no two intervals have identical start and finish times.

The key idea is to leave as much room as possible for future activities. By always selecting the activity that finishes earliest, we ensure that the remaining time interval is as large as possible for accommodating other compatible activities. Any activity that finishes later would only reduce the available space for subsequent choices. Thus, choosing the earliest finishing activity is the safest local decision that preserves maximum flexibility for the rest of the schedule.

Let us begin with a sequential algorithm for this problem, shown in Algorithm [Interval](#). Let G be the set of mutually compatible intervals represented as a Boolean array such that $G[i]$ is 1 iff the activity i is in the subset of mutually compatible intervals that can be selected. It is clear that the first activity (with the least finishing time) is always in G . (Why?) We now traverse over all intervals as follows. We use the variable i to denote the current activity under consideration and the variable $last$ as the last activity that was included in G . We initialize $last$ to 1. The current activity is included in G if and only if its start time is greater than or equal to the finish time of the last activity. If it is, our current activity becomes the last activity, and we explore the next activity.

Algorithm Interval: Sequential Interval Scheduling

```

Input:  $s$ : array[1.. $n$ ] of int // Start times
Input:  $f$ : array[1.. $n$ ] of int // Finish times
Output:  $G$ : array[1.. $n$ ] of 0..1 initially 0 // Selected intervals

1  $G[1] := 1;$  // First interval always selected
2 int  $last := 1;$  // Index of last selected interval
3 for int  $i := 2$  to  $n$  do
4   if  $s[i] \geq f[last]$  then
5      $G[i] := 1;$  // Interval  $i$  is compatible
6      $last := i$ 
7   end
8 end

```

The main loop visits each interval once and performs $O(1)$ work per visit, so Algorithm [Interval](#) runs in $O(n)$ time when the intervals are already sorted by finish time. If the intervals are unsorted, the initial sort dominates and the total cost is $O(n \log n)$. No extra space beyond the output array is needed, giving $O(n)$ space.

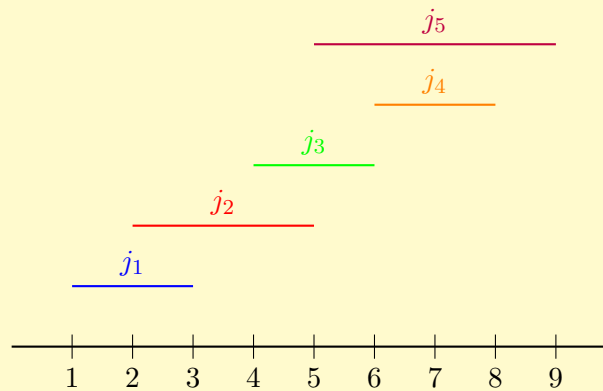
Theorem 6.1 (Interval Scheduling optimality) *Selecting activities in increasing order of finish times yields a maximum-sized set of mutually compatible activities.*

Proof: We prove the theorem using the **exchange argument**. Let G be the set of activities chosen by the greedy algorithm, which selects the activity with the earliest finish time first. Let O be an optimal solution with the maximum number of activities. The greedy algorithm selects the activity with the earliest finish time, say A_1 . Suppose O selects a different first activity A_k . Since activities are sorted by finish time, $f_1 \leq f_k$. Replacing A_k with A_1 in O still allows the remaining selections in O to proceed without reducing the total count. By repeatedly applying this argument for the next selected activity, we replace each activity in O with the corresponding activity of G without decreasing the number of selected activities.

After at most n exchanges, O is transformed into G . Since O was an optimal solution and G has the same size, G is also optimal. ■

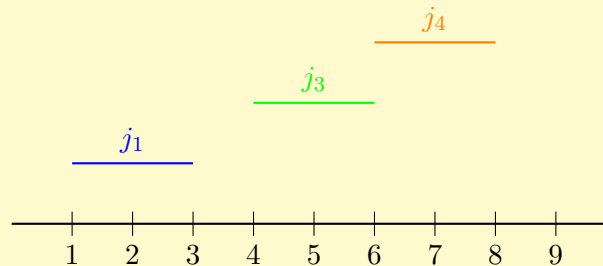
Example 6.2 Suppose that we are given five jobs, each with a start time and an end time. The goal is to select the maximum number of non-overlapping jobs.

Job	Start Time	End Time
j_1	1	3
j_2	2	5
j_3	4	6
j_4	6	8
j_5	5	9



An optimal schedule selects the maximum number of non-overlapping jobs. The optimal solution is:

$$\{j_1, j_3, j_4\}$$



6.3 Interval Partition Problem

Consider a university that offers multiple courses for students. Every course i has a fixed slot $[s_i, f_i)$ where s_i indicates the start time of the lecture in course i and f_i indicates the finish time for that lecture. There

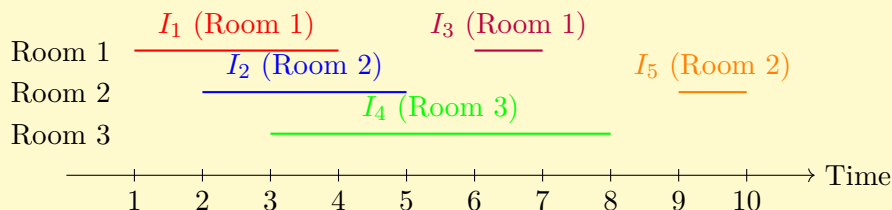
are n courses in total. The university wants to use as few classrooms as possible for lectures. Our task is to assign a room to each course so that if two lectures overlap, then they must be assigned different rooms. Since each lecture is modeled as an interval, this problem is called the *interval partition problem*.

For this problem, we first look at the lower bound on the number of rooms required. Suppose that there are m overlapping lectures. Then it is clear that any room assignment must use at least m rooms.

At any moment, the number of rooms required is determined by how many intervals overlap at that time. Therefore, the goal is simply to reuse rooms whenever possible. When a new interval begins, we assign it to the room that becomes free the earliest. If no room is free, only then do we open a new one. This greedy rule ensures that we never use more rooms than necessary, since a new room is created only when all existing rooms are simultaneously occupied.

We now show a simple greedy algorithm that achieves the lower bound on the number of rooms. We sort the courses based on the start times of the lectures. We assign courses to rooms one at a time. The first course is assigned the room 1. Now suppose that k courses have been assigned their rooms. We assign the course $k + 1$ as follows: assign it to the least numbered room that is available at the start time of the lecture. It is clear that a course is assigned a *new* room only if there are overlapping lectures for all smaller-numbered rooms. Alternatively, the intuition is as follows. When an interval begins, we already know every interval that could possibly conflict with it. We get a new room only if it is necessary. Thus, our greedy algorithm achieves the minimum number of rooms assigned.

Example 6.3 Given the intervals $I_1 = [1, 4)$, $I_2 = [2, 5)$, $I_3 = [6, 7)$, $I_4 = [3, 8)$, $I_5 = [9, 10)$, sorted by start time they become $[1, 4)$, $[2, 5)$, $[3, 8)$, $[6, 7)$, $[9, 10)$. The greedy assignment is:



The minimum number of rooms required is equal to the maximum number of overlapping intervals, which is 3.

Theorem 6.4 (Interval Partitioning optimality) *For the Interval Partitioning Problem, assigning intervals to the first available room in order of their start times produces an optimal solution, that is, it minimizes the number of rooms required.*

Proof: The greedy algorithm sorts the intervals by start times and assigns each interval to the first available room; if no room is available, it opens a new one. Let G be the number of rooms used by the algorithm, and let O be the minimum number of rooms required in any optimal solution. Let $d = \max_t D(t)$ denote the maximum number of intervals that overlap at any single time.

Lower bound: $O \geq d$. At any time t with $D(t) = d$, those d intervals pairwise overlap and must lie in d distinct rooms. Hence any valid assignment, including the optimum, uses at least d rooms, so $O \geq d$.

Upper bound: $G \leq d$. The algorithm opens a new room only when every existing room already holds an interval overlapping the current one. Suppose the algorithm opens its k -th room for interval I at time

$t = s_I$. Then each of the $k - 1$ previously opened rooms currently hosts an interval that has not yet finished, i.e., overlaps I at time t ; together with I , that gives k pairwise-overlapping intervals at time t . Therefore $k \leq D(t) \leq d$. Since this holds for the last room opened, $G \leq d$.

Combining the two bounds, $G \leq d \leq O$. Because $G \geq O$ trivially (any valid solution uses at least the optimum), we conclude $G = O = d$, and the greedy algorithm is optimal. ■

Algorithm [IntervalPartition](#) is a greedy algorithm for the Interval Partition problem.

Algorithm IntervalPartition: Sequential Interval Partition

Input: s : array[1.. n] of int // Start times (sorted: $s[1] \leq s[2] \leq \dots \leq s[n]$)

Input: f : array[1.. n] of int // Finish times

Output: $room$: array[1.. n] of int // Room assigned to each interval

Var: Q : min-priority queue of $(endTime, roomNumber)$; int $numRooms := 0$

```

1 for int i := 1 to n do
2   if Q is not empty and Q.min().endTime ≤ s[i] then
3     | r := Q.extract-min().roomNumber;           // Reuse earliest-free room
4   else
5     | numRooms := numRooms + 1;                 // Allocate a new room
6     | r := numRooms
7   end
8   room[i] := r;
9   Q.insert(f[i], r);                           // Room r busy until f[i]
10 end
11 return room, numRooms

```

Now, let us determine the time complexity of our algorithm. The sorting of intervals based on the start times can be done in $O(n \log n)$ time. Suppose that we keep *available* rooms in a heap ordered by the $(endTime, roomnumber)$. Whenever we need a room for a lecture, we simply remove the minimum available in the heap. This heap operation takes $O(\log n)$ time. Thus, we take $O(n \log n)$ time for the entire algorithm.

6.4 Minimizing Maximum Lateness of Jobs

Suppose that we have n jobs such that each job j takes time t_j to execute and should ideally be completed before its deadline d_j . Our task is to schedule these jobs on a single processor. Suppose that the job j is started at time s_j . Then, the job ends at time $s_j + t_j$. If $s_j + t_j$ is less than or equal to the deadline d_j , then this job is not late. Otherwise, this job has *lateness* equal to $s_j + t_j - d_j$. Our task is to schedule these jobs so that the maximum lateness is minimized.

Jobs with earlier deadlines are more urgent because delaying them risks incurring large lateness that cannot be corrected later. By scheduling jobs in increasing order of deadlines, we ensure that the most time-sensitive jobs are completed as early as possible. Delaying a job with a later deadline is less risky, since

it has more slack. Thus, prioritizing earliest deadlines balances the schedule to minimize the worst-case lateness.

Let us start with a sequential algorithm. Should we consider these jobs in the order of their processing times, their *slack* times ($d_j - t_j$), or their deadlines d_j ? If a job has an earlier deadline, delaying it is always more dangerous than delaying a job with a later deadline. Thus, for this problem, we should consider jobs in the order of their deadlines.

Algorithm MinMaxLateness: A Sequential Program for the Minimizing Maximum Lateness problem

Input: t : array[1... n] of int; // processing times, sorted by deadline

Input: d : array[1... n] of int; // deadlines in non-decreasing order

Output: G : array[1... n] of int initially 0; // start time of each job

```

1 int last := 0;
2 int lateness := 0;
3 for int i := 1 to n do
4   | int finish := last + t[i];
5   | G[i] := last;
6   | last := finish;
7   | if (finish > d[i]) then lateness := max(lateness, finish - d[i]);
8 end

```

Algorithm [MinMaxLateness](#) processes jobs in the order of increasing deadlines. At each step, $last$ holds the current end-of-schedule time, $G[i]$ records the starting time of job i , and $finish$ is the moment it completes. The loop performs $O(1)$ work per job, so the algorithm runs in $O(n)$ time once the jobs are sorted by deadline. If the jobs are not pre-sorted, the initial sort dominates and the total time is $O(n \log n)$. The algorithm uses $O(n)$ space for the output array G .

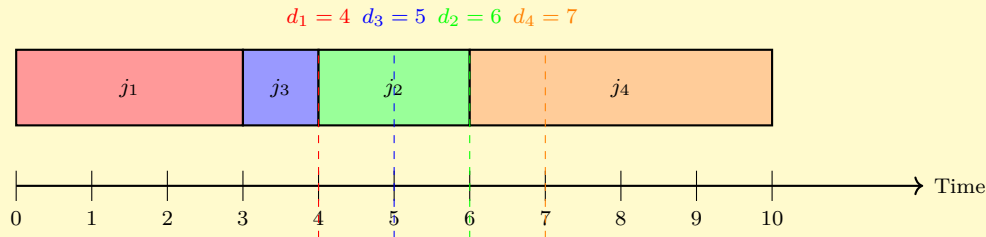
Example 6.5 Consider the following example.

Job j_i	Processing Time t_i	Deadline d_i
j_1	3	4
j_2	2	6
j_3	1	5
j_4	4	7

Sorting by deadline gives the order j_1, j_3, j_2, j_4 . Tracing the algorithm row by row:

i	job	$t[i]$	$d[i]$	$G[i]$ (= last on entry)	finish	lateness after
1	j_1	3	4	0	3	0
2	j_3	1	5	3	4	0
3	j_2	2	6	4	6	0
4	j_4	4	7	6	10	3

Only job j_4 runs past its deadline, contributing lateness $10 - 7 = 3$. The resulting schedule using Earliest Deadline First (EDF) is shown next.



The job j_2 finishes at time 6 and has a deadline of 6. The job j_4 finishes at time 10 but has a deadline of 7. Using earliest deadline first, this schedule minimizes *maximum lateness*, which is 3 units.

Theorem 6.6 (EDF optimality) *For the problem of scheduling n jobs with known processing times and deadlines on a single machine to minimize the maximum lateness, the Earliest Deadline First (EDF) algorithm produces an optimal solution.*

Proof:

The EDF algorithm schedules jobs in increasing order of their deadlines. Let G be the schedule produced by the EDF algorithm, and let O be an optimal schedule that minimizes maximum lateness. Suppose for contradiction that O is different from G and has a different order of jobs while achieving a smaller maximum lateness.

Consider the first position in which O and G differ. Let job j be scheduled earlier in O than job i , while G places job i before job j . Since EDF sorts jobs by increasing deadlines, we have $d_i \leq d_j$. Since both schedules are contiguous, swapping i and j in O does not delay the completion time of any job before i , nor does it improve the lateness of any job. Instead, it ensures that the job with an earlier deadline finishes sooner, which prevents an increase in lateness. By iterative application of this argument, we can transform O into G without increasing the maximum lateness. Since O was assumed to be optimal, it follows that G is also optimal. Therefore, EDF produces an optimal schedule.



6.5 Huffman Tree

The goal of the Huffman tree problem is to find an efficient encoding of a set of symbols. We are given n symbols. For each symbol i , the value $p[i]$ gives the probability that it appears in the given text, with $\sum_i p[i] = 1$. Our goal is to design a *prefix* code such that the given text can be translated to a binary string from which the original text can be recovered. We will create a binary tree so that the leaves correspond to the symbols and the path from the root to the leaf corresponds to the *code* for the symbol. Note that only the leaves of the tree correspond to symbols. Also note that leaves may be at different levels, and therefore this method returns a variable-length code for symbols rather than a fixed-length code.

In an optimal prefix code, symbols that occur more frequently should have shorter codewords, while rare symbols can tolerate longer ones. The greedy strategy builds the code tree from the bottom up by repeatedly merging the two least frequent symbols. This ensures that the least important symbols are pushed deeper in the tree, where longer codewords are assigned. By making the locally optimal choice of combining the least probable symbols at each step, we construct a globally optimal encoding.

Given n symbols with their probabilities of occurrence, the Huffman coding algorithm constructs an optimal prefix code using the following greedy approach. We first insert all symbols into a min-priority queue keyed by probability. As long as the queue contains more than one node, we remove the two nodes with the smallest probabilities, merge them into a new node whose probability is the sum of the two, and insert this merged node back into the queue. When only one node remains, it is the root of the Huffman tree. Finally, we obtain the code for each symbol by traversing the tree from the root, labelling the left branch with 0 and the right branch with 1.

Algorithm HuffmanCoding: Huffman Coding

Input: Set of symbols $S = \{s_1, s_2, \dots, s_n\}$ and frequencies $f(s_1), f(s_2), \dots, f(s_n)$

Output: Huffman tree T

```

1  $Q :=$  priority queue (min-heap) of nodes initially null ;
2 for each  $s_i \in S$  do
3   | Create a leaf node  $n_i$  with frequency  $f(s_i)$  and symbol  $s_i$ ;
4   | Insert  $n_i$  into  $Q$ ;
5 end
6 for  $i = 1$  to  $n - 1$  do
7   |  $x :=$  extract-min( $Q$ ) ; // Remove node with smallest frequency
8   |  $y :=$  extract-min( $Q$ ) ; // Remove next smallest
9   |  $z :=$  new internal node with children  $x$  and  $y$ ;
10  |  $f(z) := f(x) + f(y)$  ; // Sum frequencies
11  | Insert  $z$  into  $Q$ ;
12 end
13  $T :=$  extract-min( $Q$ ) ; // Final tree (root)
14 return  $T$ 
```

Example 6.7 Consider six symbols with frequencies that exercise merges of internal nodes with each other:

Symbol	A	B	C	D	E	F
Frequency	45	13	12	16	9	5

The priority queue evolves as follows, with frequencies shown in non-decreasing order at each step and internal nodes denoted by the sum of their leaves in parentheses:

Step	Extracted (smallest two)	Queue after insertion
0	—	5, 9, 12, 13, 16, 45
1	5, 9 → 14	12, 13, 14, 16, 45
2	12, 13 → 25	14, 16, 25, 45
3	14, 16 → 30	25, 30, 45
4	25, 30 → 55	45, 55
5	45, 55 → 100	100

The resulting Huffman codes are:

Symbol	Frequency	Huffman Code
A	45	0
C	12	100
B	13	101
F	5	1100
E	9	1101
D	16	111

The total frequency is $45 + 13 + 12 + 16 + 9 + 5 = 100$, so the average code length is $(45 \cdot 1 + (12 + 13) \cdot 3 + (5 + 9) \cdot 4 + 16 \cdot 3) / 100 = 224 / 100 = 2.24$ bits per symbol, compared with 3 bits per symbol for a fixed-length code, a saving of about 25%.

The time and space complexity of the Huffman coding algorithm depend on the use of a min-heap for the priority queue Q . For the time complexity, building Q from the n leaf nodes costs $O(n \log n)$ since each insertion into a heap of size at most n is $O(\log n)$. The main loop runs $n - 1$ times, and each iteration performs two extract-min operations and one insertion, all at cost $O(\log n)$; the loop therefore contributes $O(n \log n)$. The final extract-min is $O(\log n)$, which is absorbed, giving an overall running time of $O(n \log n)$. For the space complexity, the priority queue holds at most n nodes at any time, and the constructed tree has n leaves and $n - 1$ internal nodes for a total of $2n - 1$ nodes; both contribute $O(n)$, so the algorithm uses $O(n)$ space, where n is the number of symbols.

Theorem 6.8 (Huffman optimality) *Algorithm [HuffmanCoding](#) constructs a prefix code of minimum expected length $\sum_i p_i \ell_i$, where ℓ_i is the length of the codeword for symbol i .*

Proof: We prove the claim by induction on n , using an exchange argument at each step. For $n = 2$, any prefix code has $\ell_1 = \ell_2 = 1$ and Huffman produces exactly this code, which is trivially optimal.

For $n > 2$, let u and v be the two symbols with the smallest probabilities, and let z be a fresh symbol with $p_z = p_u + p_v$. Consider the reduced alphabet $C' = (C \setminus \{u, v\}) \cup \{z\}$ on $n - 1$ symbols.

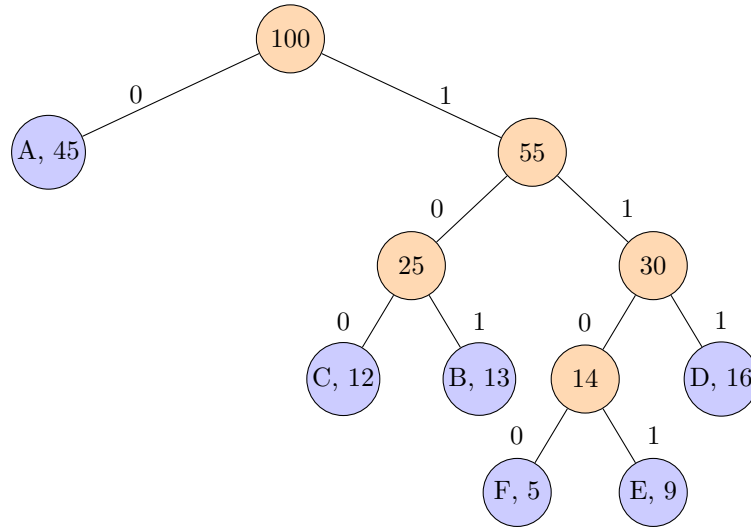


Figure 6.1: Huffman tree for symbols A, B, C, D, E, F with frequencies 45, 13, 12, 16, 9, 5. Leaves (blue) carry the source symbols; internal nodes (orange) carry the summed frequencies created by merges.

Greedy-choice property. There is an optimal tree T^* for C in which u and v are sibling leaves at the deepest level. Take any optimal tree T ; since T is full (otherwise we could shorten a codeword), it has two sibling leaves a, b at the deepest level. Swap $u \leftrightarrow a$ and $v \leftrightarrow b$. The cost change is $(p_a - p_u)(\ell_u^{\text{old}} - \ell_a^{\text{old}}) + (p_b - p_v)(\ell_v^{\text{old}} - \ell_b^{\text{old}}) \leq 0$ because $p_u \leq p_a$, $p_v \leq p_b$, and the depth differences are non-negative. The resulting tree is still optimal and has u, v as deepest siblings.

Optimal-substructure property. Let T' be the tree for C' obtained from T^* by replacing the internal node that is parent of u, v with a leaf z . Conversely, given any tree S' for C' , we can form a tree S for C by expanding the leaf z into an internal node with children u, v . In both directions, $\text{cost}(S) = \text{cost}(S') + p_u + p_v$. Hence T' is optimal for C' .

By the inductive hypothesis, Huffman's algorithm produces an optimal tree for C' in its remaining $n-2$ merges. Expanding z back into u, v yields an optimal tree for C , which is exactly what Huffman produces on the full input. Therefore Huffman is optimal. ■

6.6 Fractional Knapsack

The knapsack problem is a natural test case for the reach of greedy methods. We are given n items with values v_1, \dots, v_n and weights w_1, \dots, w_n , and a knapsack capacity W . In the *0/1 knapsack problem*, we must choose a subset $S \subseteq [n]$ with $\sum_{i \in S} w_i \leq W$ that maximises $\sum_{i \in S} v_i$; here no fixed greedy rule is optimal. For capacity $W = 5$ and items $\{(v=10, w=5), (v=7, w=3), (v=6, w=2)\}$, the rule “take the largest value first” picks item 1 alone, for value 10, while items 2 and 3 together give 13. The 0/1 knapsack problem needs dynamic programming (Chapter 10) and in general is NP-hard.

In the *fractional knapsack problem*, items are divisible: we may take a fraction $x_i \in [0, 1]$ of each item, paying weight $x_i w_i$ and earning value $x_i v_i$. The capacity constraint becomes $\sum_i x_i w_i \leq W$. Divisibility restores the power of greed.

To maximize value within a limited capacity, we should prioritize items that provide the greatest value per unit weight. Sorting items by their value-to-weight ratio ensures that each unit of capacity is used as efficiently as possible. We take as much as possible of the highest-ratio item before moving to the next. Since items are divisible, any deviation from this rule would replace higher-value density with lower-value density, leading to a worse total value.

Algorithm FractionalKnapsack: Greedy algorithm for the Fractional Knapsack problem

Input: $v[1..n]$, $w[1..n]$: reals; W : real capacity
Output: $x[1..n]$: fractional selections, initially 0

```

1 Sort items in non-increasing order of the ratio  $v[i]/w[i]$ ;
2 int  $rem := W$ ;
3 for  $i := 1$  to  $n$  do
4   if  $w[i] \leq rem$  then
5     |  $x[i] := 1$ ;
6     |  $rem := rem - w[i]$ ;
7   else
8     |  $x[i] := rem/w[i]$ ;           // take the remaining fraction
9     | break;
10  end
11 end

```

Theorem 6.9 (Fractional Knapsack optimality) *Algorithm [FractionalKnapsack](#) returns a selection of maximum total value for the fractional knapsack problem.*

Proof: We use an exchange argument. Assume items are indexed so that $v_1/w_1 \geq v_2/w_2 \geq \dots \geq v_n/w_n$, and let G be the greedy solution and O any optimal solution; without loss of generality both pack total weight exactly W (if O is short, augment it with any available item, which cannot decrease its value).

Let j be the smallest index with $O_j < G_j$. Because both solutions pack the same total weight, there must exist $j' > j$ with $O_{j'} > G_{j'}$. Shift $\epsilon > 0$ units of weight from item j' to item j : set $O_j \leftarrow O_j + \epsilon/w_j$ and $O_{j'} \leftarrow O_{j'} - \epsilon/w_{j'}$, choosing ϵ small enough that both fractions stay in $[0, 1]$. The change in total value is

$$\Delta = \epsilon \left(\frac{v_j}{w_j} - \frac{v_{j'}}{w_{j'}} \right) \geq 0,$$

since $j < j'$ implies $v_j/w_j \geq v_{j'}/w_{j'}$. The swapped solution is still feasible and no worse. Iterating eliminates all disagreements without reducing the value, turning O into G . Hence G is optimal. ■

Example 6.10 Consider four items and a knapsack of capacity $W = 10$:

Item i	Value v_i	Weight w_i	Ratio v_i/w_i
1	10	2	5
2	12	3	4
3	6	2	3
4	8	4	2

The items are already sorted by non-increasing value-per-weight ratio. Tracing the algorithm:

i	w_i	rem (before)	x_i	Value added
1	2	10	1	10
2	3	8	1	12
3	2	5	1	6
4	4	3	3/4	6

The greedy algorithm takes items 1, 2, and 3 in full and three-quarters of item 4. The total weight used is $2 + 3 + 2 + 3 = 10 = W$, and the total value is $10 + 12 + 6 + 6 = 34$.

For contrast, refusing to break item 4 and taking items 1, 2, 3 alone gives value 28; taking item 4 whole instead of item 3 yields weight $2 + 3 + 4 = 9$ and value $10 + 12 + 8 = 30$. Both alternatives are worse, confirming the optimality of the greedy fractional selection on this instance.

The sort dominates the running time: $O(n \log n)$ sequentially, with $O(n)$ additional work for the loop. In parbook, observe that the prefix sums of w_i along the sorted order determine, in parallel, the cut-off index k where the knapsack becomes full; a parallel prefix-sum computes them in $O(\log n)$ time and $O(n)$ work, and the fractional coefficient x_k is then read off in $O(1)$ time. The total parallel time is $O(\log n)$ after the sort, matching the bound for Minimizing Maximum Lateness.

6.7 LLP Perspective

In this section, we provide LLP versions of greedy algorithms.

LLP: Interval Scheduling Algorithm

We are given n jobs sorted by their finish times. Jobs with the same finish times are sorted according to their start times. We are interested in finding a maximum-sized subset of $[n]$ such that each pair of jobs is mutually compatible. We define B on a subset of jobs to be true if the subset is a maximum-sized subset of mutually compatible jobs. For example, consider the set of jobs as $\{[1, 3), [2, 4), [5, 8), [6, 9)\}$. It can be verified that there is no subset of size three that has mutually compatible jobs. However, there are multiple subsets of size two that are mutually compatible. For example, $\{[1, 3), [5, 8)\}$ is one such subset. Another example is $\{[2, 4), [5, 8)\}$. The predicate B as defined is not lattice-linear in the lattice of all subsets. It is not closed under the meet operation. The meet of the two examples is the subset $\{[5, 8)\}$, which is not a maximum-sized subset that has mutually compatible jobs. We redefine B to be the lexicographically least maximum-sized subset of mutually compatible jobs. Now, simply by the definition, B is true exactly on one subset. Hence, B is lattice-linear.

We determine the lexicographically least maximum-sized subset of mutually compatible jobs as follows. We maintain a linked list of all the jobs under consideration. These jobs are in the order of their finish times. The algorithm will maintain a Boolean $G[j]$ for each job j . The variable $G[j]$ is initially false and is set to true only if it is guaranteed to be in the maximal set of jobs that the algorithm chooses. We call a job j *fixed* if $G[j]$ is true. The *init* command always fixes the first job. We now make the following observations.

1. If any job j has its start time greater than or equal to the finish time of the previous job, then that job is always included in the final set. Thus, the job j can be fixed.
2. For any job, if the previous job in the linked list is fixed and the start time of the current job is less than the finish time of the previous job, then this job can be deleted.

It follows from the above two rules that eventually all jobs will be fixed or deleted. The algorithm terminates when all jobs are fixed. The set of all fixed jobs is a maximum-sized set of jobs that can be completed. Furthermore, it is the lexicographically least such set.

The LLP algorithm for the activity selection problem is shown in Algorithm [LLP-IntervalScheduling](#).

Algorithm LLP-IntervalScheduling: LLP Program for the Interval Scheduling problem

```
// Assume that all jobs are in a doubly linked list
// The variable prev denotes the previous node in the linked list
Input:  $s[j]$ : int,  $f[j]$ : int
Output:  $G[j]$ : boolean, initially if  $(j = 1)$  then  $G[j] := true$  else  $G[j] := false$ 
1 forbidden:  $\neg G[j] \wedge f[prev] \leq s[j]$ ;
2 advance:  $G[j] := true$ ;
3 forbidden:  $(j > 1) \wedge G[prev] \wedge s[j] < f[prev]$ ;
4 advance: delete this node from the linked list;
```

For example, consider the following four jobs: $job_1 = [1, 4)$, $job_2 = [2, 5)$, $job_3 = [4, 5)$, $job_4 = [5, 7)$. The first job gets fixed because the first job is always fixed. The fourth job also gets fixed because its starting point is at least as large as the finishing time of the previous job. Since the first job is fixed and the second job starts before the first job is finished, it is deleted from the list. Now, the third job is also fixed because it starts after the first job is finished. Each job in the linked list is either fixed by the first advance rule or deleted by the second, and both actions are $O(1)$ because the list pointers let us reach *prev* directly. Every job is touched at most twice (once to be fixed, possibly once to be deleted), so the main loop runs in $O(n)$ time once the jobs are sorted by finish time; the overall cost is therefore $O(n \log n)$ including the sort, with $O(n)$ space for the list.

LLP: Interval Partition Algorithm

We now explore an LLP version of the algorithm for Interval Partition. Let $G[i]$ denote the room assigned to course i (where courses are sorted based on start times). We initialize $G[i]$ to 1 for all i . This room assignment is feasible iff there are no overlapping intervals. We now define the feasibility of the room assignment for any set of intervals. A room assignment is feasible if for any pair of intervals i, j whenever they overlap, they are in different rooms. Formally, let B be defined as

$$\forall i, j \in [n] : (i \neq j) \wedge (s_j < f_i) \wedge (s_i < f_j) \Rightarrow G[i] \neq G[j]$$

It can again be observed that B is not lattice-linear. For example, if there are only two overlapping activities, then both G vectors, $[1, 2]$ and $[2, 1]$, satisfy B . However, their minimum $[1, 1]$ does not satisfy B . We apply the same trick as in the previous problem. Instead of finding any feasible room assignment, we find the lexicographically least room assignment. Thus, B is redefined as

$$\forall j : G[j] = \min\{r \mid \forall i < j : (s[j] < f[i]) \Rightarrow (G[i] \neq r)\}$$

Thus, $G[j]$ is assigned the least numbered free room available when the course j starts. With this new definition, it is clear that B holds only for one possible G .

The predicate B may not be true for the initial assignment when there are overlapping intervals. Our interest is in finding the least assignment to the rooms that makes the predicate true. We consider the lattice of n -dimensional vectors of natural numbers. The bottom element of this lattice is the vector with all the ones. We use $fixed[i]$ to denote that the course i has been assigned a room that will not be changed. Let $pre(i)$ denote all courses numbered less than i that start prior to course i or at the same time as course i and finish later than s_i .

We say that index j is forbidden if all courses in $pre(j)$ are $fixed$ and $\neg fixed[j]$. Whenever index j is forbidden, it can be advanced as follows. Let r be the least room that has not been assigned to any overlapping course prior to j . Then $G[j]$ is set to r and $fixed[j]$ is set to true.

Algorithm LLP-IntervalPartition: An LLP Program for the Interval Partition Problem

Input: $s[j]$: int, $f[j]$: int; $pre[j]$: set of $1..n$ initially

$$pre[j] = \{i \mid (i < j) \wedge (s[i] \leq s[j]) \wedge (s[j] < f[i])\}$$

Output: $G[j]$: int initially 1; $fixed[j]$: boolean initially false

- 1 **forbidden:** $(\forall i \in pre(j) : fixed(i)) \wedge \neg fixed(j)$;
 - 2 **advance:** $G[j] := \min_{k \in pre(j)} \{r \mid r \neq G[k]\}$; $fixed[j] := true$;
-

As before, we assume that the intervals are provided to us, sorted by their start times. Each index j keeps track of the number of intervals in $pre(j)$ that have been fixed, so checking whether j is forbidden is $O(1)$. Finding the least free room in the advance step uses a heap keyed by the occupied rooms at time $s[j]$; each heap operation costs $O(\log n)$. There are exactly n advances, one per course, so the main loop contributes $O(n \log n)$ time, which dominates the initial sort and gives an overall sequential running time of $O(n \log n)$. The space complexity is $O(n)$: the arrays G and $fixed$ use $O(n)$ each, and the heap stores at most n rooms.

LLP: Minimizing Maximum Lateness of Jobs

We now consider the LLP version of computing a job schedule that minimizes the maximum lateness of jobs. The greedy approach is to schedule jobs in order of increasing deadline. For simplicity, assume that all deadlines are distinct and that the jobs are indexed so that $d[1] < d[2] < \dots < d[n]$.

Let $G[j]$ denote the starting time of job j , initially 0. The lattice is \mathbb{N}^n with componentwise order, and the bottom element is the all-zeros vector. The target vector G^* is characterised by

$$B \equiv \forall j : G[j] = \sum_{i < j} t[i].$$

Because G^* is uniquely determined, B is lattice-linear on its own.

We take the forbidden condition to be the direct value comparison

$$\text{Forbidden}(j) \equiv G[j] < \sum_{i < j} t[i],$$

and the advance step simply raises $G[j]$ to that target value. Initially every $j \geq 2$ is forbidden (since the right-hand side is positive while $G[j] = 0$), so all $n - 1$ indices can advance in the same parallel round.

Algorithm LLP-MinMaxLateness: LLP Program for the Minimizing Maximum Lateness problem

Input: $t[j]$: int, processing time of job j ; $d[j]$: int, deadline of job j (jobs indexed in non-decreasing deadline order)

Output: $G[j]$: int, initially 0, the starting time of job j ; *lateness*: int, initially 0

- 1 **forbidden**(j): $G[j] < \sum_{i < j} t[i]$;
 - 2 **advance**: $G[j] := \sum_{i < j} t[i]$;
 - 3 *lateness* := $\max_j \max(0, G[j] + t[j] - d[j])$;
-

Example 6.11 Continuing Example 6.5, we trace MINMAXLATELLP on the same four jobs with processing times $t = (3, 2, 1, 4)$ and deadlines $d = (4, 6, 5, 7)$. Reindex them in deadline order: $j_1 \rightsquigarrow 1$, $j_3 \rightsquigarrow 2$, $j_2 \rightsquigarrow 3$, $j_4 \rightsquigarrow 4$, so the deadline-sorted processing times are $(3, 1, 2, 4)$ and deadlines are $(4, 5, 6, 7)$. The targets $\sum_{i < j} t[i]$ are $(0, 3, 4, 6)$. Initially $G = (0, 0, 0, 0)$ and indices 2, 3, 4 are all forbidden. One parallel round advances $G[2] := 3$, $G[3] := 4$, $G[4] := 6$ simultaneously, after which no index is forbidden. The final *lateness* reduction reads $G + t - d = (-1, -1, -2, 3)$, giving *lateness* = 3, attained by j_4 .

round	forbidden set	advances applied	G after
0	$\{2, 3, 4\}$	—	$(0, 0, 0, 0)$
1	\emptyset	$G[2] := 3, G[3] := 4, G[4] := 6$ (parallel)	$(0, 3, 4, 6)$

This matches the schedule drawn in Example 6.5: the jobs start at times 0, 3, 4, 6 in deadline order, and the maximum lateness is 3 units.

Sequentially, computing each target $\sum_{i < j} t[i]$ takes $O(n)$ via a left-to-right scan, and the lateness reduction is another $O(n)$, so the total time is $O(n)$ once the deadlines are sorted and $O(n \log n)$ otherwise. The arrays G and the scalar *lateness* use $O(n)$ space.

LLP: Fractional Knapsack

We now give an LLP version of the fractional knapsack algorithm. Assume the items are indexed in non-increasing order of v_j/w_j . Let $G[j] \in [0, 1]$ denote the fraction of item j included in the knapsack, initially 0. The lattice is $[0, 1]^n$ with componentwise order and bottom the all-zeros vector.

Let $S_j = \sum_{k \leq j} w[k]$ denote the cumulative weight up through item j , and define the target fraction

$$G^*[j] = \begin{cases} 1 & \text{if } S_j \leq W, \\ (W - S_{j-1})/w[j] & \text{if } S_{j-1} < W < S_j, \\ 0 & \text{if } S_{j-1} \geq W. \end{cases}$$

The predicate $B \equiv (\forall j : G[j] = G^*[j])$ is satisfied by a unique vector G^* and is therefore lattice-linear on its own. We take the forbidden condition to be the direct value comparison

$$\text{forbidden}(j) \equiv G[j] < G^*[j],$$

and the advance step raises $G[j]$ to $G^*[j]$. Initially every j with $G^*[j] > 0$ is forbidden, so all such indices can advance in a single parallel round.

Algorithm LLP-FractionalKnapsack: LLP Program for Fractional Knapsack

Input: $v[1..n]$, $w[1..n]$: reals (sorted so that $v[j]/w[j]$ is non-increasing); W : capacity

Output: $G[j]$: real in $[0, 1]$, initially 0

// Let $S_j = \sum_{k \leq j} w[k]$ and $G^*[j]$ be the target value defined above.

1 **forbidden**(j): $G[j] < G^*[j]$;

2 **advance**: $G[j] := G^*[j]$;

Sequentially, computing the prefix sums S_j takes $O(n)$ time once the items are sorted, and each target $G^*[j]$ is obtained in $O(1)$ from S_{j-1}, S_j . The total running time is therefore $O(n \log n)$ (dominated by the sort), with $O(n)$ space.

6.8 When Greedy Fails

The five problems we have studied share a feature that makes the greedy approach succeed: each one has an *exchange property* strong enough to carry the correctness proof. Once we identify the “right” local rule (earliest finish time, earliest available room, earliest deadline, merge the two rarest symbols, highest value-per-weight), any solution that deviates from it can be transformed into the greedy solution without losing value — that is the content of the exchange argument.

Greedy algorithms rely on a structural property that allows local choices to be extended to global optimal solutions. When this structure is absent, a locally optimal decision may block better global configurations. For example, in the 0/1 knapsack problem, choosing items with the highest value or value-to-weight ratio can prevent selecting a combination of items with higher total value. In such cases, the lack of a suitable exchange property means that greedy decisions cannot be safely corrected, and more powerful techniques such as dynamic programming are required.

Many closely related problems lack this property, and for them greedy fails. Replacing divisibility of items with indivisibility turns Fractional Knapsack into the 0/1 Knapsack problem, which is NP-hard and has no known greedy rule that guarantees an optimum: the counterexample in the introduction (capacity $W = 5$ with items $(v, w) \in \{(10, 5), (7, 3), (6, 2)\}$) shows that the “highest-value first” rule gives 10 while an optimal pair gives 13. Replacing compatibility with a weight on activities turns Interval Scheduling into Weighted Interval Scheduling, for which no fixed greedy rule is optimal and dynamic programming is

required. Making the tasks non-preemptive on multiple machines changes the scheduling landscape in a way that local-choice heuristics can only approximate. Even the unifying language of *matroid theory* — which formalises when the greedy-choice property provably works — captures only some of the problems where greedy succeeds; Huffman coding, for instance, is not a matroid problem but is still solved greedily thanks to its own exchange structure.

The cautionary tale, then, is that greedy is a consequence of structure, not a method of last resort: when the problem admits a sound exchange argument, greedy produces an exact optimum in low polynomial time; when it does not, even the most intuitive local rule can be arbitrarily far from optimal, and a richer technique such as dynamic programming (Chapter 10) or approximation is needed.

6.9 Summary

The greedy strategy for *Interval Scheduling* prioritizes activities with the earliest finish time to maximize the number of non-overlapping intervals, ensuring that more activities can be accommodated. In *Interval Partitioning*, assigning each interval to the earliest available room ensures minimal room usage by efficiently tracking overlapping intervals. For *Minimizing Maximum Lateness*, scheduling jobs by their earliest deadline prevents unnecessary delays, minimizing the worst-case lateness across all jobs. For *Huffman Tree Construction*, merging the least probable symbols first ensures that frequent symbols receive shorter codes, leading to an optimal encoding that minimizes the total cost of transmission. Finally, for the *Fractional Knapsack Problem*, selecting items by highest value-to-weight ratio fills the knapsack with the most valuable mix per unit weight, yielding the maximum achievable total value.

Table 6.1 summarizes all five problems considered in this chapter. In the table, n denotes the number of intervals (for interval scheduling and interval partitioning), the number of jobs (for minimizing lateness), the number of symbols (for Huffman coding), or the number of items (for fractional knapsack). The time complexity includes the time to sort the input as required by the strategy.

Problem	Strategy	Time Complexity
Interval Scheduling	Earliest finish time	$O(n \log n)$
Interval Partitioning	Earliest available room	$O(n \log n)$
Minimizing Lateness	Earliest deadline	$O(n \log n)$
Huffman Tree	Merge smallest probabilities	$O(n \log n)$
Fractional Knapsack	Highest value/weight ratio	$O(n \log n)$

Table 6.1: Summary of greedy algorithms.

6.10 Problems

1. A railway station needs to schedule maintenance for the incoming trains before their next departure. Each train has a maintenance duration and a deadline. Find the optimal order of maintenance to minimize the worst delay.
2. Given n intervals and an integer k , find the maximum subset of intervals such that at most k intervals overlap at any time.

3. Each task has a required start time and finish time, but there must be a mandatory idle gap of at least g units between consecutive tasks in the same room. Find the minimum number of rooms required.
4. Each task has a start and finish time, but also requires a *specific type of machine*. Different machine types may be limited in number. Find the minimum number of machines needed.
5. Prove that the Huffman coding algorithm produces a prefix code of minimum expected length. State the key greedy-choice and optimal-substructure properties used in the proof.
6. Given n closed intervals on the real line, find the minimum number of points required so that every interval contains at least one chosen point. Give a greedy algorithm, prove its correctness, and state its time complexity.

6.11 Bibliographic Remarks

This chapter on greedy algorithms covers several classic problems: Interval Scheduling, Interval Partitioning, Minimizing Lateness, Huffman Tree construction, and Fractional Knapsack. The Interval Scheduling Problem and Interval Partitioning Problem are covered in [CLRS09] and [KT06b]. For the Minimizing Lateness problem, scheduling jobs by earliest deadline (EDF) to minimize maximum lateness is a well-known greedy solution, as shown in Algorithm [MinMaxLateness](#). The optimality of this algorithm, proven via an exchange argument, is a standard result in scheduling theory. [LRK76] provides one of the first rigorous treatments of this problem, establishing the effectiveness of the EDF rule for scheduling a single machine with deadlines. The Huffman tree construction, which builds an optimal prefix code by greedily merging the least probable symbols, is a cornerstone of data compression. Introduced by [Huf52], this algorithm's greedy nature and optimality for variable-length coding are foundational to information theory. Our sequential description mirrors the classic priority-queue implementation in $O(n \log n)$ time, as detailed in [CLRS09]. The fractional knapsack problem and the contrast with the NP-hard 0/1 variant are treated in [CLRS09, KT06b].

Chapter 7

The Shortest Path Problem

7.1 Introduction

The single source shortest path (SSSP) problem has wide applications in transportation, networking, route-planning, and many other fields. It asks, given a weighted directed graph with n vertices and m edges and a distinguished *source* vertex v_0 , for $cost[x]$: the minimum total weight of any directed path from v_0 to x , where the cost of a path is the sum of edge weights along it. A related and equally fundamental problem is the *all-pairs shortest-path (APSP) problem*, which asks for the matrix of shortest-path costs between every pair of vertices.

Three classical design techniques dominate this chapter. *Greedy* algorithms exploit the fact that when edge weights are non-negative, the next-smallest distance among undiscovered vertices is correct and can be fixed irrevocably — the idea that powers Dijkstra’s algorithm. *Dynamic programming* removes the non-negativity restriction by iterating a relaxation recurrence over all edges (BellmanFord) or over all intermediate-vertex sets (FloydWarshall). *Reweighting* combines the two: Johnson’s algorithm first runs BellmanFord to derive a potential function that makes every edge non-negative, then runs Dijkstra from every source on the reweighted graph to obtain all-pairs distances efficiently on sparse graphs.

All of these sequential algorithms admit reformulations in the *lattice-linear predicate (LLP)* framework introduced in Chapter 2. In an LLP formulation, a single generic algorithm — advance every forbidden index until the predicate B holds — subsumes Dijkstra’s single-fix rule, BellmanFord’s edge-relaxation loop, and FloydWarshall’s triple-nested update. The LLP reformulations expose parallelism (many forbidden indices can advance concurrently).

This chapter is organized as follows. Section 7.2 describes Dijkstra’s algorithm for SSSP on graphs with non-negative edge weights. Section 7.3 describes the BellmanFord algorithm, which lifts the non-negativity restriction and also detects negative-weight cycles. Section 7.4 presents the FloydWarshall algorithm for APSP in $\Theta(n^3)$ time via the intermediate-vertex recurrence. Section 7.5 develops Johnson’s algorithm, which uses the BellmanFord result to reweight edges and then applies Dijkstra from every source, yielding APSP in $O(mn + n^2 \log n)$ time — faster than FloydWarshall on sparse graphs.

The remaining sections reformulate these algorithms in the LLP framework. Section presents the LLP-BellmanFord algorithm for graphs that may have negative edges. Section gives the LLP version of Johnson’s algorithm, which uses an LLP pricing algorithm in place of BellmanFord to derive the reweighting

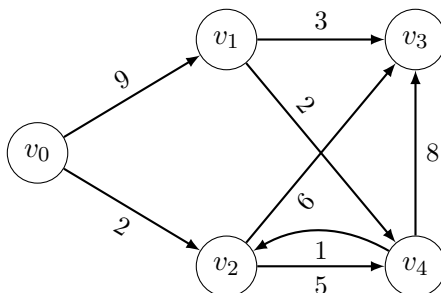


Figure 7.1: A Weighted Directed Graph

potentials. Finally, Section develops a matrix-multiplication-based algorithm for the transitive closure of a binary matrix and extends it to an LLP-FloydWarshall algorithm for all-pairs shortest paths with $O(\log^2 n)$ parallel time. The chapter closes with a summary, problems, and bibliographic remarks.

7.2 Dijkstra's Algorithm

We consider a directed weighted graph (V, E, w) where V is the set of vertices, E is the set of directed edges, and w is a map from the set of edges to positive reals (see Fig. 7.1 for a running example). To avoid trivialities, we assume that the graph is loop-free and that every vertex x , except the source vertex v_0 , has at least one incoming edge.

Algorithm Dijkstra: Finding the shortest costs from v_0 .

Input: Graph (V, E) with edge weights $w[j, k]$; source vertex v_0

Output: $dist[0..n-1]$: array of integer initially $\forall i : dist[i] = \infty$

```

1 fixed: array[0..n-1] of boolean initially  $\forall i : fixed[i] = false$ ;
2 H: binary heap of  $(j, d)$  initially empty ;           // j is the vertex and d is the cost
3  $dist[0] := 0$ ;
4 H.insert((0,  $dist[0]$ ));
5 while  $\neg H.empty()$  do
6    $(j, d) := H.removeMin()$ ;
7   if (fixed[j]) then continue;
8   fixed[j] := true;
9   forall k:  $\neg fixed[k] \wedge (j, k) \in E$  do
10    if ( $dist[k] > dist[j] + w[j, k]$ ) then
11       $dist[k] := dist[j] + w[j, k]$ ;
12      H.add (k,  $dist[k]$ );
13    end
14  end
15 end

```

Dijkstra's algorithm maintains $dist[i]$, which is a cost to reach v_i from v_0 . Each vertex x in the graph has initially $dist[x]$ equal to ∞ . Whenever a vertex is discovered for the first time, its $dist[x]$ becomes

less than ∞ . We use the predicate $discovered(x) \equiv dist[x] < \infty$. The variable $dist$ decreases for a vertex whenever a shorter path is found due to edge relaxation.

In addition to the variable $dist$, a Boolean array $fixed$ is maintained. Thus, every discovered vertex is either *fixed* or *non-fixed*. The invariant maintained by the algorithm is that if a vertex x is *fixed* then $dist[x]$ gives the final shortest cost from vertex v_0 to x . If x is *non-fixed*, then $dist[x]$ is the cost of the shortest path to x that only goes through fixed vertices.

When all edge weights are non-negative, distances can only increase as we extend a path. Thus, among all currently discovered vertices, the one with the smallest tentative distance must already have its final shortest-path value. Any alternative path to this vertex would have to go through another vertex whose distance is at least as large, and then incur an additional non-negative cost, making it no better. Therefore, we can safely fix the closest unfixed vertex at each step and propagate its distance to its neighbors, growing the set of vertices with known optimal distances in a greedy manner.

The algorithm uses a heap H that contains all vertices that have been discovered but are not fixed along with their distance estimates $dist$. We view the heap as consisting of tuples of the form $(j, dist[j])$ where the heap property is with respect to the $dist$ values. The algorithm has one main *while* loop that removes the vertex with the minimum distance from the heap with the method $H.removeMin()$, say v_j , and marks it as fixed. It then *explores* the vertex v_j by relaxing all its adjacent edges going to non-fixed vertices v_k . The value of $dist[k]$ is updated to the minimum of $dist[k]$ and $dist[j] + w[j, k]$. This step is called *edge relaxation*. If v_k is not on the heap, then it is inserted; else, if $dist[k]$ has decreased, then the label associated with the vertex k is inserted in the heap. We abstract this step as the method $H.add(k, dist[k])$. Since the vertex may already be on the heap, the insertion may cause a vertex to be present in a heap with different distances. Hence, when we remove a vertex from the heap, if it is already fixed, we simply go to the next vertex in the heap. The algorithm terminates when the heap is empty. At this point there are no discovered non-fixed vertices and $dist$ reflects the cost of the shortest path to all discovered vertices. If a vertex j is not discovered then $dist[j]$ is infinity reflecting that v_j is unreachable from v_0 .

Observe that every vertex goes through the following states. Every vertex x is initially *undiscovered* (that is, $dist[x] = \infty$). If x is reachable from the source vertex, then it is eventually *discovered* (i.e., $dist[x] < \infty$). A discovered vertex is initially *non-fixed*, and is therefore in the heap H . Whenever a vertex is removed from the heap it is a *fixed* vertex. A fixed vertex may either be *unexplored* or *explored*. Initially, a fixed vertex is unexplored. It is considered explored when all its outgoing edges have been relaxed.

The following lemma simply summarizes the well-known properties of Dijkstra's algorithm. We leave them as an exercise.

Lemma 7.1 *The outer loop in Dijkstra's algorithm satisfies the following invariants.*

- (a) *For all vertices x : $fixed[x] \Rightarrow (dist[x] = cost[x])$.*
- (b) *For all vertices x : $dist[x]$ is equal to the cost of the shortest path from v_0 to x such that all vertices in the path before x are fixed.*

Theorem 7.2 (Dijkstra correctness) *On a directed graph with non-negative edge weights, Algorithm [Dijkstra](#) terminates with $dist[x] = cost[x]$ for every vertex x , where $cost[x]$ is the length of the shortest path from v_0 to x (or ∞ if x is unreachable).*

Proof: By Lemma 7.1(a), every vertex marked *fixed* receives its correct shortest-path cost. It remains to show that every reachable vertex eventually becomes *fixed*. An extracted vertex is inserted into the heap

when a relaxed edge discovers it for the first time; since the heap is not drained until it is empty and every relaxation either inserts a new vertex or replaces an old key with a smaller one, every discovered vertex is eventually extracted and fixed. When the heap empties, by Lemma 7.1(b) every unreached vertex has $dist[x] = \infty$, which matches $cost[x]$.

The greedy invariant — that the minimum-key vertex on the heap is ready to be fixed — depends on non-negativity: if v has the smallest heap key d , no path through a not-yet-fixed vertex u can improve on d , because the sub-path through u already costs at least the heap key of u , which is $\geq d$, and all remaining edge weights are non-negative. ■

For complexity analysis, let n be the number of vertices and m be the number of edges. We analyze the version of Dijkstra's algorithm in which, instead of adjusting the key in the heap for a vertex, we simply insert the vertex in the heap. As a result, the heap may have a vertex multiple times with different keys. When a vertex is removed, we check if it has already been fixed. If it is fixed, then we do not need to explore it, and we can remove the next vertex in the heap. Since a vertex can be inserted in the heap only when an edge is relaxed, we find that there are at most m insertions in the heap. We can also conclude that there are at most m deletions from the heap. Since an insertion or deletion of a heap takes $O(\log m) = O(\log n)$ time, we get the overall time complexity of $O(m \log n)$.

Example 7.3 Here is a walk-through of Dijkstra's algorithm on the graph of Figure 7.1, starting from the vertex v_0 .

1. Set v_0 as the source vertex and assign it a distance of 0. Assign all other vertices a tentative distance of infinity. We insert the source vertex v_0 in the heap H .
2. Since the heap is not empty, we remove the vertex at the top of the heap. It is v_0 with a distance of 0. We mark that vertex as *fixed*. We examine its neighbors v_1 and v_2 . For $v_0 \rightarrow v_1$, the existing distance to v_1 is infinity. The distance 0 (distance to v_0) + 9 (edge weight from v_0 to v_1) is less than infinity, so we update the distance to v_1 to 9 and add it to the heap. For $v_0 \rightarrow v_2$, the existing distance to v_2 is infinity. The distance $0 + 2$ is less than infinity, so update the distance to v_2 to 2 and add v_2 to the heap.
3. We remove the vertex at the top of the heap. The smallest distance among the unvisited vertices is 2 (for v_2), so set v_2 as the current vertex. We examine its neighbors, v_3 and v_4 . For $v_2 \rightarrow v_3$, the existing distance to v_3 is infinity. Since 2 (distance to v_2) + 6 (edge weight from v_2 to v_3) is less than infinity, we update the distance to v_3 to 8 . For $v_2 \rightarrow v_4$, the existing distance to v_4 is infinity. Since $2 + 5$ is less than infinity, we update the distance to v_4 to 7 .
4. Continuing in this manner until the heap becomes empty, we get the final distances as: v_0 : 0 , v_1 : 9 , v_2 : 2 , v_3 : 8 , v_4 : 7 .

7.3 BellmanFord Algorithm

In this section, we give an algorithm that computes the shortest paths from any given vertex even when the edge weights are negative. Why could we not apply Dijkstra's algorithm for this problem? Consider

the case where the node that can be reached from the source node on one edge with a minimum cost w has a cost of x . Let this node be v . Dijkstra's algorithm assumes that this is the shortest path to node v , because any other path would already have incurred the cost of w by going to some other vertex. However, in the presence of negative weights, there may be a longer path to v by incurring more cost initially but then traversing an edge with negative cost.

When there are negative cost edges, we have to be careful that there are no negative cost cycles. In the presence of negative cost cycles, the cost of going from s to some vertex v may be decreased in an unbounded manner by going through the cycle. For now, we will assume that the graph may have negative cost edges but no negative cost cycles.

In the 1950s, Bellman and (independently) Ford gave an algorithm for graphs with negative weights. The algorithm uses dynamic programming. To establish a recurrence relation, we need to come up with an appropriate variable with an index k such that its base case when k is small is easy and when k is large, we reach our goal. We let $d^k[v]$ be the cost of reaching v from the source vertex s with a path of at most k edges. It is clear that the base case of k equal to 0 is quite simple. The value $d^0[v]$ equals 0 when v equals s and ∞ otherwise. When k equals $n - 1$, we claim that $d^k[v]$ is exactly equal to the shortest path from s to v . Why could there not be a path with n edges or more, with a lower cost? If there is a path with n edges, then at least some vertex w appears more than once. The path from w to itself forms a cycle. From our assumption, there are no negative cost cycles. If the cycle is not negative, then we get a smaller path with equal or lower cost by removing the cycle.

The recurrence relation is as follows: $d^0[v] = 0$ if v equals s and ∞ , otherwise.

$$d^k[v] = \min\{d^{k-1}[v], \min_{(u,v) \in E} d^{k-1}[u] + w(u,v)\}$$

We let $p[v]$ denote the predecessor of node v in the shortest path. Initially, $p[v]$ is null for all vertices.

Algorithm [BellmanFord](#) shows the procedure used when the edge weights in a directed graph may be

negative.

Algorithm BellmanFord: Finding the shortest costs from v_0 when edge weights may be negative

Input: Graph $G = (V, E)$, source vertex s , edge weights $w(u, v)$
Output: Distance array $d[]$, Predecessor array $p[]$

```

1 forall vertex  $v \in V$  do
2   |  $d[v] := \infty$ ;
3   |  $p[v] := \text{null}$ ;
4 end
5  $d[s] := 0$ ;
6 for  $k = 1$  to  $|V| - 1$  do
7   | forall edge  $(u, v) \in E$  do
8     |   | if  $d[u] + w(u, v) < d[v]$  then
9       |   |   |  $d[v] := d[u] + w(u, v)$ ;
10      |   |   |  $p[v] := u$ ;
11      |   |   end
12     |   end
13 end
14 forall edge  $(u, v) \in E$  do
15   | if  $d[u] + w(u, v) < d[v]$  then return "Negative cycle detected" ;
16 end
17 return  $d[], p[]$ ;

```

The first *for* loop initializes the value of d and p for the base case corresponding to k equal to zero. We now compute d^k for k equal to $1 \dots n - 1$. Instead of computing the recurrence explicitly for each vertex, we *relax* each edge to check if $d^k[v]$ can be reduced by some edge (u, v) . Finally, we check if there is a negative cost cycle in the graph. If there is a negative cost cycle that can be reached from the source vertex, then there would be an edge (u, v) in the graph such that $d[v]$ can be reduced further by using that edge even after k has reached the value $n - 1$.

The time complexity of BellmanFord is $O(|V| \cdot |E|)$, where $|V|$ is the number of vertices and $|E|$ is the number of edges, making it less efficient than Dijkstra's algorithm for graphs with non-negative weights but more versatile due to its handling of negative weights. The space complexity is $O(|V|)$ for the arrays d and p .

In our algorithm, we have iterated k from 1 to $n - 1$. What if the values of d do not change in some iteration of k ? The reader is invited to change the algorithm so that it checks for this condition and terminates sooner.

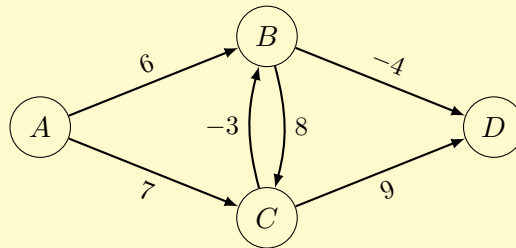
Theorem 7.4 (BellmanFord correctness) *Let $G = (V, E)$ be a directed graph with edge weights w and source s , containing no negative-weight cycle reachable from s . After the main loop of Algorithm [BellmanFord](#) completes, $d[v]$ equals the cost of a shortest path from s to v for every $v \in V$. The final scan correctly reports a negative-weight cycle if and only if one is reachable from s .*

Proof: By induction on k . Let $\delta^k(v)$ denote the cost of a shortest path from s to v with at most k edges (or ∞ if no such path exists). The initialisation gives $d[v] = \delta^0(v)$. Assume that after $k - 1$ iterations, $d[v] \leq \delta^{k-1}(v)$ for all v . Relaxing every edge once in iteration k ensures that for any edge (u, v) , $d[v] \leq d[u] + w(u, v) \leq \delta^{k-1}(u) + w(u, v)$, which is at least $\delta^k(v)$ taken over all such u . Hence $d[v] \leq \delta^k(v)$

after iteration k , and $d[v]$ is never smaller than the true shortest-path cost. Since any shortest path without a negative cycle has at most $n - 1$ edges, $n - 1$ iterations suffice.

For the negative-cycle test: if the algorithm can still relax some edge after $n - 1$ iterations, then there is a path from s to some vertex with cost less than δ^{n-1} , which is only possible if a negative-weight cycle is reachable from s . Conversely, in the presence of such a cycle, edges along it can always be relaxed. ■

Example 7.5 Consider the directed graph below on vertices A, B, C, D with source A . The cycle $B \rightarrow C \rightarrow B$ has weight $8 + (-3) = 5$, and there is no other directed cycle, so the graph has no negative-weight cycle.



Starting from $d[A] = 0$ and $d[B] = d[C] = d[D] = \infty$, the three iterations of the main loop produce:

After iteration	$d[A]$	$d[B]$	$d[C]$	$d[D]$
0 (init)	0	∞	∞	∞
1	0	4	7	2
2	0	4	7	0
3	0	4	7	0

The values stabilise at iteration 2; the subsequent iteration leaves d unchanged, and the negative-cycle check passes. The shortest-path costs from A are therefore $d[A] = 0$, $d[B] = 4$, $d[C] = 7$, $d[D] = 0$.

7.4 FloydWarshall Algorithm

Suppose that $A[i, j]$ represents the weight of the direct edge from i to j denoting the cost of going from i to j using at most one edge. We assume that all diagonal entries are zero and that there is no negative weight cycle. Our goal is to find a matrix $G[i, j]$ such that $G[i, j]$ is the least cost to go from i to j using any number of edges.

Rather than focusing on paths from a single source, we build shortest paths between all pairs of vertices by progressively allowing more intermediate vertices. At stage k , we assume that only vertices $\{1, \dots, k\}$ may appear as intermediates. For each pair (i, j) , the shortest path either avoids vertex k entirely or passes through it, in which case it splits into two shorter subpaths ($i \rightarrow k$) and ($k \rightarrow j$). By systematically considering each vertex as a possible intermediate, we explore all possible path decompositions and obtain the shortest distances between all pairs. We define $d^k(i, j)$ as the cost of going from vertex i to vertex j using intermediate vertices from $1..k$. When k is zero, it is clear that we cannot use any intermediate vertex. Thus,

$$d^0[i, j] = 0 \text{ if } (i = j), w(i, j) \text{ otherwise.}$$

Observe that if there is no edge from i to j when $i \neq j$, $w(i, j)$ is ∞ . Now, consider the case when $k \geq 1$. Then, we get the following recurrence:

$$d^k[i, j] = \min\{d^{k-1}[i, j], d^{k-1}[i, k] + d^{k-1}[k, j]\}$$

Thus, we get the FloydWarshall algorithm (Algorithm [FloydWarshall](#)).

Algorithm FloydWarshall: All pairs shortest path Algorithm

Input: A weighted graph represented by an $n \times n$ cost matrix $G[i, j]$, where $G[i, j]$ is the cost from vertex i to j (possibly ∞ if no edge exists).

Output: Updated matrix $G[i, j]$ containing shortest path costs between all pairs of vertices.

```

1 for k = 1 to n do
2   for i = 1 to n do
3     for j = 1 to n do
4       if G[i, k] + G[k, j] < G[i, j] then
5         | G[i, j] := G[i, k] + G[k, j];
6       end
7     end
8   end
9 end
```

The reader is invited to make two changes to the algorithm. First, modify the algorithm to return an error message when there is a negative cost cycle in the graph. Second, add a variable $p[u][v]$ to the algorithm to return the predecessor of v on the path from u to v .

The time complexity of the FloydWarshall algorithm is $O(n^3)$: the three nested loops iterate n times each and the body is $O(1)$. The space complexity is $O(n^2)$ for the matrix G , with updates done in place.

Theorem 7.6 (FloydWarshall correctness) *On a directed graph without negative-weight cycles, Algorithm [FloydWarshall](#) terminates with $G[i, j]$ equal to the cost of a shortest path from vertex i to vertex j , for every pair (i, j) .*

Proof: Let $d^k[i, j]$ denote the cost of a shortest path from i to j whose internal vertices all lie in $\{1, 2, \dots, k\}$, or ∞ if no such path exists. By induction on k : the base case $k = 0$ matches the input matrix (only direct edges). For the inductive step, any shortest (i, j) -path using internal vertices from $\{1, \dots, k\}$ either avoids k — in which case its cost is $d^{k-1}[i, j]$ — or uses k , in which case its two halves are shortest paths from i to k and from k to j using internal vertices from $\{1, \dots, k-1\}$, of cost $d^{k-1}[i, k] + d^{k-1}[k, j]$. The recurrence taking the minimum of the two is exactly what the algorithm computes in iteration k . After n iterations, $G[i, j] = d^n[i, j]$, which is the unrestricted shortest-path cost since all vertices are now permissible as internal. ■

Example 7.7 Consider a graph on vertices $\{1, 2, 3\}$ with direct-edge costs given by

$$G^{(0)} = \begin{pmatrix} 0 & 4 & 11 \\ 6 & 0 & 2 \\ 3 & \infty & 0 \end{pmatrix}.$$

After iteration $k = 1$ (allowing vertex 1 as an intermediate), only the $(3, 2)$ entry could potentially change: $G[3, 2] = \min(\infty, G[3, 1] + G[1, 2]) = \min(\infty, 3 + 4) = 7$. After iteration $k = 2$, the $(1, 3)$ entry becomes $\min(11, 4 + 2) = 6$ and the $(3, 3)$ entry is checked to be 0. Iteration $k = 3$ updates $(2, 1)$ via $(2, 3, 1)$ to $\min(6, 2 + 3) = 5$. The final all-pairs shortest-path matrix is

$$G = \begin{pmatrix} 0 & 4 & 6 \\ 5 & 0 & 2 \\ 3 & 7 & 0 \end{pmatrix}.$$

7.5 Johnson's Algorithm

Johnson's algorithm solves all-pairs shortest paths on directed graphs whose edge weights may be negative but contain no negative-weight cycle. It combines BellmanFord and Dijkstra: a single BellmanFord run computes a non-negative *price* vector with which the edges are reweighted, and Dijkstra is then run from every source on the reweighted graph. On sparse graphs this beats the $\Theta(n^3)$ FloydWarshall bound: using a Fibonacci-heap Dijkstra yields $O(|V| \cdot |E| + |V|^2 \log |V|)$ total time.

Negative edge weights prevent the direct use of Dijkstra's algorithm, but we can transform the graph so that all edge weights become non-negative while preserving shortest paths. This is done by assigning a potential (or price) to each vertex and adjusting edge weights accordingly. The reweighting shifts all path costs by a constant depending only on endpoints, so relative ordering of paths is unchanged. Once all edges are non-negative, we can efficiently run Dijkstra from each source. Thus, Johnson's algorithm combines the flexibility of BellmanFord (to handle negative edges) with the efficiency of Dijkstra (for fast shortest-path computation).

Let X be the input graph. The *reweighted* graph Y has the same vertices and edges, with edge weights

$$w'[i, j] = w[i, j] + p[j] - p[i], \tag{7.1}$$

where p is a non-negative *price* vector such that every $w'[i, j] \geq 0$. This reweighting preserves shortest paths:

Lemma 7.8 *Let s and t be any two vertices in the graph. The weight of any path on the graph Y from s to t equals the weight in the graph X plus $(p[t] - p[s])$.*

Proof: Let the path be $v_0, v_1, v_2, \dots, v_k$ where $v_0 = s$ and $v_k = t$. As we compute the cost of the path in Y , we add the price of every intermediate vertex once and subtract it once. Only the price of v_0 is subtracted exactly once, and the price of v_k is added exactly once, giving us the lemma. ■

Since path costs shift by a constant $p[t] - p[s]$, shortest paths are unchanged. It remains to find a feasible p , which exists exactly when X has no negative-weight cycle:

Lemma 7.9 *A feasible non-negative price vector exists iff the graph X has no negative-weight cycle. When it does, $p[v] := -\delta(s^*, v)$ is feasible, where $\delta(s^*, v)$ is the BellmanFord distance to v from a fresh auxiliary vertex s^* joined to every $x \in V$ by a 0-weight edge.*

Proof: (\Rightarrow) By Lemma 7.8 with $s = t$, any cycle has the same weight in X and Y , so a negative-weight cycle in X leaves some reweighted edge negative.

(\Leftarrow) Augmenting X with s^* cannot create a new negative cycle since s^* has no incoming edges; hence $\delta(s^*, v)$ is well-defined and ≤ 0 . Set $p[v] := -\delta(s^*, v) \geq 0$. The BellmanFord triangle inequality $\delta(s^*, j) \leq \delta(s^*, i) + w[i, j]$ rearranges to $w[i, j] + p[j] - p[i] \geq 0$, so p is feasible. ■

BellmanFord from s^* finds such a p in at most $|V|$ edge-relaxation passes (matching the longest simple path), and reports a negative cycle otherwise.

Algorithm Johnson: Johnson’s algorithm for all-pairs shortest paths on graphs with negative edges

Input: Directed graph $G = (V, E)$ with edge weights w (possibly negative)

Output: Matrix D with $D[u, v]$ the shortest-path cost from u to v ; or a report of a negative cycle

```

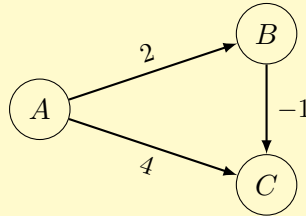
1 Add vertex  $s^*$  and an edge  $(s^*, v)$  of weight 0 for every  $v \in V$ ;
2 Run BELLMANFORD from  $s^*$ ; if it reports a negative cycle, return “negative-weight cycle”;
3  $p[v] := -\delta(s^*, v)$  for every  $v \in V$ ;
4 forall edges  $(i, j) \in E$  do
5   |  $w'[i, j] := w[i, j] + p[j] - p[i]$ ;
6 end
7 forall  $u \in V$  do
8   | Run DIJKSTRA from  $u$  on the graph with weights  $w'$  to get  $\delta'(u, v)$  for every  $v$ ;
9   | forall  $v \in V$  do
10    |  $D[u, v] := \delta'(u, v) + p[u] - p[v]$ ; // undo the reweighting
11    end
12 end
13 return  $D$ ;
```

Theorem 7.10 (Johnson correctness) *On a directed graph without a negative-weight cycle, Algorithm Johnson returns $D[u, v] = \delta(u, v)$ for every pair of vertices. If the graph has a negative-weight cycle, the algorithm reports it.*

Proof: The BellmanFord step correctly finds $\delta(s^*, v)$ or a negative cycle; by Lemma 7.9, $p[v] := -\delta(s^*, v)$ makes every reduced weight non-negative, so each Dijkstra call is valid. Lemma 7.8 gives $\delta'(u, v) = \delta(u, v) + p[v] - p[u]$, and the final line of the algorithm inverts this shift. ■

Complexity is dominated by the single BellmanFord call ($O(|V| \cdot |E|)$) and the $|V|$ Dijkstra calls. With a Fibonacci heap this gives $O(|V| \cdot |E| + |V|^2 \log |V|)$, beating FloydWarshall’s $\Theta(n^3)$ on sparse graphs.

Example 7.11 Consider the directed graph on three vertices A, B, C with edges $A \rightarrow B$ of weight 2, $B \rightarrow C$ of weight -1 , and $A \rightarrow C$ of weight 4:



The graph has no negative-weight cycle (it has no directed cycle at all).

Step 1 (BellmanFord from s^).* Add an auxiliary vertex s^* with a 0-weight edge to A, B , and C . BellmanFord returns $\delta(s^*, A) = 0$, $\delta(s^*, B) = 0$, $\delta(s^*, C) = -1$ (realised by $s^* \rightarrow B \rightarrow C$).

Step 2 (price vector). Set $p := -\delta(s^*, \cdot)$, giving $p(A) = 0$, $p(B) = 0$, $p(C) = 1$.

Step 3 (reweighting). The reduced weights $w'[i, j] = w[i, j] + p[j] - p[i]$ are $w'(A, B) = 2 + 0 - 0 = 2$, $w'(B, C) = -1 + 1 - 0 = 0$, $w'(A, C) = 4 + 1 - 0 = 5$, all non-negative.

Step 4 (Dijkstra from every source on reweighted graph). The reweighted distances are

$$\delta'(A, A) = 0, \quad \delta'(A, B) = 2, \quad \delta'(A, C) = 2 \text{ (via } A \rightarrow B \rightarrow C\text{);}$$

$$\delta'(B, B) = 0, \quad \delta'(B, C) = 0; \quad \delta'(C, C) = 0.$$

(Entries from B or C back to A are ∞ .)

Step 5 (undo the shift). Recover $D[u, v] = \delta'(u, v) + p[u] - p[v]$:

$$D = \begin{pmatrix} 0 & 2 & 1 \\ \infty & 0 & -1 \\ \infty & \infty & 0 \end{pmatrix},$$

where rows are indexed by u and columns by v . Verification: the $A \rightarrow C$ entry picks the two-edge path $A \rightarrow B \rightarrow C$ of cost $2 + (-1) = 1$, cheaper than the direct $A \rightarrow C$ edge of cost 4.

7.6 LLP Perspective

In this section we collect the LLP reformulations of the algorithms introduced earlier in the chapter.

LLP-Dijkstra Algorithm

We now cast Dijkstra's algorithm (Section 7.2) in the LLP framework. The objects being computed are the distances $d[0..n-1]$ from the source vertex v_0 . Using the same lattice as in the LLP-BellmanFord formulation, we regard distances as elements of $(\mathbb{R}_{\geq 0} \cup \{\infty\})^n$ and place the "small" distances *higher* in the lattice: $d \leq_L d' \iff \forall i : d[i] \geq d'[i]$. The feasibility predicate is the shortest-path recurrence

$$B \equiv \forall (i, j) \in E : d[j] \leq d[i] + w[i, j],$$

which is lattice-linear by the same argument used for LLP-BellmanFord (Theorem).

Dijkstra's algorithm is a particular LLP schedule that exploits non-negative edge weights to fix one vertex at a time. Let $fixed[i]$ be a Boolean flag (initially $fixed[0] = true$, $fixed[i] = false$ for $i \neq 0$), and initialise $d[0] := 0$ and $d[i] := \infty$ for $i \neq 0$. An index j is *forbidden* when it is the unique non-fixed vertex whose current $d[j]$ is the minimum among non-fixed vertices; the advance rule fixes j and relaxes its outgoing edges.

The efficient implementation maintains the set of non-fixed discovered vertices in a min-heap H keyed by d . Extracting the minimum of the heap yields the forbidden vertex at each round, and edge relaxation inserts updated keys back into the heap. The algorithm below is literally Algorithm [Dijkstra](#) rewritten in LLP notation.

Algorithm LLP-Dijkstra: Dijkstra's algorithm as an LLP schedule

Input: Graph (V, E) with non-negative edge weights $w[i, j]$; source v_0

Output: $d[0..n-1]$: real, initially $d[0] = 0$ and $d[i] = \infty$ for $i \neq 0$

```

1 fixed: array[0..n-1] of boolean, initially  $\forall i : fixed[i] = false$ ;
2 forbidden( $j$ ):  $\neg fixed[j] \wedge d[j] = \min_{k:\neg fixed[k]} d[k]$ ;
3   advance:  $fixed[j] := true$ ; forall  $(j, k) \in E$  with  $\neg fixed[k]$ :  $d[k] := \min(d[k], d[j] + w[j, k])$ ;
   // Efficient implementation maintains non-fixed vertices in a min-heap  $H$  keyed by
   //  $d$ ; extract-min returns the forbidden vertex and insertions install reduced
   // keys.
```

Correctness follows from the same invariant as in Section 7.2: whenever a vertex is extracted from the heap with key d_j , non-negativity of the edge weights guarantees that no still-undiscovered vertex can lead to a shorter path to j , so j can safely be fixed. Lemma 7.1 captures both parts of this invariant.

The LLP formulation makes clear that Dijkstra's algorithm differs from LLP-BellmanFord only in its schedule: BellmanFord advances every forbidden index concurrently, while Dijkstra advances exactly one index per round (the non-fixed vertex of minimum d). The min-heap makes this schedule efficient, yielding the same $O(m \log n)$ sequential running time as the heap-based Dijkstra of Section 7.2.

LLP-BellmanFord Algorithm

Consider the distance of other vertices from s . To apply LLP algorithm, we view finding the optimal distance vector as finding the largest $d[x]$ for every vertex $x \neq v_0$ which satisfies the following feasibility predicate:

$$\text{for all edges } (i, j) \in E : d[j] \leq d[i] + w[i, j]$$

We define B_e for any edge $e = (i, j)$ as $d[j] \leq d[i] + w[i, j]$. The feasibility predicate B can be written as

$$B \equiv \bigwedge_{e \in E} B_e$$

We will view the search for optimal distance vector as searching for d vector in a lattice defined as follows. The lattice has the bottom element as (M, M, \dots, M) where M equals $n-1$ times the largest weight edge in the graph. Since we are looking for the largest distance vector satisfying B , it can never be greater than M . In this lattice $d \leq_L d'$ iff $\forall i : d[i] \geq d'[i]$. Thus, finding the least vector in this lattice corresponds to finding the largest vector d that satisfies B .

We now claim that

Theorem 7.12 *B is a lattice linear predicate.*

Proof: It is sufficient to show that B_e is lattice linear because any conjunction of linear predicates is also lattice linear. B_e for $e = (i, j)$ is equivalent to $d[j] \leq d[i] + w[i, j]$. This is equivalent to $d[j] \geq_L d[i] + w[i, j]$. Since the right hand side is a monotonic function of d , we get that B_e is lattice linear. ■

Algorithm LLP-Edge-Relaxation: Finding the minimum distance vector satisfying B

Input: $pre(j)$: list of $1..n$; $w[i, j]$: int for all $i \in pre(j)$

Output: $d[j]$: int, initially if $(j = s)$ then 0 else *maxint*

1 **ensure:** $d[j] \leq \min\{d[i] + w[i, j] \mid i \in pre(j)\}$;

Algorithm [LLP-Edge-Relaxation](#) gives the correct answer; however, it may result in large computation complexity if d 's are lowered in a non-disciplined manner. We now optimize this algorithm. Algorithm [LLP-BellmanFord](#) chooses forbidden indices more carefully. Whenever there is a forbidden vertex, all forbidden vertices are advanced. The *while* loop checks if there is any forbidden vertex. The *forall* loop advances all forbidden vertices.

Algorithm LLP-BellmanFord: Finding the minimum distance vector satisfying B

Input: $pre(j)$: adjacency list; $w[i, j]$: edge weights

Output: $d[0..n - 1]$: array of real initially $(d[0] = 0) \wedge (\forall i \neq 0: d[i] = \infty)$

1 **forbidden**(j): $\exists i \in pre(j) : d[j] > d[i] + w[i, j]$;

2 **advance**(j): $\forall k : d[k] := \min\{d[i] + w[i, k] \mid i \in pre(k)\}$;

Let us analyze the sequential time complexity of Algorithm [LLP-BellmanFord](#). There are at most n iterations of the *while* loop. The *forall* loop can be implemented with $O(m)$ time complexity giving us the overall sequential time complexity of $O(mn)$.

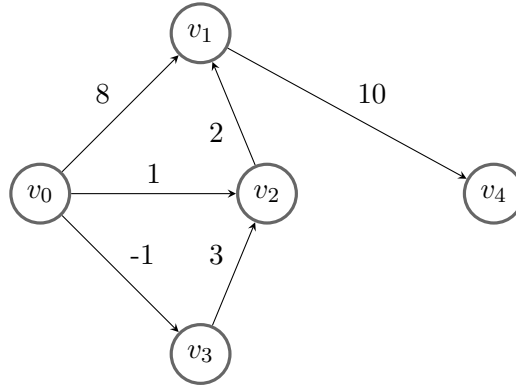
Earlier, we had assumed that the graph does not have any negative cost cycle. What if the user input is such that it has a negative cost cycle? In that case, Algorithm [LLP-BellmanFord](#) will never terminate. The standard fix, as in the classical BellmanFord algorithm of Section 7.3, is to cap the number of *while*-loop iterations at $|V| - 1$ and report “negative cost cycle” if any index is still forbidden after that.

LLP-FloydWarshall

We now give an LLP formulation of the FloydWarshall algorithm. Suppose that $A[i, j]$ represents the weight of the direct edge from i to j , denoting the cost of going from i to j using at most one edge. We assume that all diagonal entries are zero and that there is no negative weight cycle. Our goal is to find a matrix $G[i, j]$ such that $G[i, j]$ is the least cost of going from i to j by using any number of edges.

Let us formulate this problem as finding the least vector of size n^2 in a distributive lattice. We initialize G as A . This is the bottom element of our distributive lattice. Our goal is to find the least vector that satisfies the following constraint:

$$B_{FloydWarshall} \equiv \forall i, j : i \neq j : G[i, j] \leq \min_k \{G[i, k] + G[k, j]\}.$$



When i equals j , $G[i, j]$ equals 0 and will stay fixed through the execution of the algorithm. The cost of going from i to j must be smaller than the cost of going from i to k and k to j . Algorithm [LLP-FloydWarshall](#) gives a high-level LLP algorithm for all pairs shortest path.

Algorithm LLP-FloydWarshall: Computing the shortest cost matrix for a directed graph

- 1 **input:** A : matrix of real; // $A[i, j]$ is the weight of the edge from i to j ;
 - 2 **var** G : matrix of real initially $\forall i, j : G[i, j] = A[i, j]$;
 - 3 **forbidden**(i, j): $(\exists k : G[i, j] > G[i, k] + G[k, j])$
 - 4 **advance:** $G[i, j] := \min_k \{G[i, k] + G[k, j]\}$;
-

LLP-Johnson

We now revisit Johnson's reweighting framework of Section 7.5. Recall the reweighting identity $w'[i, j] = w[i, j] + p[j] - p[i]$ from Equation 7.1 and Lemma 7.8, which says that every path from s to t in the reweighted graph Y has weight $w_X(\text{path}) + p[t] - p[s]$. The sequential algorithm of the previous section uses BellmanFord from an auxiliary source s^* to obtain a feasible price vector. Here we give an LLP algorithm that finds such a price vector directly. Our feasibility predicate B for the pricing vector is

$$\forall (i, j) \in E : w[i, j] + p[j] - p[i] \geq 0$$

Furthermore, we require $p[i] \geq 0$ for all i . We first show that B is lattice linear.

Lemma 7.13 *Let X be any graph such that the edge (i, j) has weight $w[i, j]$ and every vertex i has price $p[i]$. Consider the lattice of all non-negative price vectors. Then, the predicate*

$$B \equiv \forall (i, j) \in E : w[i, j] + p[j] - p[i] \geq 0$$

is lattice linear.

Proof: Since the lattice linearity is closed under conjunction, it suffices to show that $B_e \equiv w[i, j] + p[j] - p[i] \geq 0$ is lattice linear for an arbitrary edge $e = (i, j)$. B_e can be rewritten as $p[j] \geq p[i] - w[i, j]$.

The right-hand side of this inequality is a monotone function on p and hence, from the Key Lemma of lattice-linearity, we get that B_e is lattice linear. ■

By applying the LLP algorithm, we get the following method to find the price vector.

Algorithm LLP-Johnson: Finding the minimum price vector.

Input: $pre(j)$: list of $1..n$; $w[i, j]$: int for all $i \in pre(j)$

Output: $p[j]$: int, initially 0

- 1 **forbidden**(j): $\exists i \in pre(j) : p[j] < p[i] - w[i, j]$;
 - 2 **advance**(j): $\forall k : p[k] := \max\{p[i] - w[i, k] \mid i \in pre(k)\}$;
-

Termination and iteration bounds for Algorithm [LLP-Johnson](#) match those of the sequential construction in Section [7.5](#): by Lemma [7.9](#), the advance rule converges to a feasible price vector iff the graph has no negative-weight cycle, and at most $|V|$ rounds of edge relaxation suffice. From the Key Lattice-Linearity Lemma, the set of all feasible price vectors is closed under meets, so the LLP algorithm converges to the componentwise-minimum feasible price vector; since adding a positive constant to all prices preserves feasibility and leaves reduced weights unchanged, this minimum vector has at least one zero component.

Each LLP iteration can be computed in $O(m)$ time, so the graph is “reweighted” in $O(mn)$ time — the same asymptotic bound that BellmanFord achieves in the sequential algorithm. Plugging the resulting price vector into Dijkstra from every source yields an all-pairs shortest-path algorithm with running time $O(mn + n(m + n \log n)) = O(mn + n^2 \log n)$, which is faster than the $O(n^3)$ FloydWarshall algorithm whenever the graph is not dense.

7.7 Summary

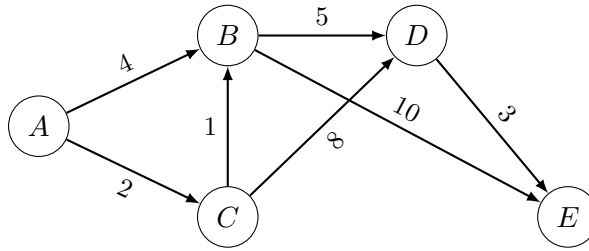
Table [7.1](#) lists all the algorithms discussed in this chapter.

Problem	Algorithm	Time Complexity
Shortest Path	Dijkstra	$O(m \log n)$
Shortest Path	BellmanFord	$O(mn)$
All Pairs Shortest Path	FloydWarshall	$O(n^3)$
All Pairs Shortest Path	Johnson	$O(mn + n^2 \log n)$
Shortest Path	LLP-Dijkstra	$O(m \log n)$
Shortest Path	LLP-BellmanFord	$O(mn)$
All Pairs Shortest Path	LLP-FloydWarshall	$O(n^3)$
Graph Conversion	LLP-Johnson	$O(mn)$

Table 7.1: Algorithms discussed in this chapter.

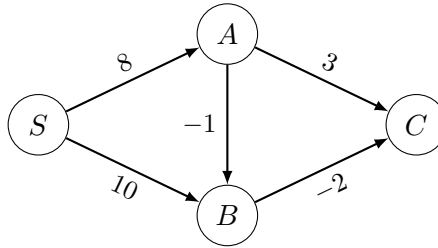
7.8 Problems

1. Consider the directed graph below:



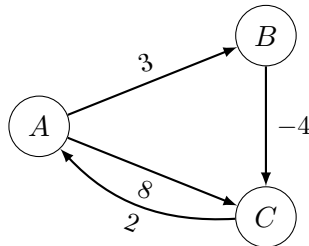
Find the shortest path from A to E using Dijkstra's algorithm.

2. Consider the directed graph below:



Find the shortest path from S to C using the BellmanFord algorithm.

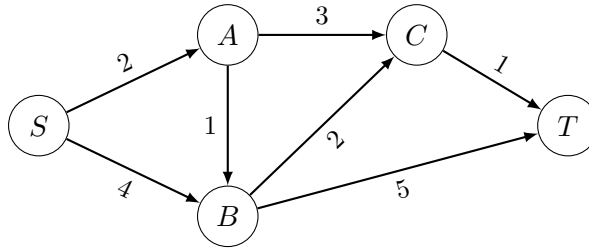
3. Consider the directed graph below:



Find the shortest path for all pairs using the FloydWarshall algorithm.

4. Prove Lemma 7.1.
5. Write an LLP algorithm that computes the reflexive transitive closure of a Boolean adjacency matrix A of an n -vertex directed graph. Specify (a) the lattice, (b) the predicate B that characterises the solution, (c) the forbidden condition, and (d) the advance rule. Argue that B is lattice-linear.
6. Give an algorithm to determine which of the edges in a directed graph are sensitive to the shortest path between s and t . An edge is sensitive, if its weight is reduced, then the shortest path from s to t is also reduced.
7. Bidirectional Search: Suppose that we want to find shortest path from s to t . We can run Dijkstra's algorithm in the forward direction from s and in the backward direction from t . If there is any vertex that is fixed in both directions, then we can terminate. Prove it. (MIT course)

8. Consider the directed graph below:



Find the shortest path from S to T such that the path does not use both edges $A \rightarrow B$ and $B \rightarrow T$ together (i.e., the pair $(A \rightarrow B, B \rightarrow T)$ is forbidden).

9. *Shortest paths in a DAG.* Let $G = (V, E, w)$ be a directed acyclic graph with weights w (possibly negative). Describe an $O(|V| + |E|)$ -time algorithm that computes the shortest-path distances from a source vertex s to every other vertex. Why does your algorithm not require the non-negativity assumption of Dijkstra's algorithm?
10. *Currency arbitrage.* Let c_1, c_2, \dots, c_n be currencies, and let $r_{ij} > 0$ denote the exchange rate from c_i to c_j (so that one unit of c_i can be exchanged for r_{ij} units of c_j). A sequence of exchanges $c_{i_0} \rightarrow c_{i_1} \rightarrow \dots \rightarrow c_{i_k} \rightarrow c_{i_0}$ is an *arbitrage* if the product of rates around the cycle exceeds 1. Describe an $O(n^3)$ -time algorithm that decides whether an arbitrage exists.
11. *Bottleneck shortest path.* Let $G = (V, E, w)$ be an undirected graph with non-negative edge weights, and let $s, t \in V$. Define the *bottleneck* of a path P as $\max_{e \in P} w(e)$. Describe an algorithm that finds an s - t path of minimum bottleneck, and state its time complexity.
12. *Shortest path with at most k edges.* Given a weighted directed graph $G = (V, E, w)$ (possibly with negative edges, but no negative-weight cycles) and an integer $k \geq 0$, describe an algorithm that computes, for every vertex v , the minimum cost of a path from a source s to v using at most k edges. State its running time.

7.9 Bibliographic Remarks

The single source shortest path problem has a rich history. For the history of Dijkstra's algorithm, the reader is referred to the book by [Eri19]. One popular research direction is to improve the worst case complexity of Dijkstra's algorithm by using different data structures. For example, by using Fibonacci heaps for the min-priority queue, Fredman and Tarjan [FT87] gave an algorithm that takes $O(e + n \log n)$. There are many algorithms that run faster when weights are small integers bounded by some constant γ . For example, Ahuja et al. [AMOT90] gave an algorithm that uses Van Emde Boas tree as the priority queue to give an algorithm that takes $O(e \log \log \gamma)$ time. Thorup [Tho00] gave an implementation that takes $O(n + e \log \log n)$ under special constraints on the weights. Raman [Ram97] gave an algorithm with $O(e + n\sqrt{\log n \log \log n})$ time. The LLP algorithm for the shortest path is taken from [AKG20]. Bellman and Ford's algorithm is from [Bel58] and [For56].

Chapter 8

The Minimum Spanning Tree Problem

8.1 Introduction

In this chapter, we discuss the problem of finding the minimum spanning tree (MST) of an undirected weighted graph. Let (V, E) be a connected undirected graph with $n = |V|$ vertices and $m = |E|$ edges, where each edge $e \in E$ has a weight $w(e) \in \mathbb{R}$. A *minimum spanning tree* (MST) of the graph is a subset $T \subseteq E$ such that:

1. T connects all vertices in V (i.e., (V, T) is connected),
2. The total weight $\sum_{e \in T} w(e)$ is minimized among all such subsets.

Equivalently, T is a tree that spans all vertices with the minimum possible total edge weight. A maximum spanning tree is defined similarly, but maximizes the total weight instead.

We assume that all edge weights are distinct. It is known that if the graph is connected and all edge weights are distinct, then there is a unique minimum spanning tree. If the graph is not connected, then there is a unique minimum spanning forest. For simplicity of exposition, we will assume that the underlying graph is connected.

The primary motivation for finding an MST is to connect all nodes in a network with the minimum total cost while ensuring connectivity and avoiding cycles. This has numerous real-world applications:

- *Network Design*: In telecommunications or electrical grids, an MST minimizes the cost of laying cables or wires to connect all points (e.g., cities or computers) without redundant loops.
- *Transportation Networks*: Building roads or pipelines between locations with minimal total length or cost.
- *Clustering and Approximation Algorithms*: MSTs are used in algorithms for clustering data points or approximating solutions to NP-hard problems like the Traveling Salesman Problem (TSP), where an MST provides a lower bound for tours.
- *Computer Networks*: Efficient broadcasting and routing in networks, reducing latency and resource usage.

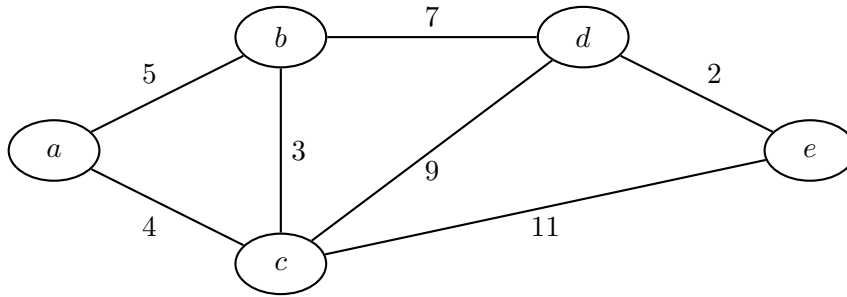


Figure 8.1: An undirected weighted graph

This chapter is organized as follows. Section 8.2 describes the key properties of a minimum spanning tree, including the notion of a *fragment*. Section 8.3 describes the Find-Union data structure, which is used by several MST algorithms to detect cycles efficiently. Section 8.4 presents Kruskal’s algorithm, a greedy algorithm that chooses the least-cost feasible edge. Section 8.5 presents Prim’s algorithm, another greedy algorithm that grows a single fragment. Section 8.6 describes Boruvka’s algorithm, which extends multiple fragments in every iteration. Section 8.8 reformulates these algorithms in the LLP framework.

8.2 Key Properties of a Minimum Spanning Tree

The notion of a *fragment* is crucial in understanding MST algorithms. A fragment is simply a subtree of the MST. Consider the graph in Fig. 8.1. The minimum spanning tree in this graph corresponds to the edges $\{2, 3, 4, 7\}$. The subtree formed by the edges 3 and 4 is a fragment with three vertices $\{a, b, c\}$ and two edges $\{(a, c), (b, c)\}$. A crucial property of the MST is as follows.

Lemma 8.1 *Let F be a fragment. Let e be the edge with the minimum weight that is outgoing from F . Then $F \cup \{e\}$ is also a fragment.*

Proof: Let T be the minimum spanning tree. If $e \in T$, then $F \cup \{e\}$ is trivially a fragment. If $e \notin T$, adding e to T creates a unique cycle C . This cycle must contain another edge f that is outgoing from F (since the cycle enters and exits F). Since $w(e) < w(f)$ (as e has minimum outgoing weight), $T' = T \setminus \{f\} \cup \{e\}$ is a spanning tree with $w(T') < w(T)$, contradicting the minimality of T . Hence $e \in T$ and $F \cup \{e\}$ is a fragment. ■

In the fragment formed by edges with weights 3 and 4, there are three outgoing edges — edges with weights 7, 9, and 11. The edge 5 is not outgoing, as it connects vertices that are part of the fragment. According to Lemma 8.1, the edge with weight 7 can be added to the edges with weights 3 and 4 to grow the fragment.

The following two properties, phrased in terms of edge-weight extremes across a cut and around a cycle, are the workhorses behind every correctness proof in this chapter.

Lemma 8.2 (Cut Property) *Let $S \subsetneq V$ and let e be the minimum weight edge crossing the cut $(S, V \setminus S)$. Then e belongs to some MST; under distinct weights, e belongs to every MST.*

Proof: Let T be any MST and suppose $e \notin T$. Adding e to T creates a unique cycle, and since the cycle starts and ends in S it must contain another edge $f \neq e$ that also crosses the cut. By hypothesis $w(e) \leq w(f)$, so $T' = T \cup \{e\} \setminus \{f\}$ is a spanning tree with $w(T') \leq w(T)$; hence T' is an MST containing e . Under distinct weights, $w(e) < w(f)$ and the inequality is strict, forcing e to be in every MST. ■

Lemma 8.3 (Cycle Property) *Let C be a cycle in the graph and let e be the strictly maximum weight edge in C . Then e does not belong to any MST.*

Proof: Suppose, for contradiction, that $e \in T$ for some MST T . Removing e partitions V into two components S and $V \setminus S$. Because C starts and ends on the same side, some other edge $f \in C \setminus \{e\}$ also crosses the cut $(S, V \setminus S)$. By hypothesis $w(f) < w(e)$, so $T' = T \setminus \{e\} \cup \{f\}$ is a spanning tree with $w(T') < w(T)$, contradicting the optimality of T . ■

8.3 The Find-Union Data Structure

A central operation in several MST algorithms — and in Kruskal’s algorithm in particular — is to test whether the next edge under consideration would close a cycle in the partially built forest. The Find-Union data structure (also known as Disjoint Set Union) supports this test efficiently. It maintains a collection of disjoint sets and supports two primary operations:

- **Find:** Determines which set a particular element belongs to by returning its representative (or “root”). With path compression, this operation is nearly constant time, $O(\alpha(n))$, where $\alpha(n)$ is the inverse Ackermann function.
- **Union:** Merges two sets into one by linking their roots, typically using rank or size heuristics to keep trees balanced, also achieving $O(\alpha(n))$ amortized time.

Initially, each element is in its own set. When this structure is used to build a spanning forest, the *Union* operation merges sets of vertices connected by selected edges, while the *Find* operation checks if adding an edge would form a cycle (i.e., if its endpoints are already in the same set).

Each set is maintained as a rooted tree in which every node has a *parent* pointer. The root of the tree serves as the identity of the set. The *Find* operation finds the root of the tree. Observe that finding the root of the tree by traversing the parent pointer takes time proportional to the distance of the node from the root. One method of reducing this distance in future traversals is *path compression*: we make all the nodes on the find path point directly to the root, so subsequent traversals from those nodes are essentially constant time.

The *Union* operation merges two trees by making the root of one of them point to the other root. For correctness, it does not matter which root points to the other root. However, to keep the lengths of the

longest paths small, we use the notion of *rank* to decide. Our rule is as follows: the root of the tree with the lower rank points to the root with the higher rank. The rank of the new tree is the same as that of the bigger rank. If both roots have the same rank, then it does not matter which one becomes the new root, and the rank of the new tree is one more than the rank of the old trees. It is a simple exercise to show that the rank of any tree with n nodes is at most $O(\log n)$ and the height of the tree is at most its rank. With these optimizations, any sequence of n Find and Union operations takes $O(n\alpha(n))$ amortized time.

Algorithm Find: Finding the root of x 's set with path compression

Input: Vertex x

Output: Root of the set containing x

```

1 parent: array[1.. $n$ ] of int initially  $\forall y : \text{parent}[y] = y$ ;
2 if  $\text{parent}[x] \neq x$  then  $\text{parent}[x] := \text{Find}(\text{parent}[x])$  //path compression ;
3 return  $\text{parent}[x]$ 

```

Algorithm Union: Computing Union by rank of sets containing x and y

Input: Vertices x, y

Output: Unified set containing x and y

```

1 parent: array[1.. $n$ ] of int initially  $\forall v : \text{parent}[v] = v$ ;
2 rank: array[1.. $n$ ] of int initially  $\forall v : \text{rank}[v] = 0$ ;
3  $\text{root}X := \text{Find}(x)$ ;
4  $\text{root}Y := \text{Find}(y)$ ;
5 if  $\text{root}X = \text{root}Y$  then return // already in same set, no union needed ;
6 if  $\text{rank}[\text{root}X] < \text{rank}[\text{root}Y]$  then  $\text{parent}[\text{root}X] := \text{root}Y$  ;
7 else if  $\text{rank}[\text{root}X] > \text{rank}[\text{root}Y]$  then  $\text{parent}[\text{root}Y] := \text{root}X$  ;
8 else  $\text{parent}[\text{root}Y] := \text{root}X$ ;
9  $\text{rank}[\text{root}X] := \text{rank}[\text{root}X] + 1$  // increment rank if equal ;

```

8.4 Kruskal's Algorithm

Kruskal's algorithm is a canonical greedy algorithm for the minimum-spanning tree problem. It chooses edges one at a time in a greedy fashion. Let T be the set of edges chosen by the algorithm at any stage. The algorithm maintains the invariant that T does not have any cycle. Initially T is empty and therefore trivially satisfies the invariant. The algorithm considers the edges in increasing order of weights. For this reason, it is sometimes also known as the *shortest-edge-next* algorithm. Suppose that the algorithm has chosen the edges in T so far. To grow T , it finds the edge with the least weight that does not form a cycle with existing edges in T . If any edge e forms a cycle with T , it is rejected, and the algorithm considers the least weight edge of the remaining edges. We use the Find-Union data structure of Section 8.3 to detect cycles efficiently: an edge (u, v) closes a cycle in T iff $\text{Find}(u) = \text{Find}(v)$.

In Algorithm [Kruskal](#), the variable T keeps the set of all edges chosen. At every step, (V, T) is an acyclic subgraph — that is, a spanning forest. The Find-Union structure represents the connected components of this forest: $\text{Find}(u)$ returns the representative of u 's component, and $\text{Union}(u, v)$ merges two components when an edge is added. Adding the next lightest edge either joins two distinct components (reducing the number of components by one) or closes a cycle within a single component (in which case the edge is rejected). The algorithm terminates when $|T| = n - 1$, at which point the forest has coalesced into a

Algorithm Kruskal: Computing Minimum Spanning Tree**Input:** Undirected Weighted Graph (V, E, w) with $n = |V|$ **Output:** Minimum Weight Spanning Tree T

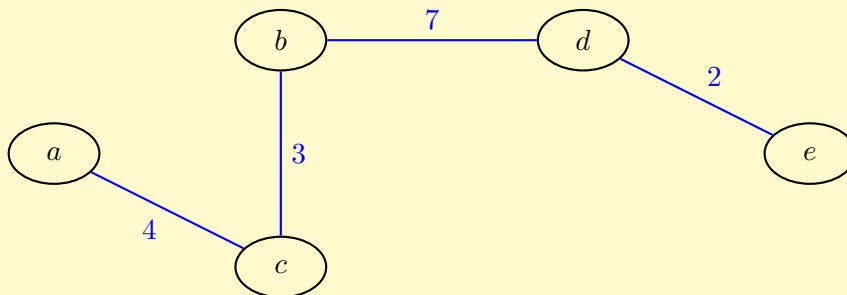
```

1  $T$ : set of edges, initially  $\{\}$ ;
2 forall  $v \in V$  do
3    $parent[v] := v$ ;  $rank[v] := 0$ ;           // MakeSet for each vertex
4 end
5 sort the edges of  $E$  in non-decreasing order of weight;
6 foreach  $edge (u, v) \in E$  in sorted order do
7   if  $Find(u) \neq Find(v)$  then
8      $T := T \cup \{(u, v)\}$ ;
9      $Union(u, v)$ ;
10    if  $|T| = n - 1$  then return  $T$ ;
11  end
12 end
13 return  $T$ ;                               // graph is disconnected if  $|T| < n - 1$ 

```

spanning tree, or when the edges have all been processed and $|T| < n - 1$, indicating that the input graph is disconnected.

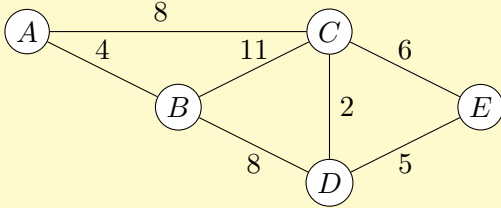
Example 8.4 Consider the graph shown in Fig. 8.1. Kruskal's algorithm first chooses the edge with weight 2 since it has the least weight. Then, it chooses the edges with weight 3 and 4 because these edges do not form any cycle. The next edge has weight 5; this edge is rejected because it forms a cycle with the already chosen edges with weights 3 and 4. The algorithm then chooses the edge with weight 7 and terminates with $T = \{2, 3, 4, 7\}$. The resulting MST is shown below.



Correctness. We argue by induction that T can always be extended to an MST. Initially $T = \emptyset$ extends trivially. Suppose T extends to some MST T^* and the algorithm is about to add edge $e = (u, v)$. If $e \in T^*$, we are done. Otherwise, let S be the set of vertices in u 's tree in the forest (V, T) . Since $Find(u) \neq Find(v)$, $v \notin S$, so e crosses the cut $(S, V \setminus S)$. The unique path from u to v in T^* must also cross this cut on some edge f ; this edge has not yet been processed by the algorithm (any earlier edge has been either accepted, which would have put v in S , or rejected, which requires it to close a cycle with edges in T — contradicting $f \in T^*$). Hence $w(f) \geq w(e)$, and $T^* \cup \{e\} \setminus \{f\}$ is an MST containing $T \cup \{e\}$. By induction, the final T is an MST.

Complexity. The algorithm is dominated by the cost of sorting the edges: $O(m \log n)$. Each *Find* or *Union* takes $O(\alpha(n))$ amortized time, so the total cost of the $O(m)$ Find-Union operations is $O(m\alpha(n))$, which is dominated by sorting. The overall complexity of Kruskal's algorithm is therefore $O(m \log n)$.

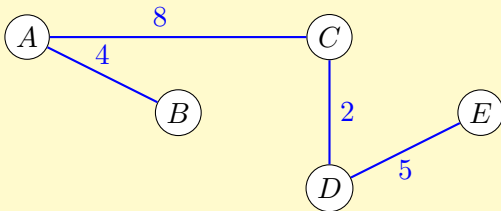
Example 8.5 Consider a graph with five vertices A, B, C, D, E and edges $A-B: 4$, $A-C: 8$, $B-C: 11$, $B-D: 8$, $C-D: 2$, $C-E: 6$, $D-E: 5$. We apply Kruskal's algorithm.



Edges in sorted order: $C-D$ (2), $A-B$ (4), $D-E$ (5), $C-E$ (6), $A-C$ (8), $B-D$ (8), $B-C$ (11). Initially each vertex is its own set. Process each edge with *Find* and *Union*:

1. $C-D$ (2): $Find(C) \neq Find(D)$, so $Union(C, D) \rightarrow \{A\}, \{B\}, \{C, D\}, \{E\}$.
2. $A-B$ (4): $Find(A) \neq Find(B)$, so $Union(A, B) \rightarrow \{A, B\}, \{C, D\}, \{E\}$.
3. $D-E$ (5): $Find(D) \neq Find(E)$, so $Union(D, E) \rightarrow \{A, B\}, \{C, D, E\}$.
4. $C-E$ (6): $Find(C) = Find(E)$; skip.
5. $A-C$ (8): $Find(A) \neq Find(C)$, so $Union(A, C) \rightarrow \{A, B, C, D, E\}$. MST complete; the remaining edges $B-D$ (8) and $B-C$ (11) would close cycles.

The MST consists of $C-D$ (2), $A-B$ (4), $D-E$ (5), $A-C$ (8), with total weight 19.



8.5 Prim's Algorithm

Prim's algorithm is also a greedy algorithm. It builds the minimum spanning tree by increasing the size of a single fragment by adding the minimum weight outgoing edge of the fragment. It simply exploits Lemma 8.1 to increase the size of the fragment until it becomes the MST. At any stage, Prim's algorithm has a fragment F . It finds the minimum outgoing edge of that fragment e . This edge can be viewed as the edge from the fragment to its nearest neighbor. Therefore, this algorithm is sometimes also known as the *nearest-neighbor-next* algorithm. To find the nearest neighbor, every vertex v maintains a label d that corresponds to the cost of adding v to the fragment. At each iteration, the algorithm chooses the vertex v with the minimum value d and adds it to the fragment. The array *fixed* keeps track of the vertices in the fragment. Whenever a new vertex v is *fixed* and added to the fragment, the d values for any adjacent

vertex v' are updated as follows. We check whether the weight of the edge (v, v') is lower than the previous value of $d[v']$. If this is true, then $d[v']$ is updated to $w[v, v']$. We also use a *parent* pointer with each node, which keeps track of the node v that is responsible for the d value of v' .

Algorithm Prim: Finding a Minimum Spanning Tree (MST)

Input: Undirected Weighted Graph: (V, E, w)
Output: Minimum Weight Spanning Tree T

```

1  $d$ : array[0.. $n - 1$ ] of real initially  $\infty$ ; //  $d[v]$  is the cost to add vertex  $v$ ;
2  $parent$ : array[0.. $n - 1$ ] of int initially  $-1$ ; //  $parent[v]$  is the node that corresponds to  $d[v]$  cost;
3  $fixed$ : array[0.. $n - 1$ ] of boolean initially false; // vertices whose  $d$  value is fixed;
4  $T$ : set of edges initially  $\{\}$ ;
5  $d[0] := 0$ ;
6 while  $|T| < n - 1$  do
7    $v := arg\ min_i \{d[i] \mid \neg fixed[i]\}$ ;
8   if  $d[v] = \infty$  then return null // no spanning tree in the graph;
9   if  $parent[v] \neq -1$  then add  $(v, parent[v])$  to  $T$ ;
10   $fixed[v] := true$ ;
11  forall  $(v, v') \in E$  do
12    if  $w[v, v'] < d[v']$  then
13       $d[v'] := w[v, v']$ ;
14       $parent[v'] := v$ ;
15    end
16  end
17 end

```

Example 8.6 Consider again the graph shown in Fig. 8.1. For Prim's algorithm, we start from a fixed node. Suppose that we start from the vertex a . Then the nearest neighbor is c with a cost of 4. The next nearest neighbor to the fragment with vertices $\{a, c\}$ is the vertex b . The cost of adding b is 3. At this point, we have the vertices $\{a, b, c\}$ in the fragment. The cost of adding vertex d is 7 and the cost of adding vertex e is 11. We add the vertex d to our fragment with the cost 7. Finally, e is added with the cost 2. Hence, the edges are added to the tree in the order 4, 3, 7, 2. The set of edges chosen is identical to that of Kruskal's algorithm, even though the order in which they are chosen is different. This is not surprising because there is a unique minimum spanning tree when all edge weights are distinct (Problem 2).

Correctness. Prim's algorithm maintains the invariant that the fragment F (the set of fixed vertices together with T) is contained in some MST. Initially $F = (\{v_0\}, \emptyset)$, which is trivially contained in every MST. Suppose the invariant holds just before an iteration that fixes vertex v . By construction, the edge $(parent[v], v)$ is the lightest edge crossing the cut $(F, V \setminus F)$: its weight is $d[v]$, and for any other edge (u', v') with $u' \in F, v' \notin F$ we have $d[v'] \geq w(u', v')$ and $d[v] \leq d[v']$. By the cut property (Lemma 8.2), this edge belongs to some MST that extends F . After $n - 1$ iterations, F is a spanning tree, and hence an MST.

Complexity. The step of finding the vertex v with a minimum value d can be performed by simply traversing the array d or maintaining d in a heap. If we simply traverse the array d to find the minimum, the work complexity of the above algorithm is $O(n^2 + m)$. In every iteration of the while loop, we perform $O(n)$ work to find the minimum, and there are $O(n)$ iterations of the loop. The work for processing edges over all iterations is $O(m)$ because every edge is processed at most once. If we use a heap to store d values of all vertices that are not fixed, we require $O(m)$ insertions in the heap resulting in $O(m \log n)$ work complexity.

Comparison with Dijkstra's algorithm. Prim's and Dijkstra's algorithms (Chapter 7) share the same control structure: both maintain a priority-keyed set of frontier vertices, repeatedly extract the one with the smallest key, fix it, and relax the keys of its neighbours. The difference lies entirely in what the key means. In Prim, $d[v']$ is the weight of the *single* lightest edge from the current fragment to v' , so the relaxation rule reads $d[v'] := \min(d[v'], w(v, v'))$. In Dijkstra, $d[v']$ is the length of the shortest path from the source to v' through already-fixed vertices, so the relaxation rule becomes $d[v'] := \min(d[v'], d[v] + w(v, v'))$. As a consequence, Prim's correctness rests on the cut property and holds for arbitrary real edge weights (including negative), whereas Dijkstra's correctness additionally requires that extending a path cannot shorten it, which is true only for non-negative weights.

8.6 Boruvka's Algorithm

In Prim's algorithm, we started with a trivial fragment that included only the vertex v_0 . We kept increasing the size of the fragment until it became a spanning tree. In Boruvka's algorithm, we may have more than one fragment. We increase the size of all fragments by adding the minimum outgoing edge for each fragment.

Algorithm [Boruvka](#) presents the sequential Boruvka's algorithm to find the MST. We use T to denote the set of tree edges. Initially, T is empty. When we determine the components in (V, T) , we find that there are n components, since each vertex is a component by itself when T is empty. The algorithm finds the minimum-weight outgoing edge for each component as follows. In any iteration, we use BFS to find the least numbered vertex to which any vertex is connected in the graph (V, T) . This vertex serves as the identifier for the component of the node i , and we use the variable $cid[i]$ to store it. Once we have determined the component identity of all nodes, we move to the next step of determining the minimum weight outgoing edge for each component. We traverse all edges, and for each edge that connects two different components, we check whether it is cheaper than the previously known outgoing edge for the component on either side. Once we have determined all minimum weight edges for each component, we add these to T and start the next iteration.

Example 8.7 Consider the graph in Fig. 8.1. Initially, T is empty and there are 5 components. We compute mwe for each component and get the edges 4, 3, 3, 2, 2 as the minimum weight edges of a, b, c, d, e , respectively. Once these edges are added, we have two components: $\{a, b, c\}$ and $\{d, e\}$. We then find mwe of these two components as the edge 7. On adding this edge, we have chosen $(n - 1)$ edges, and the algorithm terminates with the edges $\{2, 3, 4, 7\}$.

After every iteration, the number of connected components in (V, T) is reduced by at least a factor of two. This is because every component gets attached to some other component. The worst case is when every least weight edge chosen by any component is also chosen as the least weight edge by the component on the other side of the edge. Hence, the algorithm takes at most $O(\log n)$ iterations of the while loop. It

Algorithm Boruvka: Finding Minimum Spanning Tree using Boruvka's Algorithm

Input: Undirected *connected* Weighted Graph: (V, E, w)
Output: Minimum Weight Spanning Tree T

```

1  $T$ : { set of edges } initially {};
2  $cid$ : array[1.. $n$ ] of 0.. $n$  initially all 0;
3 while  $|T| < n - 1$  do
4    $visited$ : array[1.. $n$ ] of boolean initially all false;
5    $mwe$ : array[1.. $n$ ] of edge initially all null;
6    $dist$ : array[1.. $n$ ] of real initially all  $\infty$ ;
7   for  $i := 1$  to  $n$  do
8     if  $(\neg visited[i])$  then // do a BFS in the graph  $(V, T)$  from vertex  $i$  setting  $cid$  of every
9        $BFS(i)$ ; ;
10  end
11  for  $(i, j) \in E$  such that  $(cid[i] \neq cid[j])$  do
12    if  $w[i, j] < dist[cid[i]]$  then
13       $dist[cid[i]] = w[i, j]$ ;
14       $mwe[cid[i]] = (i, j)$ ;
15    end
16    if  $w[i, j] < dist[cid[j]]$  then
17       $dist[cid[j]] = w[i, j]$ ;
18       $mwe[cid[j]] = (i, j)$ ;
19    end
20  end
21  forall  $i$  do
22     $T := T \cup mwe[cid[i]]$ 
23  end
24 end
25 return  $T$ ;

```

is easy to see that the least weight edge outgoing from each component is found in $O(m)$ work. Thus, the algorithm takes $O(m \log n)$ work.

8.7 Comparison of Algorithms

All three algorithms of this chapter run in $O(m \log n)$ time, but they differ substantially in the data structures they rely on, the input formats they prefer, and the kind of parallelism they expose. This section summarises those practical trade-offs so that a reader faced with a concrete instance can pick the right algorithm.

Input representation. Kruskal’s algorithm is most natural when the edges are already available as a sorted list — the only operations it performs on the graph are “scan the next edge” and “are these two endpoints already connected”. In particular, Kruskal’s is well suited to streaming settings, where edges arrive one at a time in weight order and the graph need not be kept in memory. Prim’s algorithm, on the other hand, needs random access to each vertex’s neighbours and is the natural choice when the graph is represented as an adjacency list. Boruvka’s algorithm operates on both edge and vertex sets and can be implemented with either representation.

Graph density. When the graph is very dense ($m = \Theta(n^2)$), the $O(m \log n)$ bound of all three algorithms is dominated by the edges themselves. Prim’s algorithm implemented with an array-based scan (rather than a heap) achieves $O(n^2)$ work, which beats $O(m \log n) = O(n^2 \log n)$ in this regime. For sparse graphs ($m = O(n)$), all three algorithms are within a $\log n$ factor of linear, and the $O(n^2)$ variant of Prim’s becomes strictly worse than Kruskal’s.

Required auxiliary structures. Kruskal’s algorithm requires a union-find structure to test cycle-closing in $O(\alpha(n))$ time per query, plus a sort on the edge weights. Prim’s algorithm requires a priority queue keyed on the cost-to-fragment of each fringe vertex. Boruvka’s algorithm, by contrast, needs neither a global sort nor a priority queue: each round is a parallel scan of the edges followed by a connected-component relabelling.

Parallelism. This is where the three algorithms diverge the most. Kruskal’s algorithm is inherently sequential: edges must be committed in weight order so that the cut property can be applied, and each commit may reject its predecessors in the sort. Prim’s algorithm is also essentially sequential because it grows one fragment at a time from a single root, though modest parallelism is available in the edge-relaxation step. Boruvka’s algorithm, finally, is the parallelism-friendly one: every component picks its minimum outgoing edge independently in each round, and the whole round can be carried out in $O(\log n)$ time with $O(m)$ processors. This is why the parallel MST algorithms in the literature — and the LLP-Boruvka algorithm of the next section — are all Boruvka variants.

8.8 LLP Perspective

In this section we collect the LLP reformulations of the algorithms introduced earlier in the chapter.

LLP-Kruskal Algorithm

We now give an LLP version for Kruskal's algorithm based on the simple Boolean lattice of all edges. Our distributive lattice is the Boolean lattice formed from all edges. We assume that the edges are given to us in increasing order. Then, an edge e_j between vertices v_k and v_l is in the minimum spanning tree iff there is no path between vertices v_k and v_l using edges $e_1 \dots e_{j-1}$. Note that when j is 1, e_j corresponds to the lightest edge and is always in the minimum spanning tree. An LLP version is presented as Algorithm [LLP-Kruskal](#).

Algorithm LLP-Kruskal: Finding the minimum spanning tree in a graph.

Input: Undirected Weighted Graph: (V, E, w) , edges in increasing order of weights

Output: G : array[1.. m] of $\{0, 1\}$, indicating MST edges

1 **var** G : array[1.. m] of $\{0, 1\}$;

2 **init** $\forall j : G[j] := 0$;

3 **forbidden**(j): ($G[j] = 0$) and there exists no path from v_k to v_l with edges 1.. $j - 1$ for the edge $e_j = (v_k, v_l)$;

4 **advance**(j): $G[j] := 1$;

Example 8.8 Consider the graph in Fig. 8.1. Sorting the edges by weight gives

$$e_1 = (d, e, 2), e_2 = (b, c, 3), e_3 = (a, c, 4), e_4 = (a, b, 5), \\ e_5 = (b, d, 7), e_6 = (c, d, 9), e_7 = (c, e, 11).$$

Starting from $G = (0, 0, 0, 0, 0, 0, 0)$, an index j is forbidden when $G[j] = 0$ and e_j 's endpoints are not yet connected by an edge e_i with $i < j$ and $G[i] = 1$.

- $j = 1$: no prior edges, so d and e are not connected. Advance $G[1] := 1$.
- $j = 2$: b and c are not connected by $\{e_1\}$. Advance $G[2] := 1$.
- $j = 3$: a and c are not connected by $\{e_1, e_2\}$. Advance $G[3] := 1$.
- $j = 4$: a and b are already connected via $a-c-b$ using $\{e_2, e_3\}$. Not forbidden; $G[4]$ stays 0.
- $j = 5$: b and d are not connected by $\{e_1, e_2, e_3\}$. Advance $G[5] := 1$.
- $j = 6$: c and d are connected via $c-b-d$. Not forbidden.
- $j = 7$: c and e are connected via $c-b-d-e$. Not forbidden.

The algorithm terminates with $G = (1, 1, 1, 0, 1, 0, 0)$. The MST consists of the edges e_1, e_2, e_3, e_5 , that is, $(d, e, 2), (b, c, 3), (a, c, 4), (b, d, 7)$, with total weight 16.

LLP-Prim Algorithm

We now show an LLP algorithm to find a minimum spanning tree rooted at a fixed vertex v_0 . As before, we assume that all edge weights are unique. Every node other than v_0 has to choose an edge that we call

the G edge. A node keeps all its edges in the sorted order and begins by choosing its least edge as its G edge. For the graph in Fig. 8.1, with root as the vertex a , we get the sorted adjacency lists shown in Fig. 8.2 for the other vertices.

Vertex	Sorted adjacent edge weights
b	3, 5, 7
c	3, 4, 9, 11
d	2, 7, 9
e	2, 11

Figure 8.2: Sorted adjacency lists for each non-root vertex of the graph in Fig. 8.1, with root a .

We let $G[i]$ be the edge chosen by node i . Thus, initially, the vector G is $\{3, 3, 2, 2\}$. Observe that the set of all possible combinations of edges forms a distributive lattice. Since every node except v_0 has exactly one outgoing edge, following the chosen edge from every node, we either reach v_0 , or end up in a cycle. We define a vertex as *fixed* if traversing the path starting from the edge proposed by that vertex leads to v_0 . The vertex v_0 is trivially fixed. It is clear that the edge proposed by a non-fixed vertex can only lead to a non-fixed vertex, and the edge proposed by a fixed vertex can only lead to a fixed vertex. Thus, any G partitions the set of vertices into *fixed* and *non-fixed* vertices. In our example, the initial vector $\{3, 3, 2, 2\}$ makes the vertex a fixed and all other vertices $\{b, c, d, e\}$ non-fixed.

We define the set of edges between these two partitions as follows.

$$E'(G) := \{(i, k) \in E \mid \text{fixed}(i, G) \wedge \neg \text{fixed}(k, G)\}$$

In our example, we get the following edges as $E'(G) = \{(a, b), (a, c)\}$.

Let $x = (i, j)$ be the edge in E' with minimum $w[i, j]$. For our example, it is (a, c) with an edge weight of 4. We claim that the process j is forbidden in G . Consider any H such that $H[j] < w[i, j]$. Suppose, if possible, that H forms the minimum spanning tree. Any spanning tree must have an edge y between the set $\text{fixed}(G)$ and $\text{non-fixed}(G)$. We claim that H cannot be the minimum spanning tree. This claim holds because the edge set $H - \{y\} + \{x\}$ is also a spanning tree and has a lower weight than H . Therefore, unless the process j advances to the edge (i, j) , G cannot be the minimum spanning tree.

When we advance $G[j]$ to that edge, v_j becomes fixed. Observe that additional nodes may become fixed if their proposed edge leads to v_j directly or indirectly. The algorithm continues to advance G until all vertices become fixed or the edge set $E'(G)$ is empty. In the latter case, the graph is not connected and there is no minimum spanning tree.

Algorithm [LLP-Prim](#) shows the forbidden condition and the advance statement.

Algorithm LLP-Prim: Finding the minimum spanning tree in a graph.

Input: Undirected Weighted Graph: (V, E, w)

Output: G : array[1.. $n - 1$] of real, encoding MST edges

```

1 var  $G$ : array[1.. $n - 1$ ] of real initially  $\forall i : G[i] =$  minimum edge adjacent to  $i$ ;
2 always;
3    $fixed(j, G) \equiv$  there exists a directed path from  $v_j$  to  $v_0$  using edges in  $G$ ;
4    $E' := \{(i, k) \in E \mid fixed(i, G) \wedge \neg fixed(k, G)\}$ ;
5 forbidden( $j$ ):  $\exists i : (i, j) \in E'$  such that it has minimum weight  $w[i, j]$  of all edges in  $E'$ ;
6 advance( $j$ ):  $G[j] := \min\{w[i, j] \mid (i, j) \in E'\}$ ;

```

Example 8.9 Take the graph of Fig. 8.1 with root $v_0 = a$. Each non-root vertex j initializes $G[j]$ to the weight of its lightest adjacent edge, giving $G = (G[b], G[c], G[d], G[e]) = (3, 3, 2, 2)$, i.e., $b \rightarrow c$, $c \rightarrow b$, $d \rightarrow e$, $e \rightarrow d$. Only a is fixed; the other four vertices form two disconnected cycles, so none is fixed.

Iteration 1. The cross-cut set is $E' = \{(a, b, 5), (a, c, 4)\}$, with minimum $(a, c, 4)$, so $j = c$ is forbidden. Advance $G[c] := 4$. Now c is fixed, and b becomes fixed transitively via $b \rightarrow c \rightarrow a$. Fixed = $\{a, b, c\}$; $G = (3, 4, 2, 2)$.

Iteration 2. Now $E' = \{(b, d, 7), (c, d, 9), (c, e, 11)\}$, with minimum $(b, d, 7)$, so $j = d$ is forbidden. Advance $G[d] := 7$. Now d is fixed, and e becomes fixed transitively via $e \rightarrow d \rightarrow b \rightarrow c \rightarrow a$. Fixed = $\{a, b, c, d, e\}$; $G = (3, 4, 7, 2)$.

All vertices are fixed; the algorithm terminates with the MST edges $\{(b, c, 3), (a, c, 4), (b, d, 7), (d, e, 2)\}$, total weight 16.

LLP-Boruvka Algorithm

In this section, we give a recursive version of the algorithm and implement each recursive instance of Boruvka's algorithm with the LLP algorithm. As before, we assume that the input to our algorithm is a connected graph without any self-loops. The LLP algorithm uses the single variable G . For each instance, we let the minimum weight edge (mwe) adjacent to each vertex v be denoted by $mwe[v]$. We denote this directed graph by H : the set of vertices is V and the set of edges is $\{(v, w) \mid (v, w) = mwe[v]\}$. We initialize $G[v]$ with the node w when $mwe[v] = (v, w)$ except when $mwe[v] = (w, v)$ and $v < w$. For the latter case, we make $G[v]$ equal to v . As a result of this initialization, the structure $G[v]$ forms a rooted tree. (The same initialization underlies the parallel Boruvka algorithm of Section 8.6.) We say that the edge (v, w) is *incident to v* when $G[v]$ equals w for w different from v .

Once we have rooted trees for the graph, we convert the rooted trees to rooted stars using pointer-jumping. A node j is considered forbidden if $G[j] \neq G[G[j]]$. It is advanced by setting $G[j]$ to $G[G[j]]$. The algorithm stops this iteration when no node is forbidden. We show that each instance is indeed an LLP algorithm. First note that $G[v]$ is always reachable from v in the directed graph H . Initially, this claim holds because $G[v]$ is set to w where (v, w) is an edge in H . The only statement executed in the program is $G[v] := G[G[v]]$ which preserves the claim.

We also observe that H is a collection of rooted trees and if w is reachable from v , then there is a unique path from v to w . Furthermore, the weight of edges along any path is strictly decreasing. This is because if v points to w and w points to u then the weight of the edge (w, u) must be strictly smaller than

Algorithm LLP-Boruvka: Finding MST

Input: Undirected *connected* Weighted Graph: (V, E, w)

Output: Minimum Weight Spanning Tree T

```

1 function Boruvka( $V, E$ );
2 var  $G$ : array[1.. $n$ ] of 1.. $n$ ;
3 var  $G[v]$ : int initially  $w$  such that  $mwe[v] = (v, w)$  and  $mwe[w] \neq (v, w)$ , or
    $mwe[v] = mwe[w] = (v, w)$  and  $v < w$ ;  $v$ , otherwise.
4 forbidden( $j$ ):  $G[j] \neq G[G[j]]$ ;
5 advance( $j$ ):  $G[j] := G[G[j]]$ ;
6 if  $|V| = 1$  then return  $\{\}$ ;
7 else return  $T \cup \text{Boruvka}(V', E')$  where;
8  $T := \{(v, w) \mid (v, w) \text{ is the minimum weight edge of } v\}$ ;
9  $V' := \{v \in V \mid G[v] = v\}$ ;
10  $E' := \{(G[v], G[w]) \mid ((v, w) \in E) \wedge (G[v] \neq G[w])\}$ ;

```

the weight of the edge (v, w) by the property that each vertex points to the minimum weight edge incident to it. We are now ready to show the following lemma.

Lemma 8.10 *Each instance of finding connected components is an LLP algorithm that terminates.*

Proof: We consider the lattice of vectors. For component j , we keep the weight of the minimum weight edge in the path from j to $G[j]$. We define the predicate B as

$$B \equiv \forall j : G[j] = G[G[j]]$$

We show that the predicate is lattice-linear. Suppose the predicate B is not true. This implies that there exists j such that $G[j]$ is different from $G[G[j]]$. We show that the component j is forbidden. $G[j]$ cannot be the root of a tree in H because the root points to itself (and therefore $G[j]$ equals $G[G[j]]$). Let k be equal to $G[j]$. We just showed that k is not equal to the root. Therefore, $G[k] \neq k$ even if $G[k]$ changes. Hence, $G[j] \neq G[G[j]]$ continues to hold. The component j in G vector is advanced by setting $G[j]$ to be $G[G[j]]$. ■

Algorithm [LLP-Boruvka](#) shows the entire algorithm. Observe that there are no synchronization requirements for any instance of the LLP algorithm. Any thread j that finds $G[j]$ different from $G[G[j]]$ can set it to $G[G[j]]$. In particular, $G[G[j]]$ may have changed between the observation of $(G[j] \neq G[G[j]])$ and setting of $G[j]$ to $G[G[j]]$, but the algorithm still works correctly.

Example 8.11 Consider the graph in Fig. 8.1. The minimum-weight adjacent edge of each vertex is

$$mwe[a] = (a, c, 4), \quad mwe[b] = (b, c, 3), \quad mwe[c] = (b, c, 3), \quad mwe[d] = (d, e, 2), \quad mwe[e] = (d, e, 2).$$

Vertices b and c share the same minimum edge (b, c) , as do d and e via (d, e) . Breaking each 2-cycle in favor of the smaller-numbered vertex, the initialization gives

$$G[a] = c, \quad G[b] = b, \quad G[c] = b, \quad G[d] = d, \quad G[e] = d,$$

which corresponds to two rooted trees: $\{a \rightarrow c \rightarrow b\}$ rooted at b , and $\{e \rightarrow d\}$ rooted at d . The contributed MST edges so far are $\{(a, c, 4), (b, c, 3), (d, e, 2)\}$.

Pointer jumping. Only a is forbidden, since $G[a] = c \neq b = G[G[a]]$. Advancing sets $G[a] := b$. Now every vertex points directly to its root: $G = (b, b, b, d, d)$, two rooted stars $\{a, b, c\} \rightarrow b$ and $\{d, e\} \rightarrow d$.

Recursive call. Contract each star to a single vertex, leaving the two-vertex graph $V' = \{b, d\}$ with the cross-star edges $(b, d, 7), (c, d, 9), (c, e, 11)$ reprojected onto (b, d) as weights 7, 9, 11. Both components' minimum outgoing edge is $(b, d, 7)$, yielding initialization $G[b] = b, G[d] = b$ — already a rooted star. The edge $(b, d, 7)$ is added, and the recursion terminates with a single vertex.

The final MST is $\{(d, e, 2), (b, c, 3), (a, c, 4), (b, d, 7)\}$, total weight 16.

8.9 Summary

Three classical algorithms compute a minimum spanning tree, each exploiting the cut and cycle properties in a different way. *Kruskal's* algorithm scans edges in non-decreasing weight order and admits each edge that does not close a cycle; the find-union data structure reduces the cycle check to near-constant amortized time. *Prim's* algorithm grows a single fragment from an arbitrary root, repeatedly adding the lightest edge that leaves the fragment — an instance of the nearest-neighbor-next strategy. *Boruvka's* algorithm extends all fragments in parallel: in each iteration, every current component contributes its minimum outgoing edge, so the number of components is at least halved, and the algorithm terminates in $O(\log n)$ iterations.

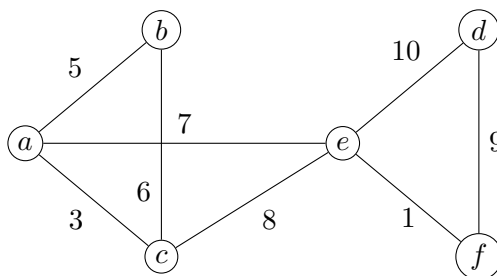
Table 8.1 summarizes the time complexity of the sequential algorithms discussed in this chapter.

Problem	Algorithm	Time Complexity
Minimum Spanning Tree	Kruskal	$O(m \log n)$
Minimum Spanning Tree	Prim	$O(m \log n)$
Minimum Spanning Tree	Boruvka	$O(m \log n)$
Minimum Spanning Tree	LLP-Kruskal	$O(m \log n)$
Minimum Spanning Tree	LLP-Prim	$O(m \log n)$
Minimum Spanning Tree	LLP-Boruvka	$O(m \log n)$

Table 8.1: Algorithms discussed in this chapter.

8.10 Problems

1. Consider the undirected weighted graph on six vertices $\{a, b, c, d, e, f\}$ shown below.



Find the minimum spanning tree using Kruskal's algorithm, Prim's algorithm, and Boruvka's algorithm.

2. Show that there is a unique minimum spanning tree in any weighted undirected graph when all edge weights are distinct.
3. Show that replacing every edge weight $w(e)$ by $-w(e)$ and then running Kruskal's (or Prim's) algorithm yields a *maximum* weight spanning tree of the original graph.
4. A *bottleneck spanning tree* of a weighted graph is a spanning tree that minimizes the weight of its heaviest edge. Show that every minimum spanning tree is also a bottleneck spanning tree.
5. Let T be the MST of an undirected weighted graph $G = (V, E, w)$. Suppose the weight of exactly one edge $e = (u, v) \in E$ is decreased. Describe an $O(n)$ -time algorithm to update T (whether or not e was in the original MST), and justify its correctness.
6. An undirected weighted graph G has a set $F \subseteq E$ of *excluded* edges that must not appear in the spanning tree. Adapt Kruskal's algorithm to compute a minimum spanning tree of $(V, E \setminus F)$ (if one exists) and state a necessary and sufficient condition on F for a feasible spanning tree to exist.
7. Both Prim's algorithm and Dijkstra's algorithm repeatedly pick a vertex outside the current frontier whose "cost" is minimum and add it to the frontier. State precisely the *cost* each algorithm minimizes, and use this difference to explain why Prim's algorithm requires only non-negative edge weights to be irrelevant to correctness (it works for any real weights), whereas Dijkstra's requires non-negative edge weights.
8. Show that in Boruvka's algorithm, regardless of implementation, the number of connected components strictly decreases by at least a factor of two after each iteration. Deduce that the algorithm terminates in at most $\lceil \log_2 n \rceil$ iterations.
9. Assume all edge weights are distinct. Let T be the MST of $G = (V, E, w)$ and let $e \in T$ be the *heaviest* edge in T . Removing e from T partitions V into two components S and $V \setminus S$. Prove that e is the *lightest* edge crossing the cut $(S, V \setminus S)$ in G .

8.11 Bibliographic Remarks

The minimum spanning tree problem has an unusually long and well-documented history. Otakar Borůvka published the first MST algorithm in 1926 in connection with an electrical-grid layout problem in Moravia [Bor26]; the paper also contains the first proof that the MST is unique when edge weights are distinct. A detailed historical survey and an English translation of Borůvka's original papers is given

by Nešetřil, Milková, and Nešetřilová [NMN01]. The classical sequential algorithms bearing the names of Kruskal [Kru56] and Prim [Pri57] were developed independently in the 1950s. Prim's algorithm was in fact discovered earlier by Vojtěch Jarník in 1930, and later independently by Dijkstra [Dij59] as a specialisation of his shortest-path algorithm — a coincidence that is not accidental: the two algorithms have the same control structure and differ only in the relaxation rule (see Section 8.5).

The efficient implementation of Kruskal's algorithm hinges on the union-find data structure. Tarjan [Tar75] proved the near-constant $O(\alpha(n))$ amortized bound for union-find with path compression and union by rank used in Algorithms [Find](#) and [Union](#). An $O(m + n \log n)$ variant of Prim's algorithm, based on Fibonacci heaps, is due to Fredman and Tarjan [FT87]. For a textbook treatment of the three sequential algorithms and their complexity, see Cormen et al. [CLRS01] or Kleinberg and Tardos [KT06a].

A line of subsequent research has pushed the sequential complexity below $O(m \log n)$. Karger, Klein, and Tarjan [KKT95] gave a randomized algorithm running in expected $O(m)$ time — the first linear-time MST algorithm in any model. On the deterministic side, Chazelle [Cha00] obtained $O(m \alpha(m, n))$, inverse-Ackermann in the edge count, using an intricate soft-heap data structure. Pettie and Ramachandran [PR02] then gave an optimal algorithm whose running time matches the minimum number of comparisons needed to decide any MST instance, though its exact asymptotic complexity remains open.

In the parallel and distributed setting, the textbook of Sanders, Mehlhorn, Dietzfelbinger, and Dementiev [SMDD19] is our reference for the parallel Borůvka algorithm of Section 8.6. The pointer-jumping contraction technique that lies at its heart was popularised for parallel graph algorithms by Tarjan and Vishkin [TV85]. Practical shared-memory implementations for sparse graphs are described by Bader and Cong [BC06]. The classical distributed MST algorithm of Gallager, Humblet, and Spira [GHS83] computes an MST in a message-passing model with $O(n \log n)$ message complexity.

The LLP formulations presented in this chapter recast the three classical algorithms as instances of a single lattice-linear predicate framework. The LLP-Prim algorithm in particular is due to Alves and Garg [AG22], where the parallel early-fixing variant is also developed.

Chapter 9

Divide and Conquer

9.1 Introduction

Divide-and-conquer is one of the oldest and most widely used algorithm-design strategies. The idea is to break a problem into a small number of independent sub-problems of the same kind, solve the sub-problems recursively, and then combine their solutions to produce the solution of the original problem. The strategy predates modern computer science: von Neumann's merge sort (1945) is already a textbook example, and the approach underlies algorithms in virtually every area of computing — sorting, searching, numerical computation, computational geometry, and signal processing.

Three features make divide-and-conquer particularly attractive. First, the recursive structure often yields surprisingly tight complexity analyses through the Master Theorem (Section 9.2). Second, because the sub-problems are independent of each other, every divide-and-conquer algorithm is inherently parallel: the recursive calls can be carried out simultaneously on independent processors. Third, and more subtly, decomposing a problem into smaller sub-problems sometimes exposes algebraic identities — as in Karatsuba's multiplication and Strassen's matrix multiplication — that lower the asymptotic complexity below what naive approaches can achieve.

We have already seen two instances of this pattern in Chapter 4: Quicksort, which partitions an array about a pivot and recurses on each part, and Mergesort, which splits the array in half and merges the sorted halves. In both cases the two sub-problems are independent of each other, which makes the approach naturally parallel — a theme that recurs throughout this chapter.

Viewing a problem as finding an appropriate element in a distributive lattice, the divide-and-conquer approach corresponds to finding two appropriate elements in two sub-lattices independently (and in parallel), then combining them to find the required element in the original lattice. The notion of a splittable predicate, which formalises this idea, is defined in Section 2.6 of Chapter 2.

This chapter is organized as follows. Section 9.2 gives the Master Theorem for analysing the time complexity of divide-and-conquer algorithms. Section 9.3 discusses the problem of determining nearest neighbors in the Euclidean plane. Section 9.4 discusses the problem of counting inversions in an array. Section 9.5 describes a solution to the problem of computing the planar convex hull of a set of points. Section 9.6 describes Karatsuba's method for multiplying two natural numbers using the divide-and-conquer approach. Section 9.7 gives Strassen's method for matrix multiplication. Section 9.8 presents

the Fast Fourier Transform and its application to polynomial multiplication. Section 9.9 discusses LLP algorithms based on divide and conquer.

9.2 The Master Theorem

When we apply the divide-and-conquer paradigm, the time complexity of the algorithm is usually analysed through a recurrence relation on the input size. The Master Theorem solves a broad family of such recurrences in closed form.

Theorem 9.1 (Master Theorem) *Let $a \geq 1$, $b > 1$, and $c \geq 0$, and suppose that $T(n) = aT(n/b) + O(n^c)$ with $T(1) = O(1)$. Then*

$$T(n) = \begin{cases} O(n^{\log_b a}) & \text{if } c < \log_b a, \\ O(n^c \log n) & \text{if } c = \log_b a, \\ O(n^c) & \text{if } c > \log_b a. \end{cases}$$

Here a is the number of sub-problems, b is the factor by which the sub-problem size shrinks, and $O(n^c)$ is the cost of work done outside the recursive calls.

Proof: Expanding the recurrence k times gives

$$T(n) = a^k T(n/b^k) + \sum_{i=0}^{k-1} O(a^i (n/b^i)^c) = a^k T(n/b^k) + n^c \sum_{i=0}^{k-1} r^i,$$

where $r = a/b^c$. The recursion terminates at $k = \log_b n$, where $a^k = n^{\log_b a}$ and $T(1) = O(1)$, so the leaf contribution is $O(n^{\log_b a})$. The remaining geometric sum in r is handled by three cases.

- If $c < \log_b a$ then $r > 1$; the sum is $\Theta(r^k)$, and $n^c r^k = n^{\log_b a}$, so $T(n) = O(n^{\log_b a})$ (work is concentrated at the leaves).
- If $c = \log_b a$ then $r = 1$; the sum is $\Theta(k) = \Theta(\log n)$, so $T(n) = O(n^c \log n)$ (work is balanced across levels).
- If $c > \log_b a$ then $r < 1$; the sum is $\Theta(1)$, so the root cost n^c dominates and $T(n) = O(n^c)$.

Substituting these bounds into the expansion gives the three cases of the theorem. ■

Fig. 9.1 illustrates the proof for Merge Sort, where $a = b = 2$ and $c = 1$. Every level of the recursion tree costs n (four levels for $n = 8$), and the $1 + \log_2 n$ levels together give $T(n) = O(n \log n)$. This is the balanced case $c = \log_b a$ of the theorem; in the other two cases the level costs form a geometric sequence rather than being constant, and either the leaves or the root dominate.

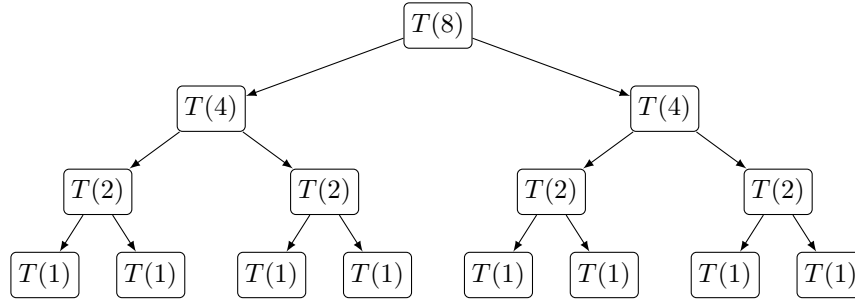


Figure 9.1: Recursion tree for $T(n) = 2T(n/2) + O(n)$ with $n = 8$. Every level costs $n = 8$, and there are $1 + \log_2 n = 4$ levels, giving $T(n) = O(n \log n)$.

Applications

We apply the Master Theorem to several recurrences that arise elsewhere in the book.

1. *Merge Sort*. $T(n) = 2T(n/2) + O(n)$: $a = 2, b = 2, c = 1 = \log_2 2$; case 2 gives $T(n) = O(n \log n)$.
2. *Binary Search*. $T(n) = T(n/2) + O(1)$: $a = 1, b = 2, c = 0 = \log_2 1$; case 2 gives $T(n) = O(\log n)$.
3. *Strassen's matrix multiplication*. $T(n) = 7T(n/2) + O(n^2)$: $a = 7, b = 2, c = 2 < \log_2 7 \approx 2.81$; case 1 gives $T(n) = O(n^{2.81})$.
4. *Karatsuba's algorithm*. $T(n) = 3T(n/2) + O(n)$: $a = 3, b = 2, c = 1 < \log_2 3 \approx 1.58$; case 1 gives $T(n) = O(n^{1.58})$.
5. *Case 3 example*. $T(n) = 2T(n/2) + O(n^2)$: $a = 2, b = 2, c = 2 > \log_2 2 = 1$; case 3 gives $T(n) = O(n^2)$.
6. *Case 1 example*. $T(n) = 9T(n/3) + O(n)$: $a = 9, b = 3, c = 1 < \log_3 9 = 2$; case 1 gives $T(n) = O(n^2)$.
7. *Case 2 example*. $T(n) = T(2n/3) + O(1)$: $a = 1, b = 3/2, c = 0 = \log_{3/2} 1$; case 2 gives $T(n) = O(\log n)$.

9.3 Nearest Neighbors in the Euclidean Space

We are given n points in the two-dimensional Euclidean space. Our goal is to find a pair of points with the smallest distance between them. That is, determine two points (p_i, p_j) such that the Euclidean distance between them is minimized:

$$d(p_i, p_j) = \sqrt{(x_i - x_j)^2 + (y_i - y_j)^2}$$

The easiest algorithm would be to compute the distance between every pair of points and choose a pair with the least distance. This approach requires $O(n^2)$ sequential time. Our goal is to reduce the sequential time complexity for this problem.

The divide-and-conquer approach for this problem is as follows. We divide the set of points into two sets S_1 and S_2 . Suppose that we have the nearest neighbors in S_1 and S_2 . Now, if we had nearest neighbors such that one point is in S_1 and the second point is in S_2 , then we simply have to choose the smallest of these three distances: nearest neighbors in S_1 , nearest neighbors in S_2 , and nearest neighbors across the partition. Suppose that the distance of the nearest neighbors in S_1 , or in S_2 is δ . Then, we only need to consider the nearest neighbors across the partition if they are closer than δ . Let S_y be the set of points within δ of the dividing line sorted in the y -coordinate. We claim that if $s, s' \in S$ are such that $d(s, s') < \delta$,

then s and s' are within 15 positions of each other in the sorted list S_y . Let L be the vertical line that divides the set of points according to the coordinate x . We consider boxes of size $\delta/2$ around L . We first show that there cannot be two points from the same side within a box. The largest distance within a box is $\sqrt{2}(\delta/2)$. This is equal to $\delta/\sqrt{2}$, which is less than δ .

We next claim that there cannot be two points that are sixteen positions apart in S_y and still have distance less than δ . Since the points are sixteen positions apart and at most one point in each box, we get that there are at least three rows of boxes between these two points. However, this means that the distance between them is at least $3\delta/2$, which is greater than δ .

Hence, it is sufficient to consider points that are at most 15 positions apart in S_y .

Algorithm ClosestPair: Closest Pair of Points

Input: Set of n points P in 2D space

Output: The closest pair of points

```

1 Function ClosestPair( $P$ ):
2   if  $n \leq 3$  then
3     | return BruteForce( $P$ ) // Base case: use brute-force for small input
4   end
5   Sort  $P$  by  $x$ -coordinate;
6   Split  $P$  into left ( $P_L$ ) and right ( $P_R$ ) halves;
7    $(p_1, p_2) :=$  ClosestPair( $P_L$ );
8    $(p_3, p_4) :=$  ClosestPair( $P_R$ );
9    $\delta := \min(d(p_1, p_2), d(p_3, p_4))$ ;
10   $S :=$  points within  $\delta$  of the dividing line;
11  Sort  $S$  by  $y$ -coordinate;
12  for each point  $p$  in  $S$  do
13    | for each of the next 15 points  $q$  do
14      | | if  $d(p, q) < \delta$  then
15        | | | Update  $\delta$  and closest pair;
16      | | end
17    | end
18  end
19  return closest pair;

```

Example 9.2 Consider the set of points

$$P = \{(1, 2), (3, 7), (5, 4), (8, 9), (9, 6), (10, 3)\}.$$

Sorting. Arrange by x -coordinate: $[(1, 2), (3, 7), (5, 4), (8, 9), (9, 6), (10, 3)]$. *Divide.* Split into $P_L = [(1, 2), (3, 7), (5, 4)]$ and $P_R = [(8, 9), (9, 6), (10, 3)]$. *Recurse.* The closest pair in P_L is $\{(3, 7), (5, 4)\}$ with $d = \sqrt{13} \approx 3.61$; in P_R it is $\{(8, 9), (9, 6)\}$ with $d = \sqrt{10} \approx 3.16$. *Combine.* Let $\delta = \min(3.61, 3.16) = 3.16$ and place the dividing line at $x = 5$. Only $(3, 7), (5, 4), (8, 9)$ lie within δ of it; the pair $(9, 6)$ is outside the strip since $|9 - 5| = 4 > 3.16$. Checking strip pairs (sorted by y -coordinate) yields distances $\sqrt{13}, \sqrt{34}, \sqrt{29}$, none improving on δ . *Output.* The closest pair is $\{(8, 9), (9, 6)\}$ at distance $\sqrt{10} \approx 3.16$.

The time complexity is as follows. For sorting the points, we spend $O(n \log n)$ time. To recursively solve subproblems, we use $O(n \log n)$ time. For checking points in the strip we spend $O(n)$ time. Thus, the overall time complexity is $O(n \log n)$.

9.4 Counting Inversions in an Array Using Divide and Conquer

Given an array $A[1 \dots n]$, an **inversion** is a pair of indices (i, j) such that $i < j$ and $A[i] > A[j]$. The goal is to count the total number of inversions in the array efficiently. A brute-force approach iterates through all pairs and checks if $A[i] > A[j]$, requiring $O(n^2)$ time. Using Merge Sort with an additional counting step, we can solve this problem in $O(n \log n)$ time.

We modify the *Merge Sort* algorithm to count inversions efficiently: (1) Recursively count inversions in the left and right halves. (2) Count the number of inversions across the two halves while merging. (3) Sum up all these counts.

Algorithm CountingInversions: Finding the number of inversions in an array.

```

Input: Array  $A$  of size  $n$ 
Output: Total number of inversions
1 Function CountInversions( $A, left, right$ ):
2   if  $left \geq right$  then
3     return 0 // Base case: single element has no inversions
4   end
5    $mid := (left + right) / 2;$ 
6   // Sort the left half and count the inversions
7    $invLeft := \text{CountInversions}(A, left, mid);$ 
8   // Sort the right half and count the inversions
9    $invRight := \text{CountInversions}(A, mid + 1, right);$ 
10   $invMerge := \text{MergeAndCount}(A, left, mid, right);$ 
11  return  $invLeft + invRight + invMerge;$ 
12 Function MergeAndCount( $A, left, mid, right$ ):
13   // left as an exercise

```

Example 9.3 Consider the array $A = [5, 3, 7, 1, 8, 2]$ of size 6. The recursion proceeds as follows.

- Left half $[5, 3, 7]$. Recursing further: $[5, 3]$ contributes the single inversion $(5, 3)$; $[7]$ contributes none. Merging the sorted halves $[3, 5]$ and $[7]$ adds no inversion since every left element is smaller than 7. Left-half total: 1.
- Right half $[1, 8, 2]$. Recursing further: $[1, 8]$ contributes none; $[2]$ contributes none. Merging $[1, 8]$ and $[2]$: $1 \leq 2$ is emitted, then $8 > 2$ contributes the single inversion $(8, 2)$. Right-half total: 1.
- Merging the sorted halves $[3, 5, 7]$ and $[1, 2, 8]$ while counting cross-half inversions: whenever an element from the right half is emitted with left elements still pending, the number of pending left elements counts as inversions. Emitting 1 leaves 3 left elements pending; emitting 2 leaves 3 left elements pending; the remaining comparisons emit from the left. Cross-half total: $3 + 3 = 6$.

The grand total is $1 + 1 + 6 = 8$ inversions. A direct enumeration of pairs (i, j) with $i < j$ and $A[i] > A[j]$ confirms 8: $(5, 3), (5, 1), (5, 2), (3, 1), (3, 2), (7, 1), (7, 2), (8, 2)$.

It is easy to see that the merge sort recursion takes $O(n \log n)$ time. The Merge step inversion counting takes $O(n)$ time. Thus, the overall time complexity is $O(n \log n)$.

9.5 Planar Convex Hull

Suppose that we are given S , a set of points in a plane. The convex hull of S is the smallest convex polygon that contains all points of S . The convex polygon can be described by the ordered list of points (in the clockwise direction) that defines the boundary of the polygon. Our task is to compute the convex hull of S .

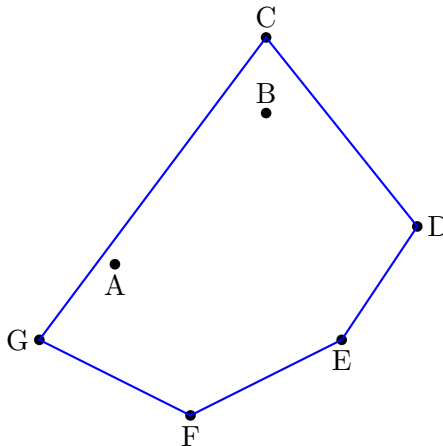


Figure 9.2: An example of Planar Convex Hull

Let points $\{p_1, \dots, p_n\}$ be sorted in their x -coordinate. For simplicity, we assume that x coordinates are unique. It is easy to verify that the point with the smallest x -coordinate, p_1 , and the largest x -coordinate, p_n , are always in the convex hull. These two points also divide up the convex hull into two sets: the upper

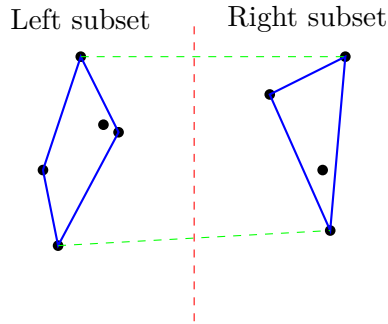


Figure 9.3: Using Divide and Conquer for Planar Convex Hull

hull and the lower hull. The upper hull consists of all points in the convex hull starting from p_1 and ending at p_n (but not including p_n) in the clockwise direction. The lower hull consists of all points starting from p_n and ending at p_1 (but not including p_1).

To compute the convex hull of S , we divide the set of points into two sets S_1 and S_2 such that S_1 has first $n/2$ points with smaller x coordinate and S_2 has $n/2$ points with the larger x coordinate. Similar to mergesort, we can compute convex hulls of S_1 and S_2 . The only task left is to merge the convex hull of S_1 and S_2 to get the convex hull of S . We first get the upper hulls of S_1 and S_2 and merge them to get the upper hull of S . This procedure can be repeated for lower hulls. To merge upper hulls, we need to determine the upper common tangent of $UH(S_1)$ and $UH(S_2)$. It is known that the upper common tangent can be determined sequentially in $O(n)$ time. Hence, we get the following recurrence:

$$T(n) \leq 2T(n/2) + O(n)$$

Solving this recurrence, we get $T(n) = O(n \log n)$.

Example 9.4 Consider the seven points shown in Fig. 9.2: $A(0, 0)$, $B(2, 2)$, $C(2, 3)$, $D(4, 0.5)$, $E(3, -1)$, $F(1, -2)$, $G(-1, -1)$. Sorting by x -coordinate gives G, A, F, B, C, E, D , which splits into $P_L = \{G, A, F, B\}$ and $P_R = \{C, E, D\}$. Recursing on P_L yields the hull G, A, F, B (all four points are on it), and on P_R the hull is the triangle C, E, D . The upper common tangent of the two hulls connects C to B , and the lower common tangent connects E to F (through G, A). Merging along the tangents gives the final hull C, D, E, F, G of five points, exactly the blue polygon of Fig. 9.2. The interior points A and B are discarded at the merge step.

9.6 Karatsuba's Multiplication Algorithm

A notable application of the divide-and-conquer approach is Karatsuba's multiplication algorithm, which computes the product of two large integers faster than the classical grade-school method. Introduced by Anatoly Karatsuba in 1962 [KO62], it reduces the complexity from $O(n^2)$ to $O(n^{\log_2 3}) \approx O(n^{1.585})$, where n is the number of digits.

Consider multiplying two polynomials $A(x) = a_0 + a_1x$ and $B(x) = b_0 + b_1x$, where coefficients are single-digit numbers (e.g., base-10 digits). Their product is:

$$C(x) = A(x) \cdot B(x) = a_0b_0 + (a_0b_1 + a_1b_0)x + a_1b_1x^2.$$

Naive multiplication computes each term $(a_0b_0, a_0b_1, a_1b_0, a_1b_1)$, requiring 4 multiplications and 3 additions, yielding $O(n^2)$ for degree- n polynomials.

Now, represent two n -digit numbers $X = x_1 \cdot 10^{n/2} + x_0$ and $Y = y_1 \cdot 10^{n/2} + y_0$ (assuming n is even), where x_1, x_0, y_1, y_0 are $n/2$ -digit numbers. Their product is:

$$X \cdot Y = (x_1 \cdot 10^{n/2} + x_0)(y_1 \cdot 10^{n/2} + y_0) = x_1y_1 \cdot 10^n + (x_1y_0 + x_0y_1) \cdot 10^{n/2} + x_0y_0,$$

akin to polynomial multiplication with $x = 10^{n/2}$. Naive computation requires 4 multiplications of $n/2$ -digit numbers, suggesting $O(n^2)$ overall. Karatsuba's algorithm optimizes this to 3 multiplications.

Karatsuba's insight reduces the number of multiplications by computing: 1. $p_1 = x_0y_0$, 2. $p_2 = x_1y_1$, 3. $p_3 = (x_0 + x_1)(y_0 + y_1)$.

Then:

$$X \cdot Y = p_2 \cdot 10^n + [p_3 - (p_1 + p_2)] \cdot 10^{n/2} + p_1.$$

$p_3 = x_0y_0 + x_0y_1 + x_1y_0 + x_1y_1 = p_1 + p_2 + (x_1y_0 + x_0y_1)$, Thus, $p_3 - (p_1 + p_2) = x_1y_0 + x_0y_1$.

This requires only 3 multiplications (p_1, p_2, p_3) instead of 4, plus additional additions/subtractions, which are $O(n)$.

For n -digit numbers:

- Split: $X = x_1 \cdot 10^{n/2} + x_0$, $Y = y_1 \cdot 10^{n/2} + y_0$.
- Recursively compute p_1, p_2, p_3 on $n/2$ -digit numbers.
- Combine using the formula above.

The recurrence is:

$$T(n) = 3T(n/2) + O(n),$$

solving to $T(n) = O(n^{\log_2 3})$ via the Master Theorem (Case 1: $a = 3, b = 2, \log_b a \approx 1.585$).

Algorithm Karatsuba: Karatsuba's Multiplication Algorithm

Input: Numbers X, Y , digit length n (power of 2)

Output: Product $X \cdot Y$

```

1 if  $n = 1$  then
2   | return  $X \cdot Y$  // base case: single-digit multiplication
3 end
4  $x_1 := \lfloor X/10^{n/2} \rfloor$ ,  $x_0 := X \bmod 10^{n/2}$  // high and low halves of  $X$ 
5  $y_1 := \lfloor Y/10^{n/2} \rfloor$ ,  $y_0 := Y \bmod 10^{n/2}$  // high and low halves of  $Y$ 
6  $p_1 := \text{Karatsuba}(x_0, y_0, n/2)$  // low product
7  $p_2 := \text{Karatsuba}(x_1, y_1, n/2)$  // high product
8  $p_3 := \text{Karatsuba}(x_0 + x_1, y_0 + y_1, n/2)$  // middle term helper
9 return  $p_2 \cdot 10^n + [p_3 - (p_1 + p_2)] \cdot 10^{n/2} + p_1$ 

```

Example 9.5 Let $X = 1234$, $Y = 5678$, $n = 4$.

Divide. With $10^{n/2} = 100$: $x_1 = 12$, $x_0 = 34$, $y_1 = 56$, $y_0 = 78$.

Recurse. $p_1 = 34 \cdot 78 = 2652$, $p_2 = 12 \cdot 56 = 672$, $p_3 = (34 + 12)(78 + 56) = 46 \cdot 134 = 6164$.

Combine. $p_3 - (p_1 + p_2) = 6164 - 3324 = 2840$, so

$$X \cdot Y = 672 \cdot 10^4 + 2840 \cdot 10^2 + 2652 = 7006652,$$

matching the direct product $1234 \cdot 5678$.

Karatsuba's algorithm reduces multiplications from 4 to 3 per level, trading them for additions/subtractions.

For $n = 2^k$:

- Levels: $k = \log_2 n$,
- Multiplications: $3^k = 3^{\log_2 n} = n^{\log_2 3}$,
- Additions: $O(n \log n)$.

Compared to FFT ($O(n \log n)$), Karatsuba is slower but avoids complex numbers, making it simpler for integer arithmetic. The example shows its practicality for small numbers, scaling efficiently for larger n .

9.7 Strassen's Matrix Multiplication

Strassen's algorithm is a divide-and-conquer approach to matrix multiplication that reduces the time complexity from the naive $O(n^3)$ to $O(n^{\log_2 7}) \approx O(n^{2.807})$ for multiplying two $n \times n$ matrices. Unlike the standard method, which performs 8 recursive multiplications for 2×2 submatrices, Strassen's method uses 7 multiplications by cleverly combining submatrix operations, trading some multiplications for additional additions.

Given two $n \times n$ matrices A and B , where n is a power of 2 (padded with zeros if necessary), Strassen's algorithm divides each matrix into four $n/2 \times n/2$ submatrices:

$$A = \begin{bmatrix} A_{11} & A_{12} \\ A_{21} & A_{22} \end{bmatrix}, \quad B = \begin{bmatrix} B_{11} & B_{12} \\ B_{21} & B_{22} \end{bmatrix}.$$

The product $C = A \cdot B$ is computed as:

$$C = \begin{bmatrix} C_{11} & C_{12} \\ C_{21} & C_{22} \end{bmatrix},$$

where traditionally:

- $C_{11} = A_{11}B_{11} + A_{12}B_{21}$,
- $C_{12} = A_{11}B_{12} + A_{12}B_{22}$,
- $C_{21} = A_{21}B_{11} + A_{22}B_{21}$,
- $C_{22} = A_{21}B_{12} + A_{22}B_{22}$.

Algorithm Strassen: Strassen's Matrix Multiplication

Input: A, B : $n \times n$ matrices, n a power of 2

Output: C : $n \times n$ matrix, $C = A \cdot B$

```

1 if  $n = 1$  then
2   |  $C[1, 1] := A[1, 1] \cdot B[1, 1]$  ;
3   | return  $C$  ;
4 end
5 Divide  $A$  into  $A_{11}, A_{12}, A_{21}, A_{22}$  ;           // Each  $n/2 \times n/2$ 
6 Divide  $B$  into  $B_{11}, B_{12}, B_{21}, B_{22}$  ;           // Each  $n/2 \times n/2$ 
7  $M_1 := \text{Strassen}(A_{11} + A_{22}, B_{11} + B_{22})$  ;
8  $M_2 := \text{Strassen}(A_{21} + A_{22}, B_{11})$  ;
9  $M_3 := \text{Strassen}(A_{11}, B_{12} - B_{22})$  ;
10  $M_4 := \text{Strassen}(A_{22}, B_{21} - B_{11})$  ;
11  $M_5 := \text{Strassen}(A_{11} + A_{12}, B_{22})$  ;
12  $M_6 := \text{Strassen}(A_{21} - A_{11}, B_{11} + B_{12})$  ;
13  $M_7 := \text{Strassen}(A_{12} - A_{22}, B_{21} + B_{22})$  ;
14  $C_{11} := M_1 + M_4 - M_5 + M_7$  ;
15  $C_{12} := M_3 + M_5$  ;
16  $C_{21} := M_2 + M_4$  ;
17  $C_{22} := M_1 - M_2 + M_3 + M_6$  ;
18 return  $C = \begin{bmatrix} C_{11} & C_{12} \\ C_{21} & C_{22} \end{bmatrix}$  ;

```

This requires 8 multiplications and 4 additions of $n/2 \times n/2$ matrices. Strassen's insight is to define 7 intermediate products (M_1 to M_7) that allow the computation of C with fewer multiplications:

The recurrence relation for Strassen's algorithm is:

$$T(n) = 7T(n/2) + O(n^2),$$

where 7 is the number of recursive multiplications, and $O(n^2)$ accounts for the additions/subtractions of $n/2 \times n/2$ matrices. Using the Master Theorem ($a = 7$, $b = 2$, $f(n) = O(n^2)$, $\log_b a = \log_2 7 \approx 2.807 > 2$):

$$T(n) = O(n^{\log_2 7}) \approx O(n^{2.807}).$$

This is an improvement over the standard $O(n^3)$.

Example 9.6 Consider the 2×2 matrices

$$A = \begin{bmatrix} 1 & 2 \\ 3 & 4 \end{bmatrix}, \quad B = \begin{bmatrix} 5 & 6 \\ 7 & 8 \end{bmatrix}.$$

The seven Strassen products evaluate to

$$\begin{aligned} M_1 &= (1 + 4)(5 + 8) = 65, & M_2 &= (3 + 4) \cdot 5 = 35, & M_3 &= 1 \cdot (6 - 8) = -2, \\ M_4 &= 4 \cdot (7 - 5) = 8, & M_5 &= (1 + 2) \cdot 8 = 24, & M_6 &= (3 - 1)(5 + 6) = 22, \\ M_7 &= (2 - 4)(7 + 8) = -30. \end{aligned}$$

Combining them:

$$\begin{aligned} C_{11} &= M_1 + M_4 - M_5 + M_7 = 19, & C_{12} &= M_3 + M_5 = 22, \\ C_{21} &= M_2 + M_4 = 43, & C_{22} &= M_1 - M_2 + M_3 + M_6 = 50. \end{aligned}$$

Hence $C = A \cdot B = \begin{bmatrix} 19 & 22 \\ 43 & 50 \end{bmatrix}$, matching the direct product. The example uses 7 scalar multiplications rather than the textbook 8.

9.8 Fast Fourier Transform

Motivation: polynomial multiplication

Given two polynomials

$$A(x) = \sum_{i=0}^{n-1} a_i x^i \quad \text{and} \quad B(x) = \sum_{j=0}^{n-1} b_j x^j,$$

their product $C(x) = A(x) \cdot B(x) = \sum_{k=0}^{2n-2} c_k x^k$ has coefficients

$$c_k = \sum_{i+j=k} a_i b_j, \quad k = 0, 1, \dots, 2n - 2.$$

Computing every c_k directly from the coefficient vectors (a_0, \dots, a_{n-1}) and (b_0, \dots, b_{n-1}) takes $\Theta(n^2)$ scalar multiplications. Polynomial multiplication is also the building block of integer multiplication (each long integer is a polynomial in the base, evaluated at the base) and of convolution (so the same algorithm computes the discrete convolution of two signals). The Fast Fourier Transform, due to Cooley and Tukey [CT65], reduces the cost to $O(n \log n)$ by exploiting an alternative representation of polynomials.

Two representations of a polynomial

A polynomial of degree at most $n - 1$ can be specified in two equivalent ways:

- the *coefficient representation* $(a_0, a_1, \dots, a_{n-1})$, and
- the *point-value representation* $((x_0, A(x_0)), (x_1, A(x_1)), \dots, (x_{n-1}, A(x_{n-1})))$ at any n distinct points x_0, \dots, x_{n-1} .

Two distinct points determine a unique line, three distinct points determine a unique parabola, and in general n distinct points determine a unique polynomial of degree at most $n - 1$ (this is the Lagrange interpolation theorem).

The coefficient representation makes it cheap to *evaluate* the polynomial at one point ($O(n)$ via Horner's rule) but expensive to multiply ($\Theta(n^2)$). The point-value representation reverses this: evaluation at the chosen sample points is free, multiplication is $O(n)$ (multiply pointwise), but recovering the coefficients of C at the end — the *interpolation* step — looks expensive at first sight.

The FFT's contribution is to choose a special set of evaluation points — the complex roots of unity — so that *both* the coefficient-to-point-value transform and its inverse can be computed in $O(n \log n)$ time.

Roots of unity

Let N be a power of two with $N \geq 2n - 1$ (so that the product polynomial C of degree at most $2n - 2$ is uniquely determined by N point values). The N -th roots of unity are the N complex numbers

$$\omega^0, \omega^1, \omega^2, \dots, \omega^{N-1}, \quad \omega = e^{2\pi i/N}.$$

They have three properties that make them the perfect evaluation set:

1. *Symmetry.* $\omega^{k+N/2} = -\omega^k$ for every k , since $\omega^{N/2} = e^{\pi i} = -1$.
2. *Recursive structure.* $(\omega^k)^2 = \omega^{2k}$, and the squares $\{\omega^{2k} : k = 0, \dots, N/2 - 1\}$ are exactly the $(N/2)$ -th roots of unity, each repeated twice.
3. *Orthogonality.* For $0 \leq j, k < N$, $\sum_{\ell=0}^{N-1} \omega^{\ell(j-k)} = N$ if $j = k$ and 0 otherwise.

Properties (1)–(2) drive the divide-and-conquer recursion below; property (3) provides a closed-form inversion formula.

The DFT and its inverse

The *discrete Fourier transform* of a coefficient vector $a = (a_0, \dots, a_{N-1})$ is the vector $\hat{a} = (\hat{a}_0, \dots, \hat{a}_{N-1})$ of values of A at the N roots of unity:

$$\hat{a}_k = A(\omega^k) = \sum_{j=0}^{N-1} a_j \omega^{jk}, \quad k = 0, 1, \dots, N-1.$$

By the orthogonality property, the *inverse* DFT is given by the same formula with $\omega^{-1} = \bar{\omega}$ in place of ω and a scaling by $1/N$:

$$a_j = \frac{1}{N} \sum_{k=0}^{N-1} \hat{a}_k \omega^{-jk}.$$

Thus, polynomial multiplication can be done as in Algorithm [FFTMultiply](#): forward DFT both inputs, pointwise multiply, inverse DFT.

Algorithm FFTMultiply: Polynomial multiplication via FFT.

Input: Coefficient vectors $a = (a_0, \dots, a_{n-1})$ and $b = (b_0, \dots, b_{n-1})$
Output: Coefficient vector $c = (c_0, \dots, c_{2n-2})$ with $C(x) = A(x)B(x)$

```

1  $N \leftarrow$  smallest power of 2 with  $N \geq 2n - 1$ ;
2 Pad  $a$  and  $b$  with zeros to length  $N$ ;
3  $\hat{a} := \text{FFT}(a, \omega)$ ; // forward DFT,  $\omega = e^{2\pi i/N}$ 
4  $\hat{b} := \text{FFT}(b, \omega)$ ;
5 for  $k = 0$  to  $N - 1$  do
6    $\hat{c}_k := \hat{a}_k \cdot \hat{b}_k$ ; // pointwise multiplication
7 end
8  $c := \text{FFT}(\hat{c}, \omega^{-1})$ , then divide every entry by  $N$ ; // inverse DFT
9 return  $(c_0, c_1, \dots, c_{2n-2})$ ;
```

The recursive FFT

To compute a DFT of size N in $O(N \log N)$ time, split the input by the parity of the index. Write

$$A(x) = A_e(x^2) + x \cdot A_o(x^2),$$

where

$$A_e(y) = \sum_{j=0}^{N/2-1} a_{2j} y^j, \quad A_o(y) = \sum_{j=0}^{N/2-1} a_{2j+1} y^j.$$

Both A_e and A_o have size $N/2$. Evaluating A at the N points ω^k ($k = 0, \dots, N - 1$) reduces to evaluating A_e and A_o at the $N/2$ points ω^{2k} , which are the $(N/2)$ -th roots of unity. The *butterfly identity* below uses the symmetry $\omega^{k+N/2} = -\omega^k$ to fold the answer back together:

$$\hat{a}_k = A_e(\omega^{2k}) + \omega^k A_o(\omega^{2k}), \quad \hat{a}_{k+N/2} = A_e(\omega^{2k}) - \omega^k A_o(\omega^{2k}).$$

Algorithm FFT: Recursive Fast Fourier Transform.

Input: Vector $a = (a_0, a_1, \dots, a_{N-1})$ with N a power of 2; primitive root ω

Output: Vector $\hat{a} = (\hat{a}_0, \dots, \hat{a}_{N-1})$ with $\hat{a}_k = \sum_j a_j \omega^{jk}$

```

1 if  $N = 1$  then
2   | return  $(a_0)$ 
3 end
4  $a^{(e)} := (a_0, a_2, \dots, a_{N-2});$    $a^{(o)} := (a_1, a_3, \dots, a_{N-1});$ 
5  $\hat{a}^{(e)} := \text{FFT}(a^{(e)}, \omega^2);$  // recurse on even-indexed
6  $\hat{a}^{(o)} := \text{FFT}(a^{(o)}, \omega^2);$  // recurse on odd-indexed
7  $z := 1;$  // the running power  $\omega^k$ 
8 for  $k = 0$  to  $N/2 - 1$  do
9   |  $\hat{a}_k := \hat{a}_k^{(e)} + z \cdot \hat{a}_k^{(o)};$ 
10  |  $\hat{a}_{k+N/2} := \hat{a}_k^{(e)} - z \cdot \hat{a}_k^{(o)};$ 
11  |  $z := z \cdot \omega;$ 
12 end
13 return  $(\hat{a}_0, \hat{a}_1, \dots, \hat{a}_{N-1});$ 

```

The recurrence is $T(N) = 2T(N/2) + O(N)$, which by the Master Theorem (Theorem 9.1, balanced case) yields $T(N) = O(N \log N)$. The same algorithm computes the inverse DFT when invoked with ω^{-1} in place of ω , after which every output entry is divided by N .

Why this beats $\Theta(n^2)$

The naive coefficient-multiplication algorithm uses $\Theta(n^2)$ scalar multiplications and additions because each output coefficient depends on a different subset of input pairs. By moving to point-value representation, multiplication becomes pointwise — only N scalar multiplications — and *all* the work is funnelled into the two DFTs and one inverse DFT. The roots of unity are the unique evaluation set whose recursive structure makes those DFTs themselves cheap.

Example 9.7 Let $A(x) = 1 + 2x$ and $B(x) = 1 + 3x$. The direct product is $C(x) = 1 + 5x + 6x^2$. We use $N = 4$, $\omega = i$, so the four roots of unity are $\{1, i, -1, -i\}$. Forward DFTs:

k	0	1	2	3
ω^k	1	i	-1	$-i$
$A(\omega^k)$	3	$1 + 2i$	-1	$1 - 2i$
$B(\omega^k)$	4	$1 + 3i$	-2	$1 - 3i$
$C(\omega^k)$	12	$-5 + 5i$	2	$-5 - 5i$

Inverse DFT: $c_k = \frac{1}{4} \sum_{\ell=0}^3 C(\omega^\ell) \omega^{-k\ell}$. For instance, $c_0 = \frac{1}{4}(12 + (-5 + 5i) + 2 + (-5 - 5i)) = \frac{4}{4} = 1$, and similarly $c_1 = 5$, $c_2 = 6$, $c_3 = 0$. We recover $C(x) = 1 + 5x + 6x^2$, agreeing with the direct computation.

To see Algorithm FFT concretely, consider the forward DFT of $A = (1, 2, 0, 0)$. The split is $a^{(e)} = (1, 0)$ and $a^{(o)} = (2, 0)$. Recursing on each (with $\omega^2 = -1$): the size-2 DFT of $(1, 0)$ is $(1, 1)$ and of $(2, 0)$ is $(2, 2)$. The butterfly with $z = 1, i$ then gives

$$\hat{a}_0 = 1 + 1 \cdot 2 = 3, \quad \hat{a}_1 = 1 + i \cdot 2 = 1 + 2i, \quad \hat{a}_2 = 1 - 1 \cdot 2 = -1, \quad \hat{a}_3 = 1 - i \cdot 2 = 1 - 2i,$$

matching the table above.

The FFT is one of the most widely used algorithms in scientific computing: it underlies fast integer multiplication (Schönhage–Strassen), large-scale signal processing (where the DFT converts time-domain samples to frequency-domain amplitudes), and numerical solution of partial differential equations.

9.9 LLP Perspective

Viewing the problem as finding an appropriate element in a distributive lattice, the divide-and-conquer approach corresponds to viewing the problem as finding two appropriate elements in two distributive lattices. This search is carried out independently (and in parallel). Once these two elements are determined, we use them to find the required element in the original lattice.

The notion of a splittable predicate is defined in Section 2.6 of Chapter 2. Quicksort and Mergesort cast as splittable predicates are revisited in Chapter 4. In this section we give LLP formulations for the remaining divide-and-conquer problems of this chapter. Each formulation recovers the correct answer as the least fixed point of a forbidden/advance rule; none of them matches the asymptotic running time of the sharpest divide-and-conquer algorithm, but each exposes the parallelism in the problem and gives a uniform template that the parallel scheduler can exploit.

Nearest Neighbors in the Euclidean Space

For each point p_j we wish to know the distance from p_j to its nearest neighbour in the remaining set. Let G be a vector of n real numbers ordered by reverse comparison (smaller values are greater in the lattice). We interpret $G[j]$ as an upper bound on the distance from p_j to its nearest neighbour. The least fixed point has $G[j] = \min_{k \neq j} d(p_j, p_k)$ for every j , from which the global nearest-pair distance is read off as

$\min_j G[j]$.

Algorithm LLP-NearestNeighbor: LLP formulation of the nearest-neighbour problem.

Input: Points $p_1, \dots, p_n \in \mathbb{R}^2$

Output: G : array[1.. n] of real; $G[j]$ = distance from p_j to its nearest neighbour

1 **var** G : array[1.. n] of real;

2 **init** $\forall j : G[j] := \infty$;

3 **forbidden**(j): $\exists k \neq j : d(p_j, p_k) < G[j]$;

4 **advance**(j): $G[j] := \min_{k \neq j} d(p_j, p_k)$;

Correctness. The predicate $B \equiv \forall j, k : k \neq j \Rightarrow d(p_j, p_k) \geq G[j]$ is lattice-linear on the reverse-ordered real lattice: if B is false because some (j, k) witnesses $d(p_j, p_k) < G[j]$, then j is forbidden, and every vector at or below G in the lattice (i.e., with a j -coordinate at least $G[j]$) has j forbidden for the same reason. The advance step moves $G[j]$ to the exact minimum over k , which is the smallest value consistent with B .

Complexity. Each advance step computes an $O(n)$ minimum, and each j is advanced at most once, so the total work is $O(n^2)$. With n processors, every $G[j]$ can be computed in parallel, giving $O(n)$ parallel time. This matches the brute-force bound and is worse than the $O(n \log n)$ divide-and-conquer algorithm of Section 9.3; it is the LLP formulation's simplicity and parallelism, not its asymptotic speed, that make it valuable.

Counting Inversions

Given an array $A[1..n]$, we assign to each index j the number of indices that are to its left and strictly larger. Let $G[j]$ denote this count; the total number of inversions is $\sum_j G[j]$.

Algorithm LLP-CountInversions: LLP formulation of the counting-inversions problem.

Input: Array $A[1..n]$

Output: G : array[1.. n] of int; total inversions = $\sum_j G[j]$

1 **var** G : array[1.. n] of int;

2 **init** $\forall j : G[j] := 0$;

3 **forbidden**(j): $G[j] < |\{i \in [1..j-1] : A[i] > A[j]\}|$;

4 **advance**(j): $G[j] := |\{i \in [1..j-1] : A[i] > A[j]\}|$;

Correctness. The target vector $G^*[j] = |\{i < j : A[i] > A[j]\}|$ depends only on the input array, so the predicate $B \equiv \forall j : G[j] = G^*[j]$ is lattice-linear. Each index is forbidden at most once and is advanced directly to $G^*[j]$.

Complexity. Advancing $G[j]$ scans $j-1$ left neighbours for a work of $O(j)$; summed over all j , the total work is $O(n^2)$. With n processors all indices can advance in parallel, giving $O(n)$ parallel time. The divide-and-conquer algorithm of Section 9.4 achieves $O(n \log n)$ sequential time by reusing the work of the merge step; the LLP formulation trades this sequential optimisation for a one-shot parallel advance.

Planar Convex Hull

The convex hull of a point set is naturally expressed by a Boolean lattice on the vertices. Let $G[j] \in \{0, 1\}$ indicate whether p_j is a vertex of the convex hull. Starting from $G[j] = 1$ for all j , we repeatedly turn off

any interior point.

Algorithm LLP-ConvexHull: LLP formulation of the convex-hull membership problem.

Input: Points $p_1, \dots, p_n \in \mathbb{R}^2$

Output: G : array[1.. n] of $\{0, 1\}$; $G[j] = 1$ iff p_j is on the convex hull

1 **var** G : array[1.. n] of $\{0, 1\}$;

2 **init** $\forall j : G[j] := 1$;

3 **forbidden**(j): $G[j] = 1 \wedge \exists i, k, \ell \neq j : G[i] = G[k] = G[\ell] = 1 \wedge p_j \in \text{triangle}(p_i, p_k, p_\ell)$;

4 **advance**(j): $G[j] := 0$;

Correctness. A point p_j is off the convex hull iff it lies strictly inside some triangle spanned by three other hull candidates. The lattice is the reverse-ordered Boolean cube (a smaller G is larger in the lattice): every advance strictly decreases $G[j]$ from 1 to 0, so the algorithm terminates in at most n rounds. At the least fixed point, G is the characteristic vector of the convex hull.

Complexity and caveat. Each forbidden check considers $O(n^3)$ triangles, for $O(n^4)$ total work — far worse than the $O(n \log n)$ divide-and-conquer algorithm of Section 9.5. Moreover, the LLP formulation returns the hull as a membership vector rather than as an ordered polygon; recovering the cyclic order requires a post-processing step. This illustrates that not every divide-and-conquer problem has a clean LLP reformulation, and that some problems — especially those whose output carries non-trivial structure beyond a per-index answer — are better served by their classical recursive algorithms.

Karatsuba and Strassen

Integer multiplication (Karatsuba) and matrix multiplication (Strassen) compute a uniquely determined output rather than searching a lattice for a predicate-satisfying element. Their speed-ups stem from algebraic identities that reduce the number of recursive multiplications, not from lattice search. Consequently neither admits a natural LLP reformulation. We mention this explicitly because it is a useful counter-example: the LLP framework is broadly applicable but is not the right tool for every divide-and-conquer problem.

9.10 Summary

In this chapter, we have shown that many problems can be solved using the divide-and-conquer approach. The following table summarizes some of the problems discussed in this chapter.

Problem	Algorithm	Time Complexity
Sorting	MergeSort	$O(n \log n)$
Nearest Neighbors	ClosestPair	$O(n \log n)$
Counting Inversions	CountingInversions	$O(n \log n)$
Convex Hull	Divide-Conquer	$O(n \log n)$
Integer Multiplication	Karatsuba	$O(n^{\log_2 3})$
Matrix Multiplication	Strassen's Algorithm	$O(n^{\log_2 7})$

Table 9.1: Divide-and-Conquer Algorithms

9.11 Problems

- Solve the problem of computing the Fast-Fourier-Transform using the divide and conquer approach.
- Use the Master Theorem to solve the following recurrences. In each case, state which case of the Master Theorem applies.
 - $T(n) = 9T(n/3) + n$
 - $T(n) = 2T(n/2) + n \log n$
 - $T(n) = 3T(n/4) + n \log n$
 - $T(n) = 7T(n/2) + n^2$
- Given an array $A[1..n]$ of integers (possibly negative), find a contiguous subarray with maximum sum. Give an $O(n \log n)$ divide-and-conquer algorithm, and prove its correctness.
- Given an array $A[1..n]$ of distinct numbers, count the number of *inversions*, i.e., pairs (i, j) with $i < j$ and $A[i] > A[j]$. Give a divide-and-conquer algorithm in time $O(n \log n)$.
- Give a divide-and-conquer algorithm to find the k -th smallest element of an unsorted array $A[1..n]$ in expected $O(n)$ time. Prove the expected time bound.
- Give the high-level idea of the closest-pair-of-points divide-and-conquer algorithm in the plane, and explain why the “strip” step requires checking only a constant number of points per point. State and prove the $O(n \log n)$ complexity.
- Consider the *skyline problem*: given n rectangular buildings defined by triples (l_i, h_i, r_i) (left, height, right), compute the silhouette they form. Give a divide-and-conquer algorithm and its complexity.
- Prove that Strassen’s algorithm multiplies two $n \times n$ matrices in $O(n^{\log_2 7})$ time, and explain why we cannot get sub-quadratic time by a divide-and-conquer approach that performs 8 recursive multiplications per level.
- Prove or disprove: the standard top-down mergesort algorithm (recursive merge-sort that splits in the middle and merges two sorted halves) is *stable*, i.e., it preserves the relative order of equal elements.
- Given two convex polygons in the plane with a total of n vertices, show how to merge them into a single convex hull in $O(n)$ time. Use this to derive an $O(n \log n)$ divide-and-conquer algorithm for the planar convex hull of n points.
- (*Splittable predicate for sorting.*) Let A be an array of length n (with n a power of two for simplicity) and let

$$B(A) \equiv \forall i \in [1..n-1] : A[i] \leq A[i+1].$$

Split A at the midpoint into A_L and A_R . Identify sub-predicates $B_L(A_L)$ and $B_R(A_R)$ and a “merge” relation m such that

$$B(A_L \cdot A_R) \iff B_L(A_L) \wedge B_R(A_R) \wedge m(A_L, A_R),$$

where $A_L \cdot A_R$ denotes concatenation. Explain how this decomposition yields the Mergesort algorithm and state its running time from the induced recurrence.

12. (*Splittable predicate for maximum subarray sum.*) Given an array $A[1..n]$ of integers (possibly negative), let

$$B(A) = \max_{1 \leq i \leq j \leq n} \sum_{k=i}^j A[k]$$

be the maximum contiguous subarray sum. Show that B is splittable by constructing a four-component recursive summary $S(A) = (\text{total}, \text{prefix}, \text{suffix}, \text{best})$ for each sub-array, where

- $\text{total}(A)$ is the sum of all elements,
- $\text{prefix}(A)$ is the maximum sum of a prefix of A ,
- $\text{suffix}(A)$ is the maximum sum of a suffix of A ,
- $\text{best}(A) = B(A)$ is the maximum contiguous subarray sum.

Describe the combine rule $S(A_L \cdot A_R)$ in terms of $S(A_L)$ and $S(A_R)$, and derive the running time.

9.12 Bibliographic Remarks

The general divide-and-conquer strategy and the Master Theorem are thoroughly treated in [CLRS09]. Strassen's matrix multiplication (Section 9.7), first proposed by [Str69], reduces the complexity of a matrix multiplication of size $n \times n$ from $O(n^3)$ to $O(n^{2.807})$. Karatsuba's multiplication introduced by [KO62] reduces the complexity of multiplying two n digit numbers from $O(n^2)$ to $O(n^{1.585})$. The Closest Pair problem uses a divide-and-conquer approach from [SH75], to obtain the time complexity $O(n \log n)$. Counting Inversions via Mergesort, achieving $O(n \log n)$, is a standard application of the divide-and-conquer approach. The Planar Convex Hull is based on [Gra79], with its $O(n \log n)$ time complexity. For further reading, we refer the readers to [Hoa62], [Str69], [KO62], [SH75], and [Gra79].

Chapter 10

Dynamic Programming

10.1 Introduction

Dynamic programming, introduced by Bellman in 1952 [Bel52], is a method for solving problems whose recursive solution involves overlapping sub-problems. Naive recursion recomputes the same sub-problems exponentially many times; dynamic programming records each sub-problem's answer in a table the first time it is computed and reuses it thereafter, replacing exponential redundancy with polynomial work. The name predates computer science: Bellman coined “dynamic programming” to describe a method for time-staged decision processes; today the term covers any algorithm built around a recurrence over a memoised table.

Three ingredients distinguish dynamic-programming problems from generic recursive ones. First, the optimal solution must satisfy a *principle of optimality*: an optimal solution to the whole problem extends an optimal solution to a smaller sub-problem. Second, the sub-problems must *overlap* so that memoisation actually saves work; if every recursive call is on a fresh sub-problem, divide-and-conquer suffices and the memo table is wasted. Third, the recursion must terminate, so the sub-problem space is finite and admits a topological order in which the table can be filled bottom-up.

The lattice-linear predicate (LLP) framework introduced in Chapter 2 reformulates these problems as searches for the least vector G in a finite distributive lattice that satisfies a lattice-linear predicate B . Dynamic-programming recurrences supply the recurrence; LLP supplies a parallel scheduler. Where the dynamic-programming algorithm fills its table in a fixed bottom-up order, the LLP algorithm advances any forbidden index in parallel, and reaches the same fixed point. The LLP view also makes it easy to add lattice-linear constraints to a problem (a feature exercised in the constrained scheduling and constrained binary-search-tree problems below) without changing the algorithm.

This chapter is organized as follows. Section 10.2 contrasts recursion with dynamic programming on the Fibonacci recurrence, motivating memoisation. Section 10.3 develops weighted interval scheduling. Section 10.4 presents the longest increasing subsequence problem. Section 10.5 gives the optimal binary search tree problem. Section 6.5 treats Huffman coding (revisited under the dynamic-programming lens). Section 10.6 addresses chain matrix multiplication. Section 10.7 treats the knapsack problem. Section 10.8 collects the LLP formulations of the algorithms in this chapter.

10.2 Recursion vs Dynamic Programming

Dynamic programming is applicable to problems in which it is easy to set up a recurrence relation so that the solution of the problem at hand can be derived from the solutions to problems with smaller sizes. The problem can be solved using recursion; however, recursion can result in many duplicate computations.

As a simple example, suppose that our goal is to compute the Fibonacci number F_i such that it satisfies the following conditions:

- $F_0 = 1$
- $F_1 = 1$
- $F_i = F_{i-1} + F_{i-2}$ for $i \geq 2$

A recursive algorithm is as follows:

Algorithm RecFib: Recursive Fibonacci Function.

Input: n : integer

Output: $F(n)$

```

1 Function  $F(n)$ :
2   if  $n < 2$  then return 1 ;
3   else return  $F(n - 1) + F(n - 2)$  ;
```

Example 10.1 To compute $F(4)$, the recursive algorithm calls $F(3)$ and $F(2)$. Now $F(3)$ in turn calls $F(2)$ and $F(1)$, so $F(2)$ is computed twice. As n grows the redundancy explodes: the recursion tree of $F(n)$ has $\Theta(\phi^n)$ nodes (where ϕ is the golden ratio), although only $n + 1$ distinct sub-problems exist.

Using *memoization* for dynamic programming, we avoid such duplicate computations. Memoization is used to recall previously computed values.

An implementation using memoization is as follows:

Algorithm RecFibMemo: Recursive Fibonacci with Memoization.

Input: n : integer

Output: $Fibonacci(n)$

```

1 var  $A$ : array[1.. $N$ ] of int initially  $\forall i : A[i] = -1$ 
2 Function  $F(n)$ :
3   if  $A[n] \neq -1$  then return  $A[n]$  ;
4   else if  $n < 2$  then  $A[n] \leftarrow 1$  ;
5   else  $A[n] \leftarrow F(n - 1) + F(n - 2)$  ;
6   return  $A[n]$  ;
```

Alternatively, instead of computing $F(n)$ in a top-down manner, one can compute it in a bottom-up fashion. Assume that we keep an array A such that $A[i]$ will eventually store $F(i)$. We start by filling in the array A from the lower to higher indices. The recurrence relation guarantees that if we have two previous entries, we can fill any entry $A[i]$. Thus, we can compute $F(n)$ in $O(n)$ time.

In this example, we had a simple array. For more nontrivial examples, we may have two- or three-dimensional tables.

10.3 Weighted Interval Scheduling

Recall the problem of Interval Scheduling. Suppose that we have n intervals with start times s_i and finish times f_i for jobs $i = 1..n$, where $s_i < f_i$. We assume that activity i occurs during the open interval $[s_i, f_i)$. Two intervals i and j are *compatible* if $f_i \leq s_j$ or $f_j \leq s_i$. Our goal is to select a maximum-sized subset of mutually compatible intervals. We assume that the intervals are sorted by their finish times. We assume that jobs with the same finish times are sorted according to their start times. Furthermore, no two intervals have identical start and finish times. For this problem, we know that a greedy algorithm works. The interval that finishes earliest and starts after the last chosen interval is always selected.

Now consider a generalization of the problem in which each interval is assigned a *weight* denoting its priority. We would like to compute a subset of intervals such that they are non-overlapping and have the maximum weight. In this generalization, the greedy algorithm does not work.

As before, we assume that the intervals are sorted according to their *finish* times. Let $Opt(j)$ denote the optimal value that can be obtained from the intervals $1..j$. When j is zero, the set of intervals selected is empty and $Opt(0)$ is zero. Let $p(j)$ be the largest interval before the interval j which is disjoint from the interval j . Then we can set up a recurrence relation as follows. The maximum value we can obtain from the intervals $1..j$ is equal to the maximum of two values obtained by including the interval j or not including the interval j . If we include the interval j , then the value obtained is $w_j + Opt(p(j))$. If we do not include the interval j , then the value obtained is $Opt(j - 1)$. Thus, we have

$$Opt(j) = \max\{w_j + Opt(p(j)), Opt(j - 1)\}$$

This recurrence relation allows us to obtain the algorithm for weighted interval scheduling as shown in Fig. [WeightedIntervalScheduling](#).

The sequential algorithm takes $O(n)$ time when p array is available. This time complexity is optimal. Computing p array takes $O(n \log n)$ time.

Algorithm WeightedIntervalScheduling: A Sequential Program for the Weighted Interval Scheduling problem

```

1 // jobs are sorted by their finish times;
  Input:  $s$ : array[0... $n-1$ ] of int (start times);  $f$ : array[0... $n-1$ ] of int (finish times);  $w$ :
           array[0... $n-1$ ] of int (weight of the interval)
  Output:  $G$ : array[0... $n-1$ ] of 0..1, indicating selected intervals;  $opt$ : array of optimal values
2 var  $G$ : array[0... $n-1$ ] of 0..1 initially 0;
3  $p$ : array[0... $n-1$ ] of int ;                               // previous non-overlapping interval
4  $opt$ : array[0... $n-1$ ] of int ;                               // optimal value

5 //  $p[j]$  is the largest interval  $i$  such that  $f[i] < s[j]$ ;
6  $p[0] := 0$ ;
7 // Compute  $p[j]$  as the largest  $i$  such that  $f[i] < s[j]$ ;
8 // Since  $f$  is sorted, we can compute all  $p[j]$  in  $O(n \log n)$  time using binary search

9  $opt[0] := 0$ ;
10 for int  $cur := 1$  to  $n-1$  do
11    $opt[cur] := opt[cur-1]$ ;
12   if ( $w[cur] + opt[p[cur]] \geq opt[cur-1]$ ) then
13      $opt[cur] := w[cur] + opt[p[cur]]$ ;
14      $G[cur] := 1$ ;
15   end
16   else  $G[cur] := 0$ ;
17 end

```

Example 10.2 Consider the following 5 intervals, sorted by finish time:

Interval	s_i	f_i	w_i
1	1	3	4
2	2	5	6
3	4	6	5
4	6	8	3
5	5	9	7

The intervals are shown in Fig. 10.1. We first compute $p(i)$ (the last nonoverlapping interval before i): $p(1) = 0$, $p(2) = 0$ ($f_1 = 3 > s_2 = 2$), $p(3) = 1$, $p(4) = 3$, $p(5) = 2$. Then we compute $OPT(i)$:

- $OPT(0) = 0$.
- $OPT(1) = \max(OPT(0), w_1 + OPT(p(1))) = \max(0, 4) = 4$ (select 1).
- $OPT(2) = \max(OPT(1), w_2 + OPT(p(2))) = \max(4, 6) = 6$ (select 2).
- $OPT(3) = \max(OPT(2), w_3 + OPT(p(3))) = \max(6, 5 + 4) = 9$ (select 1, 3).
- $OPT(4) = \max(OPT(3), w_4 + OPT(p(4))) = \max(9, 3 + 9) = 12$ (select 1, 3, 4).
- $OPT(5) = \max(OPT(4), w_5 + OPT(p(5))) = \max(12, 7 + 6) = 13$ (select 2, 5).

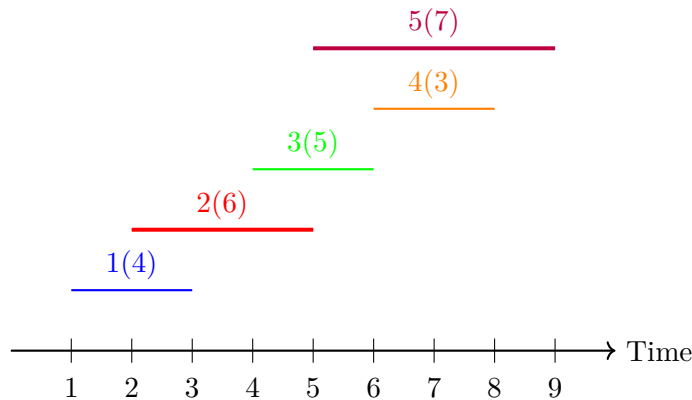


Figure 10.1: Weighted intervals (weights in parentheses) with optimal selection (thick lines).

10.4 Longest Increasing Subsequences

The Longest Increasing Subsequence (LIS) problem involves finding a subsequence of a given array of integers, where the elements are in strictly increasing order and the subsequence is as long as possible. A subsequence is derived by deleting zero or more elements from the original array without changing the order of the remaining elements.

For example, in the array $[10, 9, 2, 5, 3, 7, 101, 18]$, one possible LIS is $[2, 5, 7, 101]$ with length 4. The problem has applications in data analysis, sequence alignment, and more. We present an iterative dynamic programming solution to compute the LIS length efficiently.

Our problem can be stated as follows. Given an array $A = [a_1, a_2, \dots, a_n]$ of n integers, find the length of the longest subsequence $[a_{i_1}, a_{i_2}, \dots, a_{i_k}]$ such that:

$$i_1 < i_2 < \dots < i_k \quad \text{and} \quad a_{i_1} < a_{i_2} < \dots < a_{i_k}.$$

We use an iterative dynamic programming approach to solve the LIS problem. Define $dp[i]$ as the length of the longest increasing subsequence ending at index i (including a_i).

The idea is to build the dp array iteratively by comparing each element with all the previous elements.

- Initialize $dp[i] = 1$ for all i (each element is an LIS of length 1 itself).
- For each i , check all $j < i$. If $a[j] < a[i]$, update $dp[i]$ as $\max(dp[i], dp[j] + 1)$.
- The length of the LIS is the maximum value in the dp array.

Algorithm LIS: Longest Increasing Subsequence (Iterative)

Input: Array A of n integers

Output: Length of the LIS

```

1  $dp$ : array  $[0 \dots n - 1]$  of int initially  $\forall i : dp[i] = 1$ ;
2 for  $i = 1$  to  $n - 1$  do
3   | for  $j = 0$  to  $i - 1$  do
4   |   | if  $A[j] < A[i]$  then  $dp[i] = \max(dp[i], dp[j] + 1)$ ;
5   |   end
6 end
7 return  $\max(dp[0], dp[1], \dots, dp[n - 1])$ ;

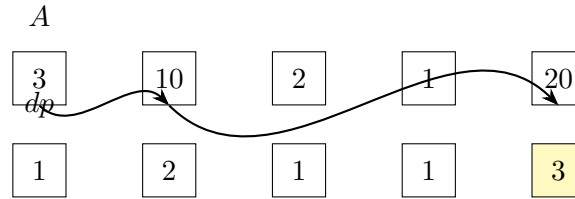
```

$dp[i]$ represents the longest increasing subsequence ending in $A[i]$, considering all prior elements. By checking all $j < i$, we ensure that every possible subsequence ending at i is evaluated. The maximum $dp[i]$ gives the overall length of the LIS, as it captures the longest chain possible.

Example 10.3 Consider the array $A = [3, 10, 2, 1, 20]$. We compute $dp[i]$ for each index:

- $A[0] = 3, dp[0] = 1$.
- $A[1] = 10, dp[1] = 2$.
- $A[2] = 2, dp[2] = 1$.
- $A[3] = 1, dp[3] = 1$.
- $A[4] = 20, dp[4] = 3$.

The final $dp = [1, 2, 1, 1, 3]$, the LIS length is $\max(dp) = 3$, and an LIS is $[3, 10, 20]$.



The highlighted $dp[4] = 3$ indicates the LIS length. Arrows show one possible LIS: $[3, 10, 20]$.

The time complexity is $O(n^2)$ and the space complexity is $O(n)$ for the dp array.

10.5 Optimal Binary Search Tree

The Optimal Binary Search Tree problem involves constructing a binary search tree (BST) for n symbols that minimizes the expected search cost, given the probability of each symbol's occurrence. Each symbol i (ranging from 0 to $n - 1$) has a probability $p[i]$ of being searched, and these probabilities sum to 1, i.e., $\sum_{i=0}^{n-1} p[i] = 1$. The objective is to arrange the symbols in a BST to minimize the total cost, defined as the sum of each symbol's probability multiplied by its depth, where the root has depth 1. For example, given three symbols A , B , and C with probabilities $p = [0.4, 0.3, 0.3]$, an optimal BST might place B as the root to reduce the average search time. This problem finds applications in areas like database indexing and compiler design, where efficient search structures are crucial. We present an iterative dynamic programming solution to compute the minimum cost.

Formally, given n symbols indexed from 0 to $n - 1$, each associated with probability $p[i]$, and defining the sum of probabilities from i to j as $s(i, j) = \sum_{k=i}^j p[k]$, the task is to construct a BST that minimizes the expected search cost, expressed as $\text{Cost} = \sum_{i=0}^{n-1} p[i] \cdot \text{depth}(i)$, where $\text{depth}(i)$ represents the depth of the symbol i in the tree.

Algorithm [OptimalBinarySearchTree](#) solves this by iterative dynamic programming. Two key arrays are defined: $dp[i][j]$ represents the minimum cost of a BST for symbols from i to j (inclusive), and $s(i, j)$ denotes the sum of probabilities from i to j , which serves as the cost increase at each level. The solution builds the dp table iteratively over increasing lengths of subarrays. When considering a single symbol ($j = i$), the cost is simply $dp[i][i] = p[i]$, reflecting the contribution of a leaf node. For larger ranges, each symbol r from i to j is tested as the root, combining the left subtree $[i, r - 1]$ and the right subtree $[r + 1, j]$. The total cost for a root r is computed as $dp[i][r - 1] + dp[r + 1][j] + s(i, j)$, where $s(i, j)$ accounts for the increased depth of all symbols in the subtree. To optimize, $s(i, j)$ is precomputed iteratively.

The correctness of this approach comes from the fact that $dp[i][j]$ computes the minimum cost by exhaustively trying each possible root r and combining the optimal costs of the resulting subproblems. The term $s(i, j)$ ensures that the cost reflects the increase in depth for all symbols in the subtree. By filling the table bottom-up, from single symbols to the full range, the iterative method guarantees that $dp[0][n - 1]$ yields the optimal cost.

To analyze the time complexity, consider the three nested loops in the algorithm. The outer loop iterates over lengths l from 1 to $n - 1$, which takes $O(n)$ time. For each l , the middle loop iterates over the starting indices i from 0 to $n - 1 - l$, also $O(n)$ per length. Within these, the inner loop tests each root r from i to j , which can be up to $O(n)$ iterations. This results in a total of $O(n) \cdot O(n) \cdot O(n) = O(n^3)$ operations. Precomputing $s[i][j]$ requires $O(n^2)$ time, as each entry builds on the previous, but this is dominated by the main computation. The total time complexity is therefore $O(n^3)$, with a space complexity of $O(n^2)$ for the tables dp and s .

Algorithm OptimalBinarySearchTree: Optimal Binary Search Tree (Iterative)

Input: Array $p[0 \dots n - 1]$ of probabilities

Output: Minimum cost of Optimal BST

```

1 Initialize  $dp[0 \dots n - 1][0 \dots n - 1]$  and  $s[0 \dots n - 1][0 \dots n - 1]$  with 0;
2 for  $i = 0$  to  $n - 1$  do
3    $dp[i][i] = p[i]$ ;
4    $s[i][i] = p[i]$ ;
5 end
6 for  $l = 1$  to  $n - 1$  do
7   for  $i = 0$  to  $n - 1 - l$  do
8      $j = i + l$ ;
9      $s[i][j] = s[i][j - 1] + p[j]$ ;
10     $dp[i][j] = \infty$ ;
11    for  $r = i$  to  $j$  do
12       $left\_cost = 0$ ;
13       $right\_cost = 0$ ;
14      if  $r > i$  then  $left\_cost = dp[i][r - 1]$  ;
15      if  $r < j$  then  $right\_cost = dp[r + 1][j]$  ;
16       $cost = s[i][j] + left\_cost + right\_cost$ ;
17       $dp[i][j] = \min(dp[i][j], cost)$ ;
18    end
19  end
20 end
21 return  $dp[0][n - 1]$ ;

```

10.6 Chain Matrix Multiplication

A problem very similar to the Optimal Binary Search Tree is that of constructing an optimal way of multiplying a chain of matrices. Since matrix multiplication is associative, the product of matrices $(M_1 * M_2) * M_3$ is equal to $M_1 * (M_2 * M_3)$. However, depending on the dimensions of the matrices, the computational effort may be different.

Example 10.4 Consider three matrices M_1, M_2, M_3 with dimensions $30 \times 10, 10 \times 30,$ and $30 \times 2,$ respectively. The product $M_1 \cdot M_2 \cdot M_3$ has dimension 30×2 regardless of the order of multiplication, but the cost differs:

- $(M_1 \cdot M_2) \cdot M_3$: computing $M_1 \cdot M_2$ takes $30 \cdot 10 \cdot 30 = 9000$ scalar multiplications and yields a 30×30 matrix; multiplying by M_3 adds $30 \cdot 30 \cdot 2 = 1800$, for a total of 10800.
- $M_1 \cdot (M_2 \cdot M_3)$: computing $M_2 \cdot M_3$ takes $10 \cdot 30 \cdot 2 = 600$ and yields a 10×2 matrix; multiplying M_1 by this adds $30 \cdot 10 \cdot 2 = 600$, for a total of 1200.

The right-associative grouping is nine times cheaper.

We let the dimension of matrix M_i be $m_{i-1} \times m_i$. Note that this keeps the matrix product well-defined because the dimension of the matrix M_{i+1} would be $m_i \times m_{i+1}$ and the product $M_i \times M_{i+1}$ is well-defined. We can view any evaluation of a chain as a binary tree where the intermediate nodes are the multiplication operation, and the leaves are the matrices themselves. Suppose that our goal is to compute the optimal binary tree for multiplying matrices in the range $M_i \dots M_j$. Taking ideas from the previous section, we let $G[i, j]$ denote the optimal cost of computing the product of matrices in the range $M_i \dots M_j$. Suppose that this product is broken into products of $M_i \dots M_k$ and $M_{k+1} \dots M_j$ and then the multiplication of these two matrices. We can compute the cost of this tree as

$$G[i, k] + G[k + 1, j] + m_{i-1}m_k m_j$$

Then, we have the following predicate on G .

$$G[i, j] \geq \min_{i \leq k < j} (G[i, k] + m_{i-1}m_k m_j + G[k + 1, j])$$

The reader will notice the similarity to the optimal binary search tree problem, and the same algorithm can be adapted to solve this problem.

10.7 Knapsack Problem

We are given n items with weights w_1, w_2, \dots, w_n and values v_1, v_2, \dots, v_n . We are also given a knapsack that has a capacity of W . Our goal is to determine the subset of items that can be carried in the knapsack and that maximizes the total value. The standard dynamic programming solution is based on the memoization of the following dynamic programming formulation [Vaz01, DPW10]. Let $G[i, w]$ be the maximum value that can be obtained by picking items from $1..i$ with the capacity constraint of w . Then, $G[i, w] = \max(G[i - 1, w - w_i] + v_i, G[i - 1, w])$. The first argument of the max function corresponds to the case when the item i is included in the optimal set from $1..i$, and the second argument corresponds

to the case when the item i is not included and hence the entire capacity can be used for the items from $1..i - 1$. If $w_i > w$, then the item i cannot be in the knapsack and can be skipped. The base cases are simple. The values of $G[0, w]$ and $G[i, 0]$ are zero for all w and i . Our goal is to find $G[n, W]$. By filling in the two-dimensional array G for all values of $0 \leq i \leq n$ and $0 \leq w \leq W$, we get an algorithm with time complexity $O(nW)$.

The following algorithm computes $G[n, W]$ and tracks the selected items:

Algorithm Knapsack: 0/1 Knapsack Dynamic Programming

Input: Weights $w[1..n]$, Values $v[1..n]$, Capacity W
Output: Maximum value and selected items

```

1 for  $w = 0$  to  $W$  do  $G[0, w] := 0$  ;
2 for  $i = 0$  to  $n$  do  $G[i, 0] := 0$  ;
3 for  $i = 1$  to  $n$  do
4   for  $w = 1$  to  $W$  do
5     if  $w_i > w$  then  $G[i, w] := G[i - 1, w]$  ;
6     else  $G[i, w] := \max(G[i - 1, w], G[i - 1, w - w_i] + v_i)$  ;
7   end
8 end
9  $selected := \emptyset$ ;
10  $i := n$ ;
11  $w := W$ ;
12 while  $(i > 0) \wedge (w > 0)$  do
13   if  $G[i, w] \neq G[i - 1, w]$  then
14      $selected := selected \cup \{i\}$ ;
15      $w := w - w_i$ ;
16   end
17    $i := i - 1$ ;
18 end
19 return  $G[n, W], selected$ ;

```

10.8 LLP Perspective

In this section we collect the LLP reformulations of the dynamic-programming algorithms introduced earlier in the chapter. Each formulation views the problem as the search for the least vector in a finite distributive lattice satisfying a lattice-linear predicate B ; the bottom-up DP table-fill is one valid schedule, but the LLP formulation also admits parallel schedules in which several forbidden indices advance concurrently.

LLP-WeightedIntervalScheduling

For weighted interval scheduling (Section 10.3), let $G[j]$ denote $Opt(j)$, the maximum weight achievable using intervals $1, \dots, j$. The lattice is $\mathbb{Z}_{\geq 0}^{n+1}$ with the usual order; j is forbidden when $j - 1$ and $p(j)$ are

both fixed but j is not yet, and the advance sets $G[j]$ to the recurrence value.

Input: p : array[1... n] of int; w : array[1... n] of int
Output: G : array[0... n] of int, with $G[n] = \text{Opt}(n)$
1 init: $G[j] := 0$, $\text{fixed}[j] := \text{false}$ for $j = 0 \dots n$; $\text{fixed}[0] := \text{true}$;
2 forbidden(j): $\neg \text{fixed}[j] \wedge \text{fixed}[j-1] \wedge \text{fixed}[p[j]]$;
3 advance(j): $G[j] := \max(G[j-1], w[j] + G[p[j]])$; $\text{fixed}[j] := \text{true}$;

LLP-LIS

For the longest increasing subsequence problem (Section 10.4), let $G[j]$ denote the length of the longest strictly increasing subsequence ending at index j . With $\text{pre}(j) = \{i < j : A[i] < A[j]\}$, the index j is forbidden when its predecessors are all fixed but j itself is not.

Input: A : array[1.. n] of real
Output: G : array[1.. n] of int; $\max_j G[j]$ is the LIS length
1 init: $G[j] := 1$, $\text{fixed}[j] := \text{false}$ for $j = 1 \dots n$;
2 forbidden(j): $\neg \text{fixed}[j] \wedge \forall i \in \text{pre}(j) : \text{fixed}[i]$;
3 advance(j): $G[j] := \max(\{1\} \cup \{G[i] + 1 : i \in \text{pre}(j)\})$; $\text{fixed}[j] := \text{true}$;

LLP-OptimalBinarySearchTree

For the optimal binary search tree problem (Section 10.5), let $G[i, j]$ denote the least cost of any binary search tree built from the keys in the range $i..j$, and let $s(i, j) = \sum_{k=i}^j p_k$ denote the sum of access frequencies in that range (with $s(i, j) = 0$ when $i > j$). If symbol k is the root of the optimal tree on $i..j$, the optimal-substructure property gives

$$G[i, j] = \min_{i \leq k \leq j} (G[i, k-1] + s(i, j) + G[k+1, j]).$$

Lemma 10.5 *The predicate $B \equiv \forall i, j : G[i, j] \geq \min_{i \leq k \leq j} (G[i, k-1] + s(i, j) + G[k+1, j])$ is lattice-linear.*

Proof: If B is false then some index (i, j) satisfies $G[i, j] < \min_{i \leq k \leq j} (G[i, k-1] + \sum_{\ell=i}^j p[\ell] + G[k+1, j])$. The right-hand side is monotone in G (smaller G values can only decrease the minimum), so unless $G[i, j]$ itself increases, B cannot become true regardless of how other components advance. Hence (i, j) is forbidden. ■

The LLP algorithm uses a single variable G initialised to zero everywhere; it advances $G[i, j]$ whenever

it is smaller than the right-hand side above. Algorithm [LLP-OptimalBinarySearchTree](#) captures this.

Algorithm LLP-OptimalBinarySearchTree: Finding an Optimal Binary Search Tree

1 $P_{i,j}$: Code for thread (i, j) ;
Input: p : array of real (frequency of each symbol)
Output: G : array of real, optimal BST costs
2 **init:** $\forall i, j : G[i, j] = 0$;
3 **ensure:** $G[i, j] \geq \min_{i \leq k \leq j} G[i, k - 1] + \sum_{\ell=i}^j p[\ell] + G[k + 1, j]$;
4 **priority:** $(j - i)$;

The **priority** statement schedules indices in non-decreasing order of $(j - i)$, processing the singleton ranges first, then ranges of width 2, and so on; this ensures that each $G[i, j]$ is updated at most once. With $O(n^2)$ index pairs and $O(n)$ work per advance, the work complexity is $O(n^3)$.

Constrained variants. The same algorithm extends to constrained versions of the problem with no change of structure:

Lemma 10.6 *The following predicates are lattice-linear:*

1. *The key x is not a parent for any key. The ensure predicate becomes $G[i, j] \geq \min_{i \leq k \leq j, k \neq x} G[i, k - 1] + s(i, j) + G[k + 1, j]$, whose right-hand side remains monotone in G .*
2. *The left and right subtrees of any internal node differ in size by at most 1. The ensure predicate restricts k to indices satisfying $||k - 1 - i| - |j - k - 1|| \leq 1$; the right-hand side remains monotone in G .*

LLP-HuffmanTree

For the Huffman tree construction (Section 6.5), the LLP formulation parallels OBST: $G[i, j]$ is the minimum total weight of an optimal merging of the (sub)alphabet whose probabilities sum to $\sum_{r=i}^j p_r$. The constraint mirrors that of OBST.

Input: p : array[1.. n] of real
Output: G : array[1.. n , 1.. n] of real
1 **init:** $G[i, i] := 0$; $G[i, j] := \infty$ for $i < j$;
2 **ensure**(i, j): $G[i, j] \leq \min_{i \leq k < j} (G[i, k] + G[k + 1, j]) + \sum_{r=i}^j p_r$

LLP-Knapsack

For the 0/1 knapsack problem (Section 10.7) with weights w_i , values v_i , and capacity W , let $G[i, c]$ be the maximum value using items $1, \dots, i$ within capacity c . The lattice is $\mathbb{Z}_{\geq 0}^{n \times (W+1)}$ with the usual order.

Input: w, v : array[1.. n] of int; W : int
Output: G : array[0.. n , 0.. W] of int, with $G[n, W]$ = optimal value
1 **init:** $G[0, c] := 0$ for all c ; $G[i, c] := 0$ otherwise;
2 **forbidden**(i, c): $G[i, c] < \max(G[i - 1, c], G[i - 1, c - w_i] + v_i)$ when $w_i \leq c$;
3 **advance**(i, c): $G[i, c] := \max(G[i - 1, c], G[i - 1, c - w_i] + v_i)$;

10.9 Summary

Dynamic programming captures problems whose optimal solutions can be assembled from optimal solutions to overlapping sub-problems. The trio of recurrence relation, memoisation, and bottom-up table fill are sufficient to convert exponential recursion into polynomial computation. The algorithms presented in this chapter share the following high-level recipe: (i) identify the sub-problem structure — typically indexed by intervals $[i, j]$, prefixes $1..j$, or item-and-capacity pairs (i, c) ; (ii) write a recurrence that expresses the optimal answer for each sub-problem in terms of strictly smaller sub-problems; (iii) fill the resulting table in any topological order respecting the recurrence. Lattice-linear predicate algorithms (LLP) provide an alternative parallel scheduler for the same computation.

Table 10.1 summarizes the time complexity of the sequential algorithms discussed in this chapter.

Problem	Algorithm	Time Complexity
Weighted interval scheduling	WeightedIntervalScheduling	$O(n \log n)$
Increasing subsequence	Longest-Increasing-Subsequence	$O(n^2)$
Optimal binary search tree	OptimalBinarySearchTree	$O(n^3)$
Huffman tree	LLP-HuffmanTree	$O(n^3)$
Chain matrix multiplication	Recurrence-based DP	$O(n^3)$
Knapsack	Knapsack	$O(nW)$

Table 10.1: Algorithms discussed in this chapter.

10.10 Problems

1. We are given a directed acyclic graph (V, E) with n nodes and m edges. Our goal is to assign a number, *label* to each vertex from $1..n$ such that for all edges $(i, j) \in E$, $label[i] < label[j]$. Define B as $\forall (i, j) \in E : label[j] \geq label[i] + 1$. Show that B is a lattice-linear predicate.
2. Modify the algorithm for Optimal Binary Search Tree to return not only the optimal cost but also the tree itself.
3. (**Longest Path in Directed Acyclic Graphs**) We are given a directed acyclic graph (V, E) with n nodes and m edges such that each edge has a positive real cost. We are also given a distinguished vertex v_0 . Give an LLP-based algorithm to find the longest path from v_0 to all vertices.
4. We are given a polygon and we are required to triangulate it optimally. We are given a weight function that takes a triangle on the vertices v_i, v_j and v_k and returns a real-valued function $w(v_i, v_j, v_k)$. Our goal is to triangulate the given polygon to minimize the sum of weights of all triangles. (*Hint: Devise a recurrence relation similar to the matrix chain product problem.*)
5. Give a dynamic programming algorithm for the *Longest Common Subsequence* (LCS) problem: given two sequences $X = x_1 \dots x_m$ and $Y = y_1 \dots y_n$, compute the length of the longest sequence that appears as a subsequence of both. State the recurrence, give the time and space complexity, and describe how to recover an optimal LCS from the DP table.

6. Give a dynamic programming algorithm for the *Edit Distance* problem: given strings X of length m and Y of length n , compute the minimum number of insertions, deletions, and substitutions to transform X into Y . Prove optimal substructure.
7. Give a dynamic programming algorithm for the *0/1 Knapsack* problem with integer weights. Prove that the running time $O(nW)$ is not polynomial in the input size, where W is the knapsack capacity.
8. Give a dynamic programming algorithm for the *Coin Change* problem: given denominations c_1, c_2, \dots, c_k (all positive integers) and a target amount N , find the minimum number of coins that sum to N (or report that N cannot be made).
9. Give an $O(n \log n)$ algorithm for the *Longest Increasing Subsequence* (LIS) problem, improving on the $O(n^2)$ dynamic programming solution. Give a brief justification of correctness.
10. Give a dynamic programming algorithm for the *Matrix Chain Multiplication* problem: given a sequence of matrices with dimensions $p_0 \times p_1, p_1 \times p_2, \dots, p_{n-1} \times p_n$, determine the parenthesization that minimizes the total number of scalar multiplications. State the recurrence and complexity.

10.11 Bibliographic Remarks

This chapter explores the paradigm of solving optimization problems by breaking them into overlapping subproblems, with applications to the Longest Increasing Subsequence (LIS), Optimal Binary Search Tree (OBST), Knapsack, Weighted Interval Scheduling, Segmented Least Squares, and Chain Matrix Multiplication. The chapter introduces dynamic programming (DP) as a method to establish recurrence relations, avoiding redundant computations through memoization or bottom-up computation, as originally formalized by Bellman [Bel52]. Cormen et al. [CLRS01] provide a comprehensive treatment of sequential DP algorithms for these problems.

The OBST problem, introduced by Knuth [Knu71], minimizes expected search cost in a binary search tree. The Knapsack problem, formulated in Horowitz and Sahni [HS74] and Ibarra and Kim [IK75a], is solved in $O(nW)$ time via DP. Vazirani [Vaz01] and Williamson [DPW10] provide modern DP formulations for Knapsack, including approximation techniques.

Weighted Interval Scheduling and Segmented Least Squares, covered in Kleinberg and Tardos [KT06b], extend greedy and DP techniques to optimization problems with weighted objectives and error minimization, respectively. The Chain Matrix Multiplication problem, structurally similar to OBST, is addressed using a recurrence akin to Knuth's formulation, as noted in Aho, Hopcroft, and Ullman [AHU74]. Papadimitriou and Steiglitz [PS82] offer additional context for combinatorial optimization problems like Knapsack and matrix multiplication.

Chapter 11

Network Flow

11.1 Introduction

How much water can flow through a network of pipes from a reservoir to a city? How many vehicles per hour can travel from a suburb to downtown through a road network? How many packets per second can be routed from a server to a client through the internet? These seemingly different questions share a common mathematical structure: they are all instances of the maximum flow problem.

The maximum flow problem is one of the most fundamental problems in combinatorial optimization and network theory. Given a directed graph where each edge has a limited capacity, the goal is to ship as much “flow” as possible from a designated source vertex s to a sink vertex t without exceeding any edge’s capacity. The problem was first studied in the 1950s in the context of Soviet railway networks, when the RAND Corporation sought to determine the maximum rate at which supplies could be transported through the Soviet rail system [JF56]. Harris and Ross formulated the problem in a secret 1955 report, and shortly thereafter Ford and Fulkerson developed both a solution algorithm and the celebrated Max-Flow Min-Cut Theorem.

Beyond its historical origins, the maximum flow problem has remarkably diverse applications:

- *Transportation and logistics.* Determining the maximum throughput of a road, rail, or shipping network between two locations.
- *Telecommunication networks.* Computing the maximum bandwidth between two nodes in a data network, or the maximum number of non-interfering calls that can be simultaneously routed.
- *Bipartite matching.* Finding a maximum matching in a bipartite graph reduces to a max-flow problem.
- *Image segmentation.* In computer vision, separating foreground from background in an image can be formulated as a minimum cut problem, which is equivalent to maximum flow.
- *Scheduling.* Determining whether a set of tasks can be assigned to machines subject to capacity constraints.

The power of maximum flow also stems from its deep connection to *minimum cuts*. The Max-Flow Min-Cut Theorem—one of the most beautiful results in combinatorics—states that the maximum amount

of flow from s to t equals the minimum capacity of any cut separating s from t . This duality has profound algorithmic consequences and connects flow theory to linear programming duality.

This chapter is organized as follows. Section 11.2 provides the definitions of flow networks and flows. Section 11.3 presents the FordFulkerson algorithm and proves its correctness with a detailed worked example. Section 11.4 establishes the Max-Flow Min-Cut Theorem. Section 11.5 presents the EdmondsKarp algorithm, which achieves a strongly polynomial running time by choosing augmenting paths carefully. Section 11.7 casts the chapter through the lattice-linear predicate framework: the lattice of minimum cuts and an algorithm for finding mincuts satisfying a lattice-linear side predicate.

11.2 Definitions

Definition 11.1 (Flow Network) *A flow network is a directed graph $G = (V, E)$ with a source vertex $s \in V$, a sink vertex $t \in V$, and a capacity function $c : E \rightarrow \mathbb{R}^+$ that assigns a non-negative capacity $c(u, v)$ to each edge $(u, v) \in E$.*

We assume that there are no edges entering s and no edges leaving t . If an edge $(u, v) \notin E$, we define $c(u, v) = 0$.

Definition 11.2 (Flow) *A flow in a flow network G is a function $f : E \rightarrow \mathbb{R}^+$ that satisfies:*

1. **Capacity constraint:** $\forall (u, v) \in E, 0 \leq f(u, v) \leq c(u, v)$.
2. **Flow conservation:** $\forall v \in V \setminus \{s, t\}, \sum_{u:(u,v) \in E} f(u, v) = \sum_{w:(v,w) \in E} f(v, w)$.

The capacity constraint says that the flow on each edge cannot exceed its capacity. The conservation constraint says that for every vertex other than s and t , the total flow entering the vertex equals the total flow leaving it—flow is neither created nor destroyed at intermediate vertices.

Definition 11.3 (Value of Flow) *The value of a flow f is defined as the total flow out of the source:*

$$|f| = \sum_{v:(s,v) \in E} f(s, v)$$

The *maximum flow problem* asks: given a flow network G with source s and sink t , find a flow f of maximum value $|f|$.

Definition 11.4 (Residual Network) Given a flow network $G = (V, E)$ and a flow f , the residual network $G_f = (V, E_f)$ consists of edges with remaining capacity, defined as:

$$E_f = \{(u, v) \in V \times V : c_f(u, v) > 0\}$$

where $c_f(u, v)$ is the residual capacity defined as:

$$c_f(u, v) = \begin{cases} c(u, v) - f(u, v) & \text{if } (u, v) \in E \\ f(v, u) & \text{if } (v, u) \in E \\ 0 & \text{otherwise} \end{cases}$$

The residual network captures the remaining capacity to push additional flow. Forward edges have residual capacity equal to the unused capacity. Backward (or reverse) edges have residual capacity equal to the current flow on the corresponding forward edge—this represents the ability to “cancel” previously sent flow, effectively rerouting it.

Definition 11.5 (Augmenting Path) An augmenting path is a simple path from the source s to the sink t in the residual network G_f .

The *bottleneck capacity* of an augmenting path p is $c_f(p) = \min_{(u,v) \in p} c_f(u, v)$, the minimum residual capacity along the path.

Definition 11.6 (Cut) An s - t cut (S, T) is a partition of V into two sets S and $T = V \setminus S$ such that $s \in S$ and $t \in T$. The capacity of the cut is:

$$c(S, T) = \sum_{\substack{u \in S, v \in T \\ (u, v) \in E}} c(u, v)$$

Note that the capacity of a cut only counts edges going from S to T , not edges from T to S .

11.3 The FordFulkerson Algorithm

The FordFulkerson algorithm is a greedy method that computes the maximum flow in a flow network. It was published in 1956 by L. R. Ford, Jr. and D. R. Fulkerson. The algorithm works by repeatedly finding augmenting paths in the residual network and increasing the flow along these paths until no more

augmenting paths exist.

Algorithm FordFulkerson: FordFulkerson Algorithm

Input: Graph G , source s , sink t

Output: Maximum flow f

```

1 Initialize flow  $f(u, v) = 0$  for all edges  $(u, v)$  in  $G$ ;
2 while there exists an augmenting path  $p$  from  $s$  to  $t$  in the residual network  $G_f$  do
3   Let  $c_f(p) := \min\{c_f(u, v) : (u, v) \in p\}$ ;
4   // residual capacity of path  $p$  for each edge  $(u, v)$  in  $p$  do
5     if  $(u, v) \in E$  then  $f(u, v) = f(u, v) + c_f(p)$  // Increase flow ;
6     else  $f(v, u) = f(v, u) - c_f(p)$  // Decrease flow in the reverse edge ;
7   end
8 end
9 return  $f$ 

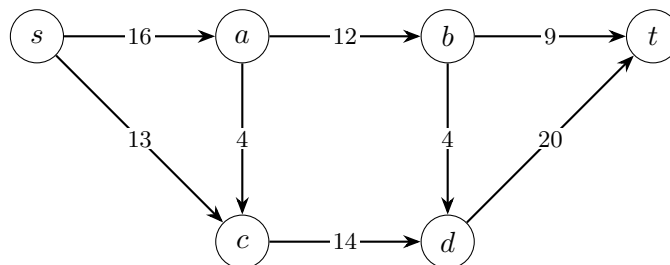
```

Theorem 11.7 *The FordFulkerson algorithm correctly computes the maximum flow in any flow network with integer capacities.*

Proof: The algorithm repeatedly increases the flow along augmenting paths found in the residual graph. Each augmentation strictly increases the total flow value by at least the minimum residual capacity of the augmenting path. Since the capacities are finite and nonnegative, the total flow is bounded above.

The algorithm terminates when there are no more augmenting paths in the residual graph. By the Max-Flow Min-Cut Theorem, a flow is maximum if and only if there is no augmenting path in the residual graph. Therefore, when the algorithm terminates, the current flow is maximal. ■

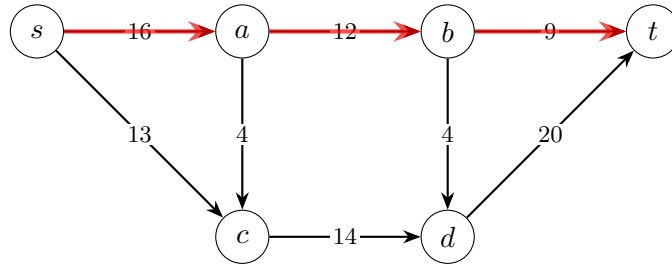
Let us demonstrate the FordFulkerson algorithm step by step on a concrete flow network. Consider a flow network with six vertices $\{s, a, b, c, d, t\}$ and the following capacities:



Initially, all flow values are 0. The edge labels show capacity.

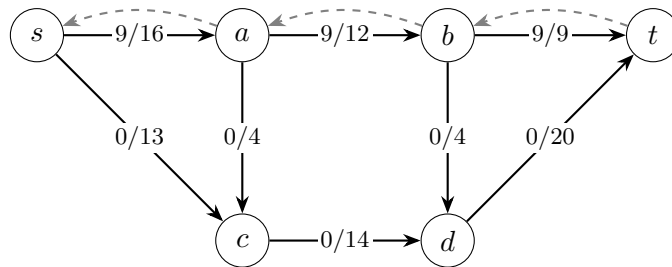
First Iteration

We look for a path from s to t in the residual network. Let us choose the path $s \rightarrow a \rightarrow b \rightarrow t$.



The minimum residual capacity on this path is $\min\{16, 12, 9\} = 9$. We augment the flow by 9 units along this path.

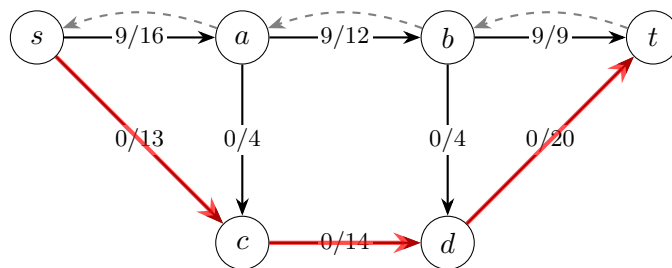
Updated Flow After First Iteration



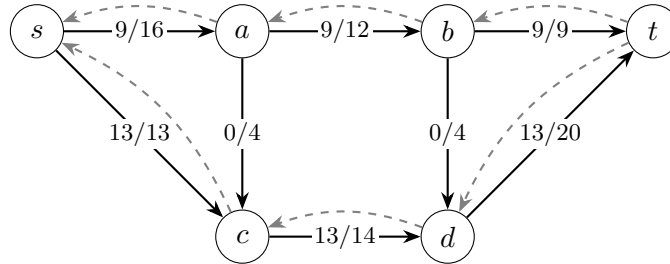
The flow value is now 9. The edge labels show flow/capacity. The residual network now includes reverse edges where the flow was pushed.

Second Iteration

We find another path in the residual network: $s \rightarrow c \rightarrow d \rightarrow t$.



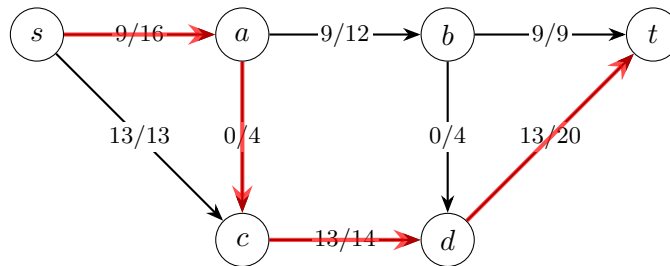
The minimum residual capacity on this path is $\min\{13, 14, 20\} = 13$. We augment the flow by 13 units.

Updated Flow After Second Iteration

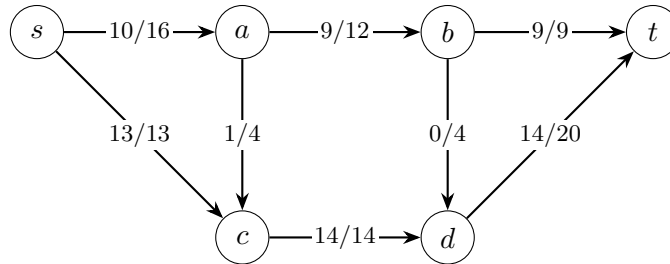
The flow value is now $9 + 13 = 22$.

Third Iteration

We find another path: $s \rightarrow a \rightarrow c \rightarrow d \rightarrow t$.



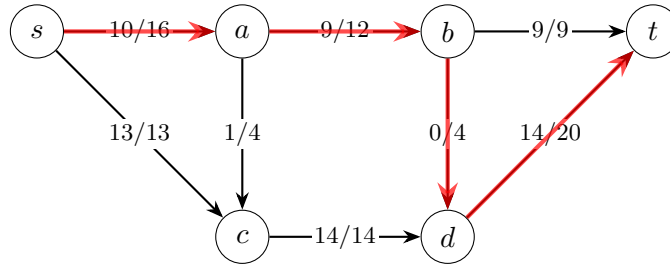
The minimum residual capacity on this path is $\min\{7, 4, 1, 7\} = 1$. We augment the flow by 1 unit.

Updated Flow After Third Iteration

The flow value is now $10 + 13 = 23$.

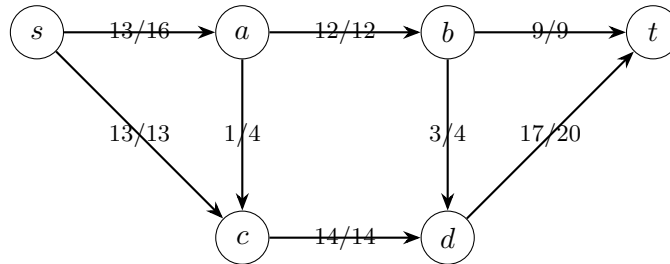
Fourth Iteration

We find one more path: $s \rightarrow a \rightarrow b \rightarrow d \rightarrow t$.



The minimum residual capacity on this path is $\min\{6, 3, 4, 6\} = 3$. We augment the flow by 3 units.

Final Flow



The flow value is now $13 + 13 = 26$. There are no more augmenting paths in the residual network, so the algorithm terminates. The maximum flow is 26.

A natural question is: why do we need reverse edges in the residual network? Could we not simply send flow greedily along forward edges? The following small example shows that without the ability to “undo” flow via reverse edges, a greedy approach can get stuck at a suboptimal solution.

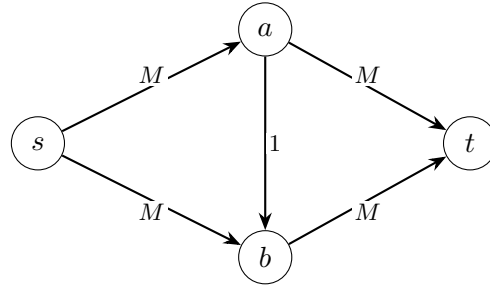
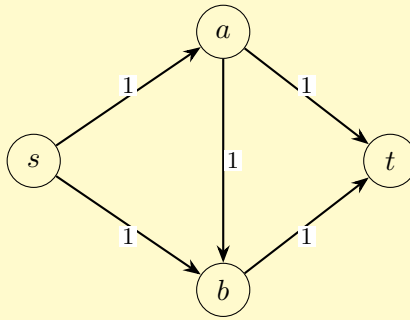


Figure 11.1: An example that shows that the FordFulkerson algorithm may take time proportional to the maxflow in the graph.

Example 11.8 Consider the following network:



The maximum flow is 2 (one unit along $s \rightarrow a \rightarrow t$ and one along $s \rightarrow b \rightarrow t$). However, suppose the algorithm first chooses the path $s \rightarrow a \rightarrow b \rightarrow t$, sending 1 unit. Now all forward edges on this path are saturated. Without reverse edges, we are stuck with flow value 1.

With the residual network, however, the reverse edge from b to a (with residual capacity 1) allows the algorithm to find the augmenting path $s \rightarrow b \rightarrow a \rightarrow t$ in the residual graph, pushing one more unit. The result: 2 units reach t , which is optimal. The reverse edge effectively “reroutes” the unit that was going through $a \rightarrow b$, redirecting the flow from a directly to t and the flow from b directly to t .

Let us determine the running time of the FordFulkerson algorithm. Suppose that all capacities in the graph are integral. Let f^* be the maximum flow in the graph. Every iteration of FordFulkerson’s algorithm finds an augmenting path in the residual network of at least one unit. This step takes $O(|E|)$ time. Hence, the total running time is $O(f^*|E|)$. This algorithm is pseudo-polynomial because the running time depends on the numeric value of the maximum flow f^* , which can be exponential in the input size. It is desirable to get an algorithm whose time complexity depends only on the number of vertices and edges, not on the capacities. For large capacities, this can lead to inefficiencies, motivating strongly polynomial algorithms like the EdmondsKarp and Dinitz algorithms.

To see how bad FordFulkerson can be with poor augmenting path choices, consider the network shown in Figure 11.1.

where M is a large integer. The maximum flow is $2M$. If the algorithm alternately chooses paths

$s \rightarrow a \rightarrow b \rightarrow t$ and $s \rightarrow b \rightarrow a \rightarrow t$ (using the reverse edge each time), each augmentation sends only 1 unit. The algorithm would require $2M$ iterations instead of the 2 iterations needed by choosing $s \rightarrow a \rightarrow t$ and $s \rightarrow b \rightarrow t$.

11.4 Min-Cut and the Max-Flow Min-Cut Theorem

There is a beautiful duality between flows and cuts. Intuitively, a cut represents a “bottleneck” in the network—removing the cut edges disconnects s from t , so no flow can pass. The capacity of the minimum cut therefore provides an upper bound on the maximum flow. Remarkably, this bound is always tight.

Theorem 11.9 (Max-Flow Min-Cut Theorem) *In any flow network, the maximum value of a flow is equal to the minimum capacity of an s - t cut.*

Proof: Let f be a flow with maximum value and let (S, T) be any cut of the network, where $s \in S$ and $t \in T$. The value of the flow $|f|$ cannot exceed the capacity of the cut $c(S, T)$, since no more than $c(S, T)$ units of flow can cross from S to T .

When the FordFulkerson algorithm terminates, there are no augmenting paths in the residual graph. Define S as the set of vertices reachable from s in the residual graph, and $T = V \setminus S$. Since no augmenting paths remain, all edges from S to T in the original graph are saturated, and all edges from T to S carry no flow.

Therefore, the value of the flow equals the capacity of the cut (S, T) , proving that the maximum flow equals the minimum cut capacity. ■

According to the Max-Flow Min-Cut theorem, the value of the maximum flow equals the capacity of the minimum cut. Let us identify the minimum cut in our example by finding the set of vertices reachable from s in the final residual network.

Let us trace the vertices reachable from s in the final residual network. From s : we can reach a because $s \rightarrow a$ has residual capacity $16 - 13 = 3 > 0$. From a : we can reach c because $a \rightarrow c$ has residual capacity $4 - 1 = 3 > 0$. From c : the edge $c \rightarrow d$ is saturated (residual 0), so d is not reachable from c . No further vertices are reachable. So $S = \{s, a, c\}$ and $T = \{b, d, t\}$.

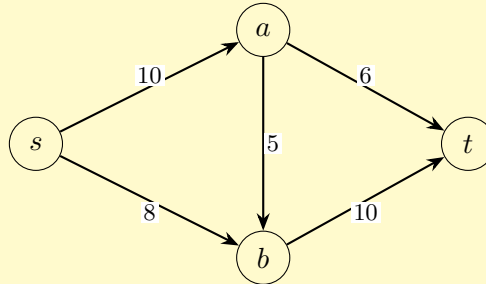
The cut edges (from S to T) are:

- (a, b) with capacity 12
- (c, d) with capacity 14

The total capacity is $12 + 14 = 26$, which is exactly equal to the maximum flow, confirming the Max-Flow Min-Cut theorem.

We give another example.

Example 11.10 Consider a simpler network with four vertices:



Iteration 1: Path $s \rightarrow a \rightarrow t$, bottleneck $\min\{10, 6\} = 6$. Flow = 6.

Iteration 2: Path $s \rightarrow a \rightarrow b \rightarrow t$, bottleneck $\min\{4, 5, 10\} = 4$. Flow = 10.

Iteration 3: Path $s \rightarrow b \rightarrow t$, bottleneck $\min\{8, 6\} = 6$. Flow = 16.

No more augmenting paths exist. Maximum flow = 16.

To find the minimum cut, we check which vertices are reachable from s in the final residual network. After the three augmentations, $s \rightarrow a$ has residual $10 - 10 = 0$ and $s \rightarrow b$ has residual $8 - 6 = 2$, so b is reachable. From b , $b \rightarrow t$ has residual $10 - 10 = 0$ and the reverse edge of $a \rightarrow b$ has residual 4, so a is reachable. From a , $a \rightarrow t$ has residual $6 - 6 = 0$. Thus $S = \{s, a, b\}$, $T = \{t\}$, and the minimum cut has capacity $c(a, t) + c(b, t) = 6 + 10 = 16$, matching the max flow.

11.5 EdmondsKarp Algorithm

Figure 11.1 showed that poor choices of augmenting paths can cause the FordFulkerson algorithm to run for $\Theta(f^*)$ iterations, where f^* can be exponential in the input size. The EdmondsKarp modification fixes this by always choosing a *shortest* augmenting path (fewest edges), found via breadth-first search (BFS) in the residual graph.

Algorithm EdmondsKarp: EdmondsKarp Algorithm

Input: Graph $G = (V, E)$, capacities c , source s , sink t

Output: Maximum flow f

```

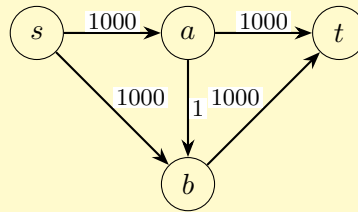
1 Initialize  $f(u, v) \leftarrow 0$  for all  $(u, v) \in E$ ;
2 while there exists an augmenting path in  $G_f$  do
3   Use BFS to find shortest path  $p$  from  $s$  to  $t$  in  $G_f$ ;
4   Compute  $c_f(p) = \min_{(u, v) \in p} c_f(u, v)$ ;
5   for each edge  $(u, v) \in p$  do
6     if  $(u, v) \in E$  then  $f(u, v) \leftarrow f(u, v) + c_f(p)$  // Increase flow ;
7     else  $f(v, u) \leftarrow f(v, u) - c_f(p)$  // Decrease flow in the reverse edge ;
8   end
9 end
10 return  $f$ ;
  
```

The key insight is that using BFS guarantees that the distance (in number of edges) from s to any vertex in the residual graph never decreases across augmentations. Moreover, after at most $O(|V| \cdot |E|)$ augmentations, the algorithm terminates.

Theorem 11.11 *The EdmondsKarp algorithm runs in $O(|V| \cdot |E|^2)$ time.*

This is a *strongly polynomial* bound—it depends only on $|V|$ and $|E|$, not on the magnitudes of capacities.

Example 11.12 Consider the network from Figure 11.1 with $M = 1000$:



With arbitrary path selection, FordFulkerson might take 2000 iterations. With BFS (EdmondsKarp), the shortest paths from s to t are $s \rightarrow a \rightarrow t$ and $s \rightarrow b \rightarrow t$, each with 2 edges. The algorithm chooses one of these, sending 1000 units, then the other, sending 1000 units. Total: 2 iterations, giving maximum flow 2000.

When all capacities in a flow network are integers, the FordFulkerson algorithm (and hence the EdmondsKarp algorithm) always produces an integer-valued maximum flow.

Theorem 11.13 (Integrality Theorem) *If all capacities in a flow network are integers, then there exists a maximum flow in which the flow on every edge is an integer.*

Proof: The FordFulkerson algorithm starts with the zero flow (integer-valued). In each iteration, the bottleneck capacity of the augmenting path is the minimum of integer residual capacities, hence an integer. Augmenting by an integer amount preserves the integrality of all flow values. Since the algorithm terminates with a maximum flow, this maximum flow is integer-valued. ■

This theorem has important combinatorial consequences. For instance, it implies that the maximum number of edge-disjoint s - t paths equals the minimum number of edges whose removal disconnects s from t (Menger's theorem), since we can model edge-disjoint paths as a max-flow problem with unit capacities.

11.6 Applications of Max-Flow

The max-flow problem reduces several classical combinatorial problems to a single algorithmic primitive. We illustrate two such reductions; both produce a flow network with unit capacities, so by the integrality theorem (Theorem 11.13) the resulting maximum flow is integer-valued and corresponds directly to a combinatorial answer.

Bipartite Matching

Given a bipartite graph $G = (L \cup R, E)$ with edges only between L and R , a *matching* is a set of edges no two of which share an endpoint. The *maximum bipartite matching* problem asks for a matching of largest size.

To reduce to max-flow, build a directed network $G' = (V', E')$ with $V' = L \cup R \cup \{s, t\}$. For every $\ell \in L$ add an edge (s, ℓ) , for every $r \in R$ add an edge (r, t) , and orient every edge $(\ell, r) \in E$ from L to R . Give every edge unit capacity (Fig. 11.2). A maximum integer-valued flow in G' corresponds exactly to a maximum matching in G : an edge (ℓ, r) is in the matching iff its flow is 1, since flow conservation and unit capacities at ℓ and r ensure that each vertex is matched at most once.

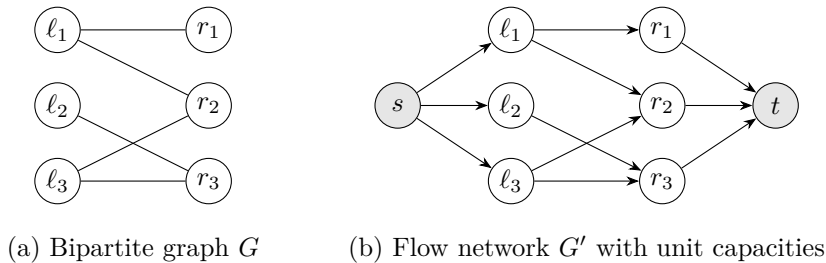


Figure 11.2: Reduction from bipartite matching to max-flow. Every edge in (b) has capacity 1. A maximum flow corresponds to a maximum matching in (a).

If the maximum flow has value $|f^*|$, then G has a matching of size $|f^*|$, and conversely. Running EdmondsKarp on G' thus solves bipartite matching in $O(VE^2)$ time, with sharper bounds available via Hopcroft-Karp.

Edge-Disjoint Paths from s to t

Given a directed graph $G = (V, E)$ and two vertices $s, t \in V$, an *edge-disjoint s - t path collection* is a set of paths from s to t no two of which share an edge. The problem asks for the largest such collection.

To reduce to max-flow, assign capacity 1 to every edge of G (Fig. 11.3). The maximum number of edge-disjoint s - t paths equals the value of a maximum integer flow: any flow of value k decomposes into k edge-disjoint paths (because each edge carries flow 0 or 1 and flow conservation routes exactly k units from s to t); conversely, k edge-disjoint paths give a flow of value k . Combined with the Max-Flow Min-Cut Theorem this is the edge form of *Menger's Theorem*: the maximum number of edge-disjoint s - t paths equals the minimum number of edges whose removal disconnects s from t .

The same reduction yields the *vertex-disjoint* version with a standard vertex-splitting trick: replace every internal vertex v by two copies $v_{\text{in}}, v_{\text{out}}$ joined by a single capacity-1 edge, and route all incoming edges into v_{in} and all outgoing edges out of v_{out} . A unit flow that uses two paths through the same vertex would saturate that internal edge, so flow conservation again limits each vertex to one passing path.

11.7 LLP Perspective

In this section we recast the maximum-flow / minimum-cut machinery in the lattice-linear predicate framework of Chapter 2. We first show that the property of being a minimum s - t cut is itself a lattice-linear

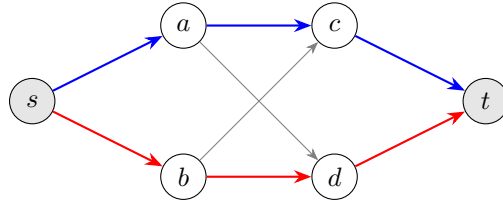


Figure 11.3: Edge-disjoint paths from s to t in a graph with all unit-capacity edges. The blue and red paths are edge-disjoint; gray edges show two further unit-capacity edges that are not used. The minimum s - t edge cut has size 2 (e.g., the two edges out of s), matching the maximum number of edge-disjoint paths.

predicate, so the set of minimum cuts forms a finite distributive lattice. This places the *constrained-mincut problem* — finding a mincut that also satisfies a side predicate B — within reach of the LLP main loop. The resulting Algorithm [LLP-MinCut](#) runs in $O(MF(n, m) + km)$ time, where $MF(n, m)$ is the time to compute one max-flow and k is a complexity parameter of B (the number of times B flips along any chain in the cut lattice).

Lattice of Mincuts

We first show that the property of a cut (S, T) being a mincut is lattice-linear. A set $S \subseteq V$ is a mincut if:

- $s \in S, t \notin S$ (i.e., S is a cut).
- $\text{val}(S) = \sum_{(u,v) \in E: u \in S, v \in V \setminus S} c(u, v) \leq \text{val}(S')$ for all cuts S' .

We can now show:

Lemma 11.14 *The predicate $B(S) = (s \in S \text{ and } t \notin S)$ and $(\text{val}(S) \text{ is minimum})$ is lattice-linear with efficient advancement.*

Proof: We use submodularity of the cut function. For min cuts S_1, S_2 , the submodularity inequality is:

$$\text{val}(S_1) + \text{val}(S_2) \geq \text{val}(S_1 \cup S_2) + \text{val}(S_1 \cap S_2).$$

If $\text{val}(S_1) = \text{val}(S_2) = \mu$ (the min cut capacity), then $\text{val}(S_1 \cup S_2) + \text{val}(S_1 \cap S_2) \leq 2\mu$. Since $\text{val}(S) \geq \mu$ for any cut, each term must equal μ . Thus, $S_1 \cap S_2$ and $S_1 \cup S_2$ are min cuts (noting $s \in S_1 \cap S_2, t \notin S_1 \cup S_2$).

For the advancement property, suppose S is not a min cut (e.g., $\text{val}(S) > \mu$). If S equals $V - \{t\}$, then we know that there is no $S' \supseteq S$ such that S' is a mincut and thus all indices are forbidden. Otherwise, there is at least some vertex v such that there exists an edge $(u, v), u \in S, v \notin S$, with residual capacity $c(u, v) > f(u, v)$. If v equals t , we know that there is no $S' \supseteq S$ such that S' is a mincut and thus all indices are forbidden. Otherwise, unless we advance by including v in S , the cut cannot be a mincut. ■

LLP-MinCut Algorithm

Since finding a mincut satisfying an arbitrary B is NP-complete in general, we focus on special cases of B . We first assume that the given predicate B is lattice-linear. This means that the set of cuts that satisfy

B form an inf-semilattice and we have the efficient advancement property. Since the set of mincuts form a sublattice, we get that the set of mincuts that satisfy B also forms an inf-semilattice.

We now have a simple algorithm to find a mincut that satisfies B .

Algorithm LLP-MinCut: To find the minimum vector that satisfies B

Input: B : lattice-linear predicate on cuts
Output: G : least mincut satisfying B , or null

```

1  $G := \text{LeastMincut}$ ;
2 while  $\neg B(G)$  do
3   while  $\exists j : \text{forbidden}(G, j, B)$  do
4     if  $G[j] = 1$  then return null;
5     else  $G[j] := 1$ ;
6   end
7   if there does not exist  $\text{LeastMincut} \geq G$  then return null;
8   else  $G := \text{LeastMincut} \geq G$ ;
9 end
10 return  $G$ ;

```

// no such cut

// the optimal solution

The algorithm alternates between finding the mincut after G and finding G that satisfies B . If it cannot find a mincut or a cut satisfying B , it returns null. Assuming that the complexity of computing $\exists j : \text{forbidden}(G, j, B)$ is $O(m)$, the above algorithm takes $O(MF(n, m) + mn)$ given the join-irreducible elements of the mincut lattice. The outer while loop executes at most n times because at least one bit in the vector G becomes 1 after every iteration.

We now show that for many lattice-linear predicates, the complexity of the above algorithm is $O(MF(n, m) + kn)$ where k is a parameter dependent upon the predicate B .

To concretize our discussion, consider the predicate $B(S)$ that holds whenever either both vertices u and v belong to S or neither belong to S . We are interested in finding S such that S is a mincut and satisfies $B(S)$.

We note here that the predicate, when evaluated over increasing sets of S , can transition at most twice. Either it is initially false and then turns true. This happens when S initially has u or v , and then later it gets both. Alternatively, it may be initially true, then turn false and then finally turn true. This happens when initially, neither u nor v is in S . Then, one of them gets added to S and finally both of them get added.

The predicate B that can transition at most k times is termed a k -Transition predicate. Formally,

Definition 11.15 A predicate $B : J(P) \rightarrow \{true, false\}$ on the ideals of a poset P is a k -Transition Predicate if, for any chain of ideals $I_1 \subseteq I_2 \subseteq \dots \subseteq I_n \in J(P)$, the sequence $B(I_1), B(I_2), \dots, B(I_n)$ has at most k transitions, where a transition occurs at index i if $B(I_i) \neq B(I_{i+1})$.

For example, the *stable* predicates [CL85] are 1-Transition predicates. Similarly, predicates that are true on a single mincut [GS24] are also 1-Transition predicates. We will assume that the algorithm for detecting whether B is true on a cut S is $O(m)$.

Thus, if the lattice-linear predicate is a k -transition predicate, then the time complexity is bounded by $O(MF(n, m) + km)$.

11.8 Summary

This chapter presented algorithms for the maximum flow problem in directed networks and established the duality between maximum flows and minimum cuts. Two unit-capacity reductions — bipartite matching and edge-disjoint s - t paths — show that max-flow is the algorithmic engine behind several classical combinatorial problems. The lattice of minimum cuts is itself a finite distributive lattice, allowing the LLP framework to extend max-flow with side constraints expressed as lattice-linear predicates.

Table 11.1 summarizes the algorithms discussed in this chapter.

Problem	Algorithm	Time Complexity
Maximum Flow	FordFulkerson	$O(f^* E)$
Maximum Flow	EdmondsKarp	$O(V \cdot E ^2)$
Bipartite Matching	via EdmondsKarp	$O(VE^2)$
Edge-disjoint s - t paths	via EdmondsKarp	$O(VE^2)$
Constrained Mincut	LLP Mincut	$O(MF(n, m) + km)$

Table 11.1: Algorithms discussed in this chapter.

Here f^* is the value of the maximum flow, $|V|$ and $|E|$ are the number of vertices and edges, $MF(n, m)$ denotes the time to compute a maximum flow, and k is a parameter dependent on the predicate.

11.9 Problems

1. Consider the flow network below, with capacities on edges. Use the FordFulkerson algorithm to compute the maximum flow from s to t . Show one possible sequence of augmenting paths and the final flow value.

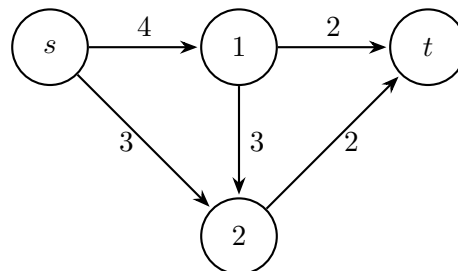
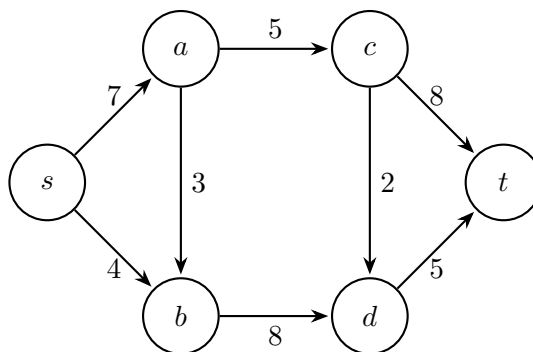


Figure 11.4: Flow network with capacities.

2. Let $G = (V, E)$ be a directed graph and let s and t be two nodes in V . Describe an efficient algorithm to find all the edge-disjoint paths from s to t in G . (Edge-disjoint means that none of the paths have any edges in common; they are allowed to have vertices in common.)

3. Suppose you are given a flow network $G = (V, E)$ with source s , sink t , integer capacities, and a maximum flow f of G along with the residual graph G_f . There are n vertices and m edges in the flow network.
 - (a) We increase the capacity of a single edge $(u, v) \in E$ by one. Give an $O(m + n)$ time algorithm to update the maximum flow.
 - (b) We decrease the capacity of a single edge $(u, v) \in E$ by one. Give an $O(m + n)$ time algorithm to update the maximum flow.
4. (**Vertex capacities.**) In some applications, vertices also have capacity constraints. Given a flow network $G = (V, E)$ where each vertex $v \in V \setminus \{s, t\}$ has a capacity $c(v)$ limiting the total flow through v , show how to reduce this problem to a standard maximum flow problem (where only edges have capacities).
5. (**Multiple sources and sinks.**) Suppose a flow network has multiple sources s_1, s_2, \dots, s_k and multiple sinks t_1, t_2, \dots, t_l . Show how to reduce this to a single-source, single-sink max-flow problem.
6. (**Minimum cut.**) Find the minimum s - t cut in the following network. Verify your answer by finding a flow of the same value.



7. (**Unique minimum cut.**) Prove or disprove: a flow network always has a unique minimum cut.
8. Let (L, \leq) be a finite distributive lattice. Show that there exists a directed graph G with distinguished vertices s and t , and a capacity function c on the edges, such that the lattice of minimum s - t cuts of G (ordered by the set of vertices reachable from s) is isomorphic to L .

11.10 Bibliographic Remarks

The maximum flow problem and its solutions have been extensively studied in combinatorial optimization and network theory. The original FordFulkerson algorithm was introduced in 1956 [JF56], together with the foundational Max-Flow Min-Cut Theorem that proves its correctness. The Dinitz algorithm [Din70] and the algorithm by Edmonds and Karp [EK72] offer strongly polynomial time guarantees. More advanced algorithms such as Goldberg and Tarjan's push-relabel method [GT88] provide even better performance in practice and theory, particularly for dense networks. For comprehensive treatments of flow algorithms, see Ahuja, Magnanti, and Orlin [AMO93], Schrijver [Sch03], and Korte and Vygen [KV18].

The applications discussed in Section 11.6 have rich histories of their own. The reduction of bipartite matching to max-flow goes back to König's theorem (1931). Hopcroft and Karp [HK73] obtained an $O(E\sqrt{V})$ algorithm for bipartite matching that exploits the unit-capacity structure to find blocking flows. The connection between max-flow and edge-disjoint paths predates max-flow itself: it is the algorithmic content of Menger's theorem [Men27]. The integrality theorem and its consequences are surveyed in detail in Schrijver [Sch03].

The lattice structure of minimum cuts of Section 11.7 was characterised by Picard and Queyranne [PQ80]; the LLP framework's treatment in this chapter views their structure through the lens of lattice-linear predicates and reduces constrained-mincut problems to a single max-flow plus lattice traversal.

Chapter 12

Bipartite Matching

12.1 Introduction

In this chapter, we discuss algorithms for matching problems in bipartite graphs and related combinatorial optimization problems. Figure 12.1 shows the relationships between various structures studied in this chapter. The three rows correspond to three classical min-max theorems: König’s theorem relates matching and vertex cover, Dilworth’s theorem relates chain decomposition and antichain, and Menger’s theorem relates vertex-disjoint paths and vertex cuts. The vertical arrows show reductions between these problems.

The matching problem is one of the most studied problems in combinatorial optimization, with applications ranging from job assignment and resource allocation to network design and computational biology. Bipartite matching, in particular, is a building block: many problems that appear unrelated — finding a minimum vertex cover, partitioning a poset into the fewest chains, exhibiting a maximum antichain, computing vertex-disjoint paths between two terminals — all reduce to bipartite matching by reductions covered in this chapter. The classical augmenting-path algorithm therefore deserves a careful treatment, both because it solves matching directly in $O(nm)$ time and because the same algorithm, run on a problem-specific transformation, settles each of those four neighbouring problems. The chapter develops the algorithm first, then makes the reductions explicit so that readers see why a single technique unlocks all of Figure 12.1.

This chapter is organized as follows. Section 12.2 describes the classical sequential augmenting path algorithm for maximum cardinality matching in bipartite graphs. Section 12.3 shows how to reduce chain partition of a poset to bipartite matching via the strict split construction, and gives an LLP algorithm for finding the maximum antichain (Dilworth’s theorem). Section 12.4 shows how to convert a maximum matching into a minimum vertex cover, establishing König’s theorem algorithmically. Section 12.7 presents Menger’s theorem on vertex-disjoint paths and vertex cuts in directed graphs, and shows how it relates to bipartite matching via poset reductions.

12.2 A Bipartite Matching Algorithm

Given an undirected graph (V, E) , a *matching* M is a subset of edges E such that no two edges are incident on the same vertex. As an example, consider a bipartite graph (L, R, E) such that the vertex set L denotes a set of men, R denotes a set of women and E denotes a symmetric “likes” relation. Then, a matching

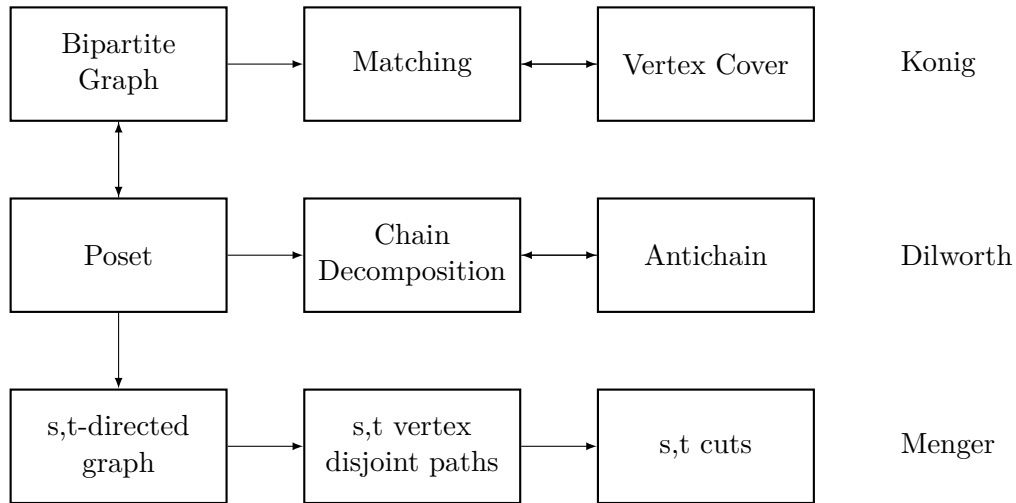


Figure 12.1: Various Structures and Transformations between them

M is simply a set of edges such that no man or woman is adjacent to two edges in M . If the number of men and the number of women are equal to n and every man is matched, then we call that matching a *perfect matching*. The perfect matching may not always exist. The following famous Theorem gives a necessary and sufficient condition for a perfect matching to exist in a bipartite graph. It is based on the notion of *overdemanded* subset. We say that $L' \subseteq L$ is overdemanded if there exists $R' \subseteq R$ such that the set of neighbors of R' in the graph is L' and the size of R' is strictly bigger than $|L'|$. Intuitively, L' is overdemanded because R' can only match with vertices in L' but the size of L' is smaller than the size of R' . We can now state the Theorem.

Theorem 12.1 (Hall's Theorem) *The necessary and sufficient condition for a perfect matching to exist in a bipartite graph (L, R, E) where $|L| = |R|$, is that L does not have any overdemanded subset L' .*

Observe that even though Hall's theorem gives us insight about the perfect matching, it cannot be directly applied to efficiently check for perfect matching because there are 2^n possible subsets of L .

We now give an efficient algorithm for a more general problem called the *maximum cardinality matching problem*. This problem requires us to find a matching of the largest size. When the number of men and the number of women are equal to n , then a perfect matching exists iff the size of the maximum cardinality matching is n . The algorithm is based on the idea of increasing the size of a matching by using an *augmenting path*. Suppose that the algorithm has a matching M_t in iteration t . The algorithm is then guaranteed to find a matching M_{t+1} of size one bigger than M_t whenever a matching with a larger size exists. If we can make this idea work, then we can start with M_0 as the empty matching and then keep applying the idea to reach a maximum cardinality matching.

To understand the idea of an augmenting path for a matching M , we define a vertex as *exposed* if it is not incident on any edge in M . Now suppose that there exists a path from an exposed vertex to another exposed vertex that alternates between unmatched and matched edges. Such a path must start with an unmatched edge and end with an unmatched edge (by the definition of exposed vertices). Furthermore,

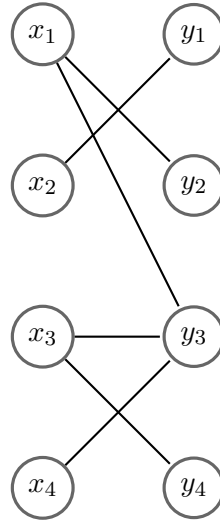


Figure 12.2: A Bipartite Graph

such a path has an odd number of edges, with the unmatched edges exactly one more than the matched edges. Then, by simply switching the matched with unmatched edges along that path, we can increase the size of matching by one.

Algorithm Seq-AugmentingPath: Finding a Maximum Cardinality Matching

Input: (L, R, E) : bipartite graph

Output: M : maximum cardinality matching

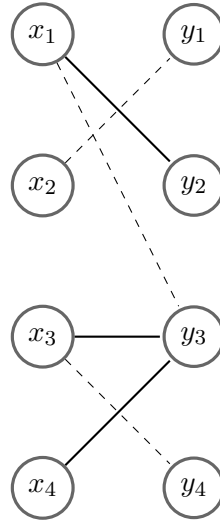
```

1  $M := \{\}$ ;
2 while there exists an alternating path  $P$  in  $(L, R, E)$  for  $M$  do
3   |  $M :=$  matching  $M$  with edges switched in the path  $P$ ;
4 end
5 return  $M$ ;

```

How do we find an alternating path? One can start with any exposed vertex and do a breadth-first-search such that the BFS tree alternates between unmatched and matched edges. Such a search will result in either finding an alternating path or an overdemand set. In our example of the matching M in Fig. 12.3, if we start from x_4 , we get the alternating path x_4, y_3, x_1, y_2 . If there are $2n$ vertices and m edges, every iteration of the *while* loop takes $O(m)$ time for breadth-first-search and there can be at most n iterations, giving us the time complexity of $O(nm)$. By augmenting along multiple paths in every iteration, the complexity can be reduced to $O(\sqrt{nm})$ [HK71].

We now give a sequential algorithm for a slight variant of the matching problem in a bipartite graph (L, R, E) . Let the vertices of L be numbered $1..n$. Let L_i denote the set of vertices $\{v_1, v_2, \dots, v_i\}$, for all $1 \leq i \leq n$. The output of our algorithm is a vector G such that $\sum_i G[i]$ is the size of the maximum matching in the graph (L_i, R, E_i) where E_i are the edges restricted to the set $L_i \cup R$. For simplicity, we set L_0 to \emptyset . We let S be the matched vertices in R . It is sufficient to describe the sequential algorithm when a new vertex v_i is added to the left side. We simply look for an alternating path from v_i to any unmatched vertex in R . If we find any such path, then the newly matched vertex in R is added to S . If there is no

Figure 12.3: A Matching M shown with dashed edges in the Bipartite Graph

path, then L_i is a constricted set, and the set of vertices in R matched to L_{i-1} is identical to the set of vertices matched to L_i . In either case, we continue the procedure to the next vertex in L , if any.

Algorithm BipartiteMatching: A Sequential Algorithm for Bipartite Matching

Input: (L, R, E) : bipartite graph

Output: $G[1..n]$: matching indicator; $S[1..n]$: partner array

```

1  $G$ : array[1.. $n$ ] of int initially  $\forall j : G[j] := 0$ ;
2  $S$ : array[1.. $n$ ] of 0.. $n$  initially  $\forall j : S[j] := 0$ ;
3 for  $j := 1$  to  $n$  do
4   if there exists an alternating path from  $v_j$  to any vertex  $k$  in  $R$  with  $(S[k] = 0)$  then
5     |   Assign  $S[]$  in the alternating path;
6     |    $G[j] := 1$ ;
7   end
8 end
9 return  $G$ ;

```

Algorithm [BipartiteMatching](#) searches the element in the boolean lattice B_n such that $G[i]$ equals 1 if the size of the bipartite matching for (L_i, R, E_i) is one more than for (L_{i-1}, R, E_{i-1}) , and 0 otherwise. The time complexity of the algorithm is $O(nm)$ because the *for* loop has n iterations and each iteration takes $O(m)$ to search for a path. The matching itself is stored in all nonzero S entries.

A key advantage of Algorithm [BipartiteMatching](#) is that it is a *deterministic* algorithm. Given a labeling of indices in L , it produces a single fixed G .

12.3 Chain Partition of a Poset

In this section, we show that maximum matching in a bipartite graph also allows us to partition a poset into the minimum number of chains.

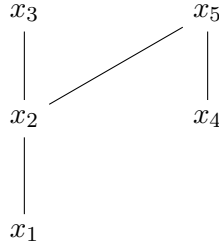


Figure 12.4: A poset

Assume that we have an algorithm for bipartite matching. How do we use it for chain decomposition? This relationship between chain decomposition and bipartite matching is based on the idea of a strict split of a poset.

Definition 12.2 (Strict Split of a Poset) *Given a poset P , the strict split of the poset P , denoted by $S(P)$, is a bipartite graph (L, R, E) where $L = \{x^- | x \in P\}$, $R = \{x^+ | x \in P\}$, and $E = \{(x^-, y^+) | x < y \text{ in } P\}$.*

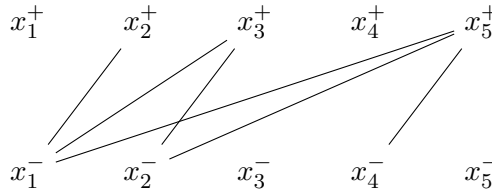


Figure 12.5: A Strict Split of the Poset in Fig. 12.4

We state the following theorem due to Fulkerson [Ful56].

Theorem 12.3 [Ful56] *Any matching in $S(P)$ can be reduced to a chain cover of P .*

Proof: We start with the trivial chain partition of P with each element in P being a chain. Thus, initially, we have n chains for n elements. We now consider each edge (x^-, y^+) in the matching. Each such edge implies that $x < y$ in P and we combine the chains corresponding to x and y . Since there can be at most one edge incident on x^- in a matching, x is the top element of its chain and similarly y is the lowest element of the chain. Therefore, these chains can be combined into one chain. If there are e edges in the matching, after this procedure we have a chain cover with $n - e$ chains. ■

at least one of the vertices in $\{u, v\}$ to cover all the edges in M . Now, the only question is which of the vertices $\{u, v\}$ do we choose to be part of the vertex cover. The answer is quite simple. If there is any vertex x adjacent to u such that x is not part of the matching M , then u would have to be part of the vertex cover. Otherwise, the edge (x, u) would not be covered. Can we have a vertex x that is adjacent to both u and v ? No, because this would form an odd size cycle that cannot be present in a bipartite graph. Finally, could we have two vertices x and y that are not part of the matching M such that x is adjacent to u and y is adjacent to v ? In this case, M is not a matching of maximum size because we can include (x, u) and (v, y) instead of just the edge (u, v) .

Algorithm VertexCoverFromMatching: A Parallel Algorithm to Convert a Matching to a Vertex Cover in a Bipartite Graph

Input: (L, R, E) : bipartite graph; M : matching in the bipartite graph (subset of E)
Output: $C[1..n]$: vertex cover set

```

1   $C$ : array[1.. $n$ ] of boolean initially  $\forall j : C[j] := 0$ ; // vertex cover set
2   $partner$ : array[1.. $n$ ] of int initially  $\forall j : partner[j] := 0$ ;
3  for  $(u, v) \in M$  in par (assume  $u < v$ ) do
4       $C[u] := true$ ;
5       $partner[u] := v$ ;
6       $partner[v] := u$ ;
7  end
8  for  $(u, v) \in E - M$  in par (assume  $u < v$ ) do
9      if  $\neg C[u] \wedge \neg C[v]$  then
10         if  $partner[u] \neq 0$  then
11              $C[partner[u]] := false$ ;
12              $C[u] := true$ ;
13         else
14             //  $partner[u] = 0$  implies  $partner[v] \neq 0$ 
15              $C[partner[v]] := false$ ;
16              $C[v] := true$ ;
17         end
18     end
19 return  $C$ ;
```

Algorithm [VertexCoverFromMatching](#) gives an algorithm to convert a bipartite matching into a vertex cover. It takes as input a bipartite graph and a matching M . The boolean array C gives the vertex cover. The variable array $partner$ is used to mark partner of any vertex in matching M . If a vertex u has no partner then $partner[u]$ is zero. We initialize the $partner$ array to zero. The algorithm first sets the partner for each vertex by going over all the edges in M . It also adds the lower numbered vertex u in the vertex cover for every edge (u, v) in M . This can be done in parallel in $O(1)$ time. At this point, we have exactly one vertex in C for every edge in M . The set C covers all edges in M , but may not cover all other edges.

In the second step, we ensure that all edges are covered. We maintain the invariant that there is exactly one vertex in C for every edge in M . We go over all edges (u, v) in parallel where u is less than v . If neither of the endpoints of the edge are in C , then we consider two cases. From the invariant, we know

that the edge (u, v) is not in matching M . If u is matched to some vertex w in M , i.e., $\text{partner}[u]$ equals w , then we remove w from C and add u to C . Otherwise, v is matched to some vertex w' in M . In this case, we remove w' from C and add v to C . If neither u nor v is matched, then M is clearly not a maximum matching since it can be increased in size by adding the edge (u, v) . Observe that we maintain the invariant that there is exactly one vertex in C for every edge in M .

We now show that in the second step it cannot happen that for a matched edge $(u, w) \in M$, some edge requires $C[u]$ be set to 1 and some other edge requires $C[w]$ to be set to 1. We set $C[u]$ to 1 only if there is an edge (u, v) which is not in M . Similarly, we set $C[w]$ to 1 only if there is an edge (w, w') which is not in M .

Example 12.6 [Running example, continued] For the strict-split graph of Example 12.4 and the matching M^* , Algorithm [VertexCoverFromMatching](#) produces a minimum vertex cover of size 3, e.g. $\{x_1^-, x_2^-, x_5^+\}$: x_1^- covers the three edges out of x_1^- , x_2^- covers the two remaining edges out of x_2^- , and x_5^+ covers (x_4^-, x_5^+) . By König's theorem, no smaller vertex cover exists, since the maximum matching has size 3.

12.5 Dilworth's Theorem and Maximum Antichain

We now establish the relationship between the chain decomposition of a poset and the maximum antichain. Recall from Section 12.3 that the minimum chain cover of a poset P with n elements can be obtained from the maximum matching in the strict split $S(P)$. If the maximum matching has e edges, then the minimum chain cover has $n - e$ chains.

Definition 12.7 (Antichain) *An antichain in a poset (P, \leq) is a set of elements $A \subseteq P$ such that no two elements in A are comparable, i.e., for all $x, y \in A$ with $x \neq y$, neither $x \leq y$ nor $y \leq x$.*

Theorem 12.8 (Dilworth's Theorem [Dil50]) *In any finite poset, the minimum number of chains needed to cover all elements equals the maximum size of an antichain.*

Proof: Let m be the minimum number of chains in a chain cover and let a be the maximum size of an antichain.

First, we show $m \geq a$. Any antichain has at most one element from each chain (since elements in the same chain are comparable). Hence, $a \leq m$.

Second, we show $m \leq a$ by constructing an antichain of size m . Given a minimum chain cover with m chains C_1, C_2, \dots, C_m , we use the LLP algorithm (Algorithm [LLP-Antichain](#)) to find an antichain. The algorithm maintains a vector G where $G[j]$ is an index into chain C_j . Let $C_j[G[j]]$ denote the $G[j]$ -th element in chain C_j from the bottom. The predicate $\text{forbidden}(j)$ is true if there exists another chain k such that $C_j[G[j]] \geq C_k[G[k]]$ in the poset. The advance operation moves $G[j]$ to the next element in the chain. When no index is forbidden, the set $\{C_1[G[1]], C_2[G[2]], \dots, C_m[G[m]]\}$ forms an antichain of size m .

We now verify that the algorithm terminates with a valid antichain. If $G[j]$ is forbidden because $C_j[G[j]] \geq C_k[G[k]]$, then advancing $G[j]$ to the successor in chain j removes this comparability (the

successor is strictly greater than $C_j[G[j]]$ and hence not below $C_k[G[k]]$. Since each chain is finite, the algorithm terminates. At termination, no two elements in the selected set are comparable, giving us an antichain of size m .

■

Example 12.9 [Running example, continued] The two-chain cover from Example 12.4 is $C_1 = (x_1, x_2, x_3)$, $C_2 = (x_4, x_5)$. Algorithm [LLP-Antichain](#) starts with $G = [1, 1]$, so the representatives are x_1 and x_4 . These two elements are incomparable in P , so neither index is forbidden, and the algorithm terminates immediately with the maximum antichain $\{x_1, x_4\}$ of size $2 = n - e$. This confirms Dilworth's theorem on the running example.

Combining Dilworth's theorem with Fulkerson's theorem (Theorem 12.3), we get that for a poset with n elements whose strict split has maximum matching of size e :

- Minimum chain cover size = $n - e$
- Maximum antichain size = $n - e$
- Minimum vertex cover in $S(P) = e$ (by König's theorem)

This establishes the correspondence shown in the second row of Figure 12.1: the chain decomposition and antichain of a poset are dual to each other via Dilworth's theorem, just as matching and vertex cover are dual via König's theorem.

12.6 Reductions Between Structures

Figure 12.1 shows multiple reductions between combinatorial structures. We now justify each arrow in the figure.

Bipartite Graph \leftrightarrow Poset

Given a bipartite graph (L, R, E) , we can construct a poset as follows. Let the elements of the poset be $L \cup R$ and define the partial order such that $x < y$ iff $x \in L$, $y \in R$, and $(x, y) \in E$. Conversely, given a poset P , its strict split $S(P)$ gives a bipartite graph.

More precisely, the reduction from posets to bipartite graphs uses the strict split (Definition in Section 12.3): given a poset (P, \leq) , we construct the bipartite graph $S(P) = (L, R, E)$ where each element x appears as x^- in L and x^+ in R , with edge (x^-, y^+) whenever $x < y$ in P .

The key property is that a matching of size e in $S(P)$ corresponds to combining e pairs of consecutive elements in chains, yielding a chain cover of size $n - e$ (Theorem 12.3).

Matching \leftrightarrow Vertex Cover (König's Theorem)

The arrow from matching to vertex cover is justified by Algorithm [VertexCoverFromMatching](#) in Section 12.4: given any maximum matching, we can construct a minimum vertex cover of the same size in $O(1)$ parallel time.

The arrow from vertex cover to matching follows because any vertex cover of size k must include at least one endpoint of each matched edge. Hence, the vertex cover size is at least the matching size. Combined with König's theorem, a minimum vertex cover immediately certifies the optimality of a maximum matching.

Chain Decomposition \leftrightarrow Antichain (Dilworth's Theorem)

Given a minimum chain cover of size m , Algorithm [LLP-Antichain](#) produces a maximum antichain of size m using an LLP computation. Conversely, an antichain of size a certifies that any chain cover requires at least a chains (since each chain contains at most one antichain element).

Poset \rightarrow s, t -Directed Graph

Given a poset (P, \leq) , we construct a directed graph as follows. For each element $x \in P$, create two vertices x_{in} and x_{out} connected by a directed edge (x_{in}, x_{out}) with capacity 1. For each relation $x < y$ in P , add an edge (x_{out}, y_{in}) with capacity 1. Add a source vertex s with edges to all x_{in} where x is a minimal element, and a sink vertex t with edges from all x_{out} where x is a maximal element.

In this construction, vertex-disjoint s - t paths correspond to chains in the poset, and a minimum s - t vertex cut corresponds to a maximum antichain. This gives an alternative proof of Dilworth's theorem via Menger's theorem.

12.7 Menger's Theorem

We now discuss the third row of Figure [12.1](#): the relationship between vertex-disjoint paths and vertex cuts in directed graphs.

Definition 12.10 (Vertex-Disjoint Paths) *Given a directed graph $G = (V, E)$ with two distinguished vertices s and t , a set of s - t paths is vertex-disjoint if no two paths share any internal vertex (i.e., any vertex other than s and t).*

Definition 12.11 (s - t Vertex Cut) *An s - t vertex cut is a set of vertices $C \subseteq V \setminus \{s, t\}$ whose removal disconnects t from s , i.e., there is no directed path from s to t in $G - C$.*

Theorem 12.12 (Menger's Theorem [[Men27](#)]) *In a directed graph $G = (V, E)$, the maximum number of vertex-disjoint s - t paths equals the minimum size of an s - t vertex cut.*

Proof: Let k be the maximum number of vertex-disjoint paths and c be the minimum vertex cut size.

First, $k \leq c$: any s - t vertex cut must contain at least one internal vertex from each vertex-disjoint path (otherwise that path would still connect s to t). Hence $c \geq k$.

Second, $k \geq c$: we reduce the problem to max-flow. Replace each internal vertex v by two vertices v_{in} and v_{out} with a directed edge (v_{in}, v_{out}) of capacity 1. Replace each original edge (u, v) by (u_{out}, v_{in}) with capacity ∞ (or n). The maximum flow from s_{out} to t_{in} in this network equals the maximum number of

vertex-disjoint s - t paths (since each internal vertex can be used at most once due to capacity 1). By the max-flow min-cut theorem, this equals the minimum cut, which corresponds to removing vertices (those with saturated vertex-edges) that disconnect s from t . ■

Given the reduction to max-flow described in the proof, we can compute the maximum number of vertex-disjoint s - t paths using any max-flow algorithm on the transformed network. Since all capacities are 0 or 1 on vertex edges, the max-flow is integral and equals the number of vertex-disjoint paths.

The sequential time complexity is $O(kn)$ where k is the number of vertex-disjoint paths (at most n), since each augmenting path takes $O(n)$ time in the unit-capacity network and there are k such paths.

Menger's theorem generalizes König's theorem. Given a bipartite graph (L, R, E) , construct a directed graph as follows: add a source s with edges to all vertices in L , direct all edges from L to R , and add a sink t with edges from all vertices in R . Split each vertex in $L \cup R$ into in/out pairs with unit capacity. Then:

- Vertex-disjoint s - t paths correspond to matched edges.
- A minimum s - t vertex cut corresponds to a minimum vertex cover.

Thus, König's theorem (max matching = min vertex cover) is a special case of Menger's theorem (max vertex-disjoint paths = min vertex cut) applied to the bipartite construction.

As described in Section 12.6, given a poset (P, \leq) , we construct a directed graph with vertex splitting. In this construction:

- Each s - t path passes through a sequence of elements $x_1 < x_2 < \dots < x_k$, corresponding to a chain.
- The maximum number of vertex-disjoint s - t paths equals the minimum chain cover size.
- A minimum s - t vertex cut corresponds to a maximum antichain: the elements whose removal disconnects all chains from source to sink.

By Menger's theorem, min chain cover = max antichain, which is exactly Dilworth's theorem. This provides a unified view: all three min-max results in Figure 12.1 (König, Dilworth, Menger) are instances of the max-flow min-cut theorem applied to appropriately constructed networks.

Edge-disjoint paths

There is a parallel statement of Menger's theorem for edges, which is in many ways the more natural form for the max-flow / min-cut framework.

Theorem 12.13 (Edge-Menger) *In a directed graph $G = (V, E)$, the maximum number of edge-disjoint s - t paths equals the minimum number of edges in an s - t edge cut.*

Proof: Assign every edge unit capacity. By integrality of the max-flow on a unit-capacity network, the maximum flow value equals the maximum number of edge-disjoint s - t paths (each edge can carry flow 0 or 1, and conservation routes integer paths). By the max-flow min-cut theorem, this also equals the minimum capacity of any s - t cut, which on a unit-capacity graph is exactly the size of the smallest edge set whose removal disconnects s from t .



The vertex version (Theorem 12.12) reduces to the edge version by the standard vertex-splitting trick of the proof above; conversely, the edge version reduces to vertex-Menger on the line graph. Both forms thus express the max-flow / min-cut duality, specialised to unit-capacity networks of two flavours.

Example 12.14 [Running example, continued] Build the s - t graph for the running poset of Example 12.4 as follows. Split each x_i into x_i^-, x_i^+ as in the strict split, add a source s with unit-capacity edges to every x_i^- and a sink t receiving unit-capacity edges from every x_i^+ , and orient the strict-split edges from L to R . The maximum vertex-disjoint s - t paths in this graph have size 3, matching the maximum-matching size of $S(P)$:

$$s \rightarrow x_1^- \rightarrow x_2^+ \rightarrow t, \quad s \rightarrow x_2^- \rightarrow x_3^+ \rightarrow t, \quad s \rightarrow x_4^- \rightarrow x_5^+ \rightarrow t.$$

A minimum vertex cut of size 3 — the same $\{x_1^-, x_2^-, x_5^+\}$ from Example 12.4 continued — disconnects s from t , confirming Menger's theorem on the running example.

12.8 Summary

This chapter presented algorithms for bipartite matching and related combinatorial optimization problems, and established the connections between matching, vertex cover, chain decomposition, antichain, and vertex-disjoint paths through König's, Dilworth's, and Menger's theorems.

Problem	Algorithm	Time Complexity
Maximum Matching	Augmenting Path	$O(nm)$
Chain Partition	Reduction to Matching	$O(nm)$
Maximum Antichain	LLP-Antichain	$O(m^2)$
Minimum Vertex Cover	Matching to Vertex Cover	$O(n)$
Vertex-Disjoint Paths	Max-Flow Reduction	$O(kn)$

Here n is the number of vertices, m is the number of edges, and k is the number of vertex-disjoint paths.

12.9 Problems

1. Show that the minimum size of a vertex cover is equal to the maximum size of a matching in a bipartite graph.
2. Give a linear program formulation for the maximum matching problem and show that its dual corresponds to the minimum vertex cover problem.
3. Prove Hall's Theorem.
4. Edmonds' matrix is defined for a bipartite graph (U, V, E) with $|U| = |V| = n$ as follows. Let the vertices in U be u_1, u_2, \dots, u_n and the vertices in V be v_1, v_2, \dots, v_n . We set $A[i, j] = x_{i,j}$ if $(u_i, v_j) \in E$ and 0 otherwise. Show that Edmonds' matrix of a balanced bipartite graph has a perfect

matching iff the polynomial corresponding to the determinant of Edmonds' matrix is not identically zero.

5. We are given an unweighted graph with two distinguished vertices s and t . We are required to find the total number of paths from s to t such that these paths do not share any edge. Give an algorithm to find this number, and justify its correctness. Suppose that the graph has n vertices and m edges, give the time complexity of your algorithm.
6. Modify Algorithm [BipartiteMatching](#) to output the vertex cover in addition to bipartite matching.
7. Prove that a matching M in a bipartite graph is maximum if and only if there is no augmenting path with respect to M .
8. Prove that every k -regular bipartite graph (with $k \geq 1$) has a perfect matching. Conclude that the edge set can be partitioned into k perfect matchings.
9. (**Assignment / Min-Cost Bipartite Matching.**) Given a bipartite graph $G = (L, R, E)$ with $|L| = |R| = n$ and non-negative edge costs c_{uv} , formulate the minimum-cost perfect matching problem as a minimum-cost flow problem. Briefly describe the resulting algorithm and its running time.
10. (**Job-Assignment Variant.**) A company has n workers and $m \geq n$ jobs, where each worker can perform only a subset of the jobs. Each worker must be assigned to exactly one job; each job at most one worker. Give an efficient algorithm to decide whether such an assignment exists and compute it when it does.

12.10 Bibliographic Remarks

The sequential augmenting path algorithm, detailed in Section [12.2](#), is a cornerstone of matching theory, achieving a time complexity of $O(nm)$ for a graph with n vertices and m edges. This algorithm is enhanced by the seminal work of Hopcroft and Karp [[HK71](#)], who introduced a method to augment multiple paths per iteration, reducing the complexity to $O(\sqrt{nm})$. Their approach remains a standard for efficient bipartite matching. Hall's Theorem, presented in the chapter, is a fundamental result characterizing the existence of a perfect matching, with roots in early combinatorial optimization. For a deeper exploration of matching theory, including Hall's Theorem and its extensions, Lovasz and Plummer [[LP86](#)] offer a classic and authoritative reference. Section [12.3](#) connects bipartite matching to poset chain partitions through strict split construction, leveraging Fulkerson's theorem [[Ful56](#)]. Fulkerson's result elegantly reduces matchings in the bipartite graph $S(P)$ to chain covers in the poset P .

The König-Egervary theorem [[Kon31](#), [Ege31](#)] establishing the equality of maximum matching and minimum vertex cover in bipartite graphs is one of the earliest results in combinatorial optimization. Dilworth's theorem [[Dil50](#)] on the duality between chain covers and antichains in posets has deep connections to matching theory via the strict split construction. Menger's theorem [[Men27](#)] on vertex-disjoint paths predates both König's and Dilworth's results and provides a unified framework through which both can be derived as special cases of the max-flow min-cut theorem [[JF56](#)].

Chapter 13

Intractability

13.1 Introduction

This chapter discusses NP-completeness, a foundational concept in computational complexity that identifies the hardest problems in NP. We define P and NP, explore polynomial-time reductions, and prove NP-completeness for a broad set of problems: 3-SAT, Clique, Vertex Cover, Hamiltonian Path, Independent Set, and TSP. We also discuss methods to deal with NP-complete problems.

The classes P and NP classify computational problems by their solvability and verifiability. P includes problems with polynomial-time deterministic algorithms, while NP includes those verifiable in nondeterministic polynomial time. The unresolved question $P = NP$ drives much of complexity theory.

The problems are specified as requiring an *output* for the given *input*. We will use n for the size of the input. Our interest would be in determining the time required to compute the output. The output we require would be *binary*. For example, our input may be a Boolean expression B on Boolean variables. We would require the output to be 1 if the Boolean expression is satisfiable and 0 otherwise. At first, it may seem that in practical applications, the user may be interested in finding a satisfying assignment rather than simply that B is satisfiable. However, if someone gave us a method f to efficiently check whether B is satisfiable, then we can use f to determine a satisfying assignment. If B is satisfiable, then we can easily compute another Boolean expression B' in which the Boolean variable x_1 is set to true. We now ask whether B' is satisfiable. If B' is satisfiable, we continue with the other variables. If B' is not satisfiable, then we know that x_1 must be false. We compute a Boolean expression B'' from B by setting x_1 to false. Thus, in n applications of f , we would have found a satisfying assignment. This idea is applicable to all problems considered in this chapter, and we will restrict ourselves to such problems.

This chapter is organized as follows. Section 13.2 defines the complexity class **P**. Section 13.3 introduces polynomial-time reductions and illustrates them with Independent Set and Vertex Cover. Section 13.4 defines **NP** and NP-completeness, and proves NP-completeness for 3-SAT, Independent Set, Vertex Cover, Clique, and Subset Sum, plus a catalogue of further problems summarized in Figure 13.8. Section 13.5 discusses the **coNP** class. Section 13.6 surveys five strategies for coping with NP-hardness in practice; the approximation strategy is developed in depth in Chapter 14.

13.2 Class P

Definition 13.1 (Class P) *The complexity class \mathbf{P} is the set of all decision problems $L \subseteq \{0,1\}^*$ for which there exists a deterministic Turing machine M and a polynomial function $p : \mathbb{N} \rightarrow \mathbb{N}$ such that:*

1. M halts on every input $x \in \{0,1\}^*$ in at most $p(|x|)$ steps, where $|x|$ denotes the length of x ,
2. M accepts x if and only if $x \in L$.

Formally, $\mathbf{P} = \{L \mid \text{there exists a deterministic Turing machine } M \text{ and a polynomial } p \text{ such that } M \text{ decides } L \text{ in } p(n) \text{ steps for all } n \text{ where } n \text{ is the input size.}\}$

The notation used in the definition is as follows:

- \mathbf{P} : The complexity class of decision problems solvable in polynomial time by a deterministic Turing machine. The boldface emphasizes it as a formal class name.
- L : A language, representing a decision problem, which is a set of strings over the binary alphabet $\{0,1\}$.
- M : A deterministic Turing machine, a theoretical model of computation with a single, fixed sequence of steps for any input. For our purposes, we can use any practical programming language such as Java or C++.
- p : A polynomial function $p : \mathbb{N} \rightarrow \mathbb{N}$, mapping natural numbers (input sizes) to natural numbers (time bounds).

Informally, a problem L is in P if we can write a program that determines if the given instance x is in L in time that is polynomial in the size of x . Most of the problems that we have seen in this book so far belong to \mathbf{P} .

Example 13.2 The problem PATH of deciding whether vertex t is reachable from s in a graph is in \mathbf{P} ; BFS decides it in $O(n + m)$ time (Section 5.3).

13.3 Polytime Reductions

We define a relation between various problems as follows. We say that a problem π_1 is at least as hard as another problem π_2 if by using solutions to the problem π_2 we can solve the problem π_1 in time polynomial in the size of the problem.

Definition 13.3 (Polynomial-Time Reduction) π_2 is polynomially reducible to π_1 , denoted $\pi_2 \leq_P \pi_1$, if arbitrary instances of π_2 can be solved using a function that is a polynomial in the size of input and makes at most polynomial calls to a function that solves π_1 .

It is easy to verify that \leq_P is a reflexive and transitive relation.

A consequence of the above definition is that if there exists an efficient algorithm to solve π_1 , then we can use that solution to solve π_2 . Another consequence, important to this chapter, is that if π_2 is *hard* to solve, then π_1 is also hard to solve.

A template for NP-completeness proofs. Every NP-completeness proof in this chapter follows the same four-step pattern:

1. **Membership in NP.** Exhibit a short certificate for yes instances of the target problem Π that can be verified in polynomial time.
2. **Choose a source.** Pick a problem Π' already known to be NP-complete.
3. **Reduction.** Describe a polynomial-time construction f that maps every instance x of Π' to an instance $f(x)$ of Π .
4. **Correctness.** Prove both directions: $x \in \Pi' \Rightarrow f(x) \in \Pi$, and $f(x) \in \Pi \Rightarrow x \in \Pi'$.

Readers are encouraged to identify these four steps in each proof that follows.

Let us show that the problem of vertex cover is harder than the problem of independent set in an undirected graph. Let $G = (V, E)$ be an undirected graph with the vertex set V and the edge set E .

Definition 13.4 (Independent Set) *The Independent Set problem asks: Given a graph G and an integer k , does there exist a subset $S \subseteq V$ of at least k vertices such that no two vertices in S are adjacent (i.e., $\forall u, v \in S, \{u, v\} \notin E$)? Such a set S is called an independent set.*

Definition 13.5 (Vertex Cover) *The Vertex Cover problem asks: Given a graph G and an integer k , does there exist a subset $C \subseteq V$ of at most k vertices such that every edge in E is incident to at least one vertex in C (i.e., $\forall \{u, v\} \in E, u \in C$ or $v \in C$)? Such a set C is called a vertex cover.*

The following theorem captures the complementary relationship between Independent Set and Vertex Cover.

Theorem 13.6 (Independent Set – Vertex Cover duality) *In any graph $G = (V, E)$ with $|V| = n$, a set $S \subseteq V$ is an independent set iff $V \setminus S$ is a vertex cover. Consequently, G has an independent set of size $\geq k$ iff it has a vertex cover of size $\leq n - k$.*

Proof:

- (\Rightarrow): Suppose $S \subseteq V$ is an independent set with $|S| \geq k$. Define $C = V \setminus S$. Since S is independent, no edge $\{u, v\} \in E$ has both $u, v \in S$. Thus, for every edge $\{u, v\} \in E$, at least one of u or v is in $C = V \setminus S$. Hence, C is a vertex cover. Moreover, $|C| = |V| - |S| \leq |V| - k = n - k$.
- (\Leftarrow): Suppose $C \subseteq V$ is a vertex cover with $|C| \leq n - k$. Define $S = V \setminus C$. For any edge $\{u, v\} \in E$, since C is a vertex cover, at least one of u or v is in C , so at most one is in S . But if both $u, v \in S$, then neither is in C , contradicting that C covers $\{u, v\}$. Thus, no edge connects vertices in S , so S is an independent set. Moreover, $|S| = |V| - |C| \geq |V| - (n - k) = k$.

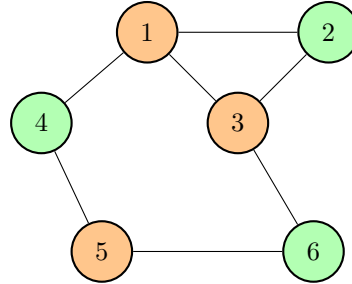


Figure 13.1: A graph $G = (V, E)$ with $V = \{1, \dots, 6\}$ and edges $\{\{1, 2\}, \{1, 3\}, \{2, 3\}, \{1, 4\}, \{4, 5\}, \{5, 6\}, \{3, 6\}\}$. The set $S = \{2, 4, 6\}$ is a maximum independent set (shaded green); its complement $C = \{1, 3, 5\}$ is the corresponding minimum vertex cover (shaded orange).

Example 13.7 In Figure 13.1, G has vertex set $V = \{1, \dots, 6\}$. The set $S = \{2, 4, 6\}$ is an independent set of size 3: no two of its vertices are joined by an edge. It is in fact a maximum independent set, since the triangle $\{1, 2, 3\}$ forces any larger independent set to pick at least two pairwise-nonadjacent vertices from $\{4, 5, 6\}$, but then one of $\{1, 2, 3\}$ becomes adjacent to a chosen pendant. Its complement $C = V \setminus S = \{1, 3, 5\}$ is a vertex cover: every edge has at least one endpoint in $\{1, 3, 5\}$. Consistent with Theorem 13.6, $|S| = 3 = n - |C|$.

Theorem 13.6 immediately yields a polynomial-time reduction $\text{Independent Set} \leq_P \text{Vertex Cover}$: given an instance (G, k) of Independent Set, output the Vertex Cover instance $(G, |V| - k)$. The reduction runs in $O(|V|)$ time, and by the theorem, (G, k) is a *yes* instance of Independent Set iff $(G, |V| - k)$ is a *yes* instance of Vertex Cover.

Conversely, we argue that the Independent Set problem is at least as hard as the Vertex Cover problem by providing a polynomial-time reduction from Vertex Cover to Independent Set.

Given an instance $(G = (V, E), k)$ of Vertex Cover, construct an instance of Independent Set as follows:

- Use the same graph $G = (V, E)$.
- Set the parameter $k' = |V| - k$.

The reduction is polynomial-time, identical to the previous reduction.

Since we have reduced Vertex Cover to Independent Set in polynomial time, Independent Set is at least as hard as Vertex Cover. Formally, $\text{Vertex Cover} \leq_P \text{Independent Set}$.

The bidirectional reductions show that Independent Set and Vertex Cover are computationally equivalent in terms of polynomial-time solvability: $\text{Independent Set} \leq_P \text{Vertex Cover}$ and $\text{Vertex Cover} \leq_P \text{Independent Set}$. The reductions exploit the complementary relationship: S is an independent set if and only if $V \setminus S$ is a vertex cover.

We now show that the problem of *Set Cover* is harder than the vertex cover.

Definition 13.8 (Set Cover) *The Set Cover problem is defined as follows: Given a universe U of n elements $\{e_1, e_2, \dots, e_n\}$, a collection $\mathcal{S} = \{S_1, S_2, \dots, S_m\}$ of subsets of U such that $\bigcup_{i=1}^m S_i = U$, and an integer k , does there exist a subcollection $\mathcal{C} \subseteq \mathcal{S}$ of at most k subsets such that $\bigcup_{S_i \in \mathcal{C}} S_i = U$? In other words, we seek a cover of U using at most k subsets from \mathcal{S} .*

It is easy to see that the vertex cover is a special case of set cover. Formally, given a Vertex Cover instance $(G = (V, E), k)$, build a Set Cover instance (U, \mathcal{S}, k') by taking $U = E$ (one universe element per edge), $\mathcal{S} = \{S_v : v \in V\}$ where $S_v = \{e \in E : v \in e\}$ is the set of edges incident to v , and $k' = k$. The construction runs in $O(|V| + |E|)$ time. Choosing $\mathcal{C} \subseteq \mathcal{S}$ of size at most k that covers U corresponds exactly to choosing $\leq k$ vertices that cover every edge. Hence Vertex Cover \leq_P Set Cover.

So far, our problems were graph-theoretic. Let us now investigate problems that relate to satisfiability of boolean expression. A Boolean expression is said to be in Conjunctive Normal Form (CNF) if it is a conjunction of *clauses* where each clause is a disjunction of *literals*. A literal is any boolean variable or its negation. We call the problem of checking satisfiability of a boolean expression in CNF the problem SAT. We define 3-SAT as the special case of SAT where each clause has exactly three literals.

Having seen several reductions between problems, we now formalize what it means for a problem to be “as hard as any problem in NP”.

Definition 13.9 (Class NP) *A decision problem $L \subseteq \{0, 1\}^*$ is in the class NP if there exists a polynomial-time deterministic verifier V and a polynomial p such that for every $x \in \{0, 1\}^*$:*

$$x \in L \iff \exists c \in \{0, 1\}^{\leq p(|x|)} \text{ with } V(x, c) = 1.$$

The string c is called a certificate (or witness) for x .

Informally, the class NP includes problems where a *yes* certificate is verifiable in polynomial time. A *yes* certificate is an input, of size polynomial in $|x|$, that certifies the given instance is a *yes* instance.

Example 13.10 SAT is in NP: given a CNF formula ϕ over n variables, a satisfying assignment $\tau \in \{0, 1\}^n$ serves as a certificate of size n . Substituting τ into ϕ and evaluating each clause takes polynomial time, so the verifier accepts iff $\phi(\tau) = 1$.

As another example, consider the Subset Sum problem: given a set S of integers and a target t , is there a subset of S summing to t ? A *yes* certificate is simply a candidate subset $S' \subseteq S$; verifying that $\sum_{a \in S'} a = t$ takes linear time.

Note that we require certificates only for *yes* instances. We claim

$$P \subseteq \text{NP}.$$

Given any problem $\pi \in P$, there is a polynomial-time algorithm for π , so certification is trivial: the verifier simply ignores the certificate and runs the algorithm. Whether the containment is strict — whether there is a problem in NP but not in P — is the most famous open problem in theoretical computer science.

Definition 13.11 (NP-hard) A decision problem L is NP-hard if for every $L' \in \text{NP}$, $L' \leq_p L$.

Definition 13.12 (NP-complete) A decision problem L is NP-complete if $L \in \text{NP}$ and L is NP-hard.

The first problem to be shown NP-complete was SAT, by Cook in 1971 and independently by Levin in 1973.

Theorem 13.13 (Cook-Levin Theorem) SAT is NP-complete.

Proof: *Sketch.* SAT is in NP: a satisfying assignment is the certificate. For NP-hardness, fix any $L \in \text{NP}$ decided by a nondeterministic polynomial-time Turing machine M running for at most $p(n)$ steps on inputs of length n . Introduce Boolean variables that describe an accepting computation *tableau* of M on input x : one variable per (tape cell, time step, symbol), per (time step, state), and per (time step, head position). Clauses encode that (i) the initial row records x and the start state, (ii) every later row follows from the previous one by one of M 's transitions, and (iii) some row enters an accepting state. The resulting CNF has $O(p(n)^2)$ variables and clauses, is constructible in polynomial time, and is satisfiable iff M accepts x . Hence $L \leq_p \text{SAT}$. A fully detailed tableau encoding is given in Sipser [Sip13] and Arora–Barak [AB09].

■

Corollary 13.14 If any NP-complete problem is in P , then $P = \text{NP}$.

Proof: Let L be an NP-complete problem with $L \in P$. For any $L' \in \text{NP}$, we have $L' \leq_p L$ by definition of NP-hardness; composing the polynomial-time reduction with the polynomial-time algorithm for L yields a polynomial-time algorithm for L' , so $L' \in P$. Hence $\text{NP} \subseteq P$. Combined with the trivial inclusion $P \subseteq \text{NP}$, we get $P = \text{NP}$.

■

Figure 13.2 illustrates the relationship between P , NP-complete, and NP. P is a subset of NP, as polynomial-time solvable problems are verifiable in polynomial time. NP-complete problems are the hardest in NP, with every NP problem reducing to them in polynomial time. If $P \neq \text{NP}$, NP-complete problems are disjoint from P .

We now investigate the complexity of 3-SAT compared to SAT.

It is clear that SAT is at least as hard as 3-SAT because 3-SAT is a special case of SAT. Somewhat surprisingly, 3-SAT is as hard as SAT.

Theorem 13.15 3-SAT is NP-complete.

Proof: 3-SAT is in NP: a satisfying truth assignment is a certificate, and evaluating a 3-CNF formula under a given assignment takes polynomial time.

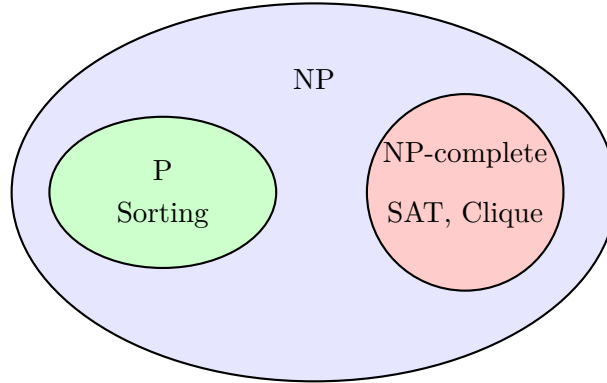


Figure 13.2: Relationship between P, NP-complete, and NP, assuming $P \neq NP$.

To establish NP-hardness we give a polynomial-time reduction $SAT \leq_P 3\text{-SAT}$; NP-hardness of 3-SAT then follows from NP-hardness of SAT (Cook–Levin, Theorem 13.13).

Given a CNF formula $\phi = C_1 \wedge \cdots \wedge C_m$ with $C_i = l_{i1} \vee \cdots \vee l_{ik_i}$, we construct a 3-CNF formula ψ clause-by-clause.

Construction. For each clause C_i :

- **Case 1:** $k_i = 1$ (clause (l_1)): introduce fresh variables y_i, z_i and replace C_i by

$$(l_1 \vee y_i \vee z_i) \wedge (l_1 \vee y_i \vee \neg z_i) \wedge (l_1 \vee \neg y_i \vee z_i) \wedge (l_1 \vee \neg y_i \vee \neg z_i).$$

- **Case 2:** $k_i = 2$ (clause $(l_1 \vee l_2)$): introduce y_i and replace by

$$(l_1 \vee l_2 \vee y_i) \wedge (l_1 \vee l_2 \vee \neg y_i).$$

- **Case 3:** $k_i = 3$: keep $(l_1 \vee l_2 \vee l_3)$ unchanged.

- **Case 4:** $k_i > 3$ (clause $(l_1 \vee \cdots \vee l_k)$): introduce $y_{i,1}, \dots, y_{i,k-3}$ and replace by

$$(l_1 \vee l_2 \vee y_{i,1}) \wedge (\neg y_{i,1} \vee l_3 \vee y_{i,2}) \wedge \cdots \wedge (\neg y_{i,k-3} \vee l_{k-1} \vee l_k).$$

Each clause of length k_i yields $O(k_i)$ clauses of length exactly 3, and no more than $O(k_i)$ fresh variables, so the overall construction is polynomial in $|\phi|$.

Correctness. Each replacement preserves satisfiability clause-by-clause. In Case 1, if l_1 is true then every one of the four clauses is already satisfied by l_1 . Conversely, if l_1 is false, the four clauses simplify to

$$(y_i \vee z_i) \wedge (y_i \vee \neg z_i) \wedge (\neg y_i \vee z_i) \wedge (\neg y_i \vee \neg z_i),$$

which is unsatisfiable: a quick truth-table check shows that every assignment of (y_i, z_i) falsifies at least one of the four clauses. Hence the conjunction is logically equivalent to l_1 . Case 2 is similar for $(l_1 \vee l_2)$. Case 4 is verified by induction: if some l_j is true in C_i , set $y_{i,1} = \cdots = y_{i,j-2} = T$ and $y_{i,j-1} = \cdots = y_{i,k-3} = F$ to satisfy every new clause; conversely, if every new clause is satisfied but all l_j were false, the chain of implications $y_{i,1} \Rightarrow y_{i,2} \Rightarrow \cdots \Rightarrow y_{i,k-3}$ together with the last clause $\neg y_{i,k-3} \vee l_{k-1} \vee l_k$ forces a contradiction. Hence ϕ is satisfiable iff ψ is, and since ψ is a 3-CNF, $SAT \leq_P 3\text{-SAT}$. ■

Example 13.16 [SAT to 3-SAT] Consider the SAT instance

$$\phi = (x_1) \wedge (x_2 \vee x_3) \wedge (x_1 \vee \neg x_2 \vee x_3 \vee x_4),$$

which has one 1-clause, one 2-clause, and one 4-clause. Applying the reduction:

- The 1-clause (x_1) uses dummy variables y_1, z_1 and becomes

$$(x_1 \vee y_1 \vee z_1) \wedge (x_1 \vee y_1 \vee \neg z_1) \wedge (x_1 \vee \neg y_1 \vee z_1) \wedge (x_1 \vee \neg y_1 \vee \neg z_1).$$

- The 2-clause $(x_2 \vee x_3)$ uses dummy y_2 and becomes

$$(x_2 \vee x_3 \vee y_2) \wedge (x_2 \vee x_3 \vee \neg y_2).$$

- The 4-clause $(x_1 \vee \neg x_2 \vee x_3 \vee x_4)$ uses dummy y_3 and becomes

$$(x_1 \vee \neg x_2 \vee y_3) \wedge (\neg y_3 \vee x_3 \vee x_4).$$

The resulting 3-CNF formula ψ has $4 + 2 + 2 = 8$ clauses, each of size exactly 3. The assignment $x_1 = \text{true}, x_2 = \text{true}, x_3 = \text{true}, x_4 = \text{false}$ satisfies ϕ ; extending it with $y_3 = \text{false}$ (any choice of y_1, z_1, y_2) satisfies ψ .

Theorem 13.17 *Independent Set is NP-complete.*

Proof: We follow the four-step template announced earlier in this section.

Step 1 (Membership in NP). A candidate subset $S \subseteq V$ is a certificate, and checking that no edge of G has both endpoints in S takes $O(|E|)$ time.

Step 2 (Source). We reduce from 3-SAT, known to be NP-complete by Theorem 13.15.

Step 3 (Reduction). Given a 3-SAT formula ϕ with variables x_1, \dots, x_n and clauses C_1, \dots, C_m , where each clause $C_j = (l_{j1} \vee l_{j2} \vee l_{j3})$, we construct an instance $(G = (V, E), k)$:

- **Vertices.** For each clause C_j create three vertices v_{j1}, v_{j2}, v_{j3} , one per literal. Thus $V = \{v_{jk} \mid 1 \leq j \leq m, 1 \leq k \leq 3\}$, and $|V| = 3m$.
- **Intra-clause edges.** For each clause C_j , form a triangle on $\{v_{j1}, v_{j2}, v_{j3}\}$. This ensures at most one vertex per clause is picked by any independent set.
- **Inter-clause edges.** For any two vertices v_{jk} and $v_{j'k'}$ with $j \neq j'$, add the edge $\{v_{jk}, v_{j'k'}\}$ iff the literals l_{jk} and $l_{j'k'}$ are *inconsistent* (one is x_i and the other $\neg x_i$).
- **Parameter.** $k = m$.

The construction runs in $O(m^2)$ time.

Step 4 (Correctness). We claim that ϕ is satisfiable iff G has an independent set of size at least m .

(\Rightarrow) Suppose ϕ has a satisfying assignment τ . In each clause C_j choose one literal l_{jk} that is true under τ , and let $S = \{v_{jk} : \text{the picked vertex of } C_j\}$. Then $|S| = m$. No intra-clause edge lies inside S (one vertex per clause), and no inter-clause edge lies inside S either: if $v_{jk}, v_{j'k'} \in S$ were joined by an inconsistency edge then l_{jk} and $l_{j'k'}$ could not both be true under τ . Hence S is independent of size m .

(\Leftarrow) Suppose G has an independent set S with $|S| \geq m$. Each clause gadget is a triangle, so at most one vertex per clause is in S ; with m clauses and $|S| \geq m$, S has exactly one vertex per clause. Set $x_i = \text{true}$ if some $v_{jk} \in S$ corresponds to $l_{jk} = x_i$, and $x_i = \text{false}$ if some $v_{jk} \in S$ corresponds to $l_{jk} = \neg x_i$. This assignment is consistent: a conflict would require both x_i and $\neg x_i$ to appear in S , but those two vertices are joined by an inconsistency edge, contradicting independence. The assignment makes some literal true in every clause, so ϕ is satisfied.

Hence $\phi \in 3\text{-SAT} \Leftrightarrow (G, m) \in \text{Independent Set}$, completing the reduction. Subsequent NP-completeness proofs in this chapter follow the same four-step template; we abbreviate the step labels for brevity. ■

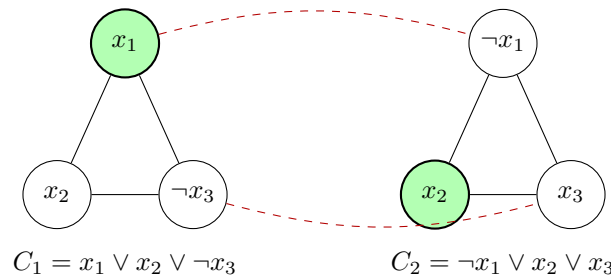


Figure 13.3: Reduction of the 3-SAT instance $\phi = (x_1 \vee x_2 \vee \neg x_3) \wedge (\neg x_1 \vee x_2 \vee x_3)$ to Independent Set. Each clause becomes a triangle (clause gadget). Dashed red edges connect inconsistent literals across clauses: $x_1 \leftrightarrow \neg x_1$ and $\neg x_3 \leftrightarrow x_3$. The shaded vertices $\{x_1 \text{ from } C_1, x_2 \text{ from } C_2\}$ form an independent set of size $2 = m$, corresponding to the satisfying assignment $x_1 = \text{true}, x_2 = \text{true}, x_3 = \text{anything}$.

Example 13.18 [3-SAT to Independent Set] Consider the 3-SAT instance $\phi = (x_1 \vee x_2 \vee \neg x_3) \wedge (\neg x_1 \vee x_2 \vee x_3)$ with $m = 2$ clauses. The reduction produces the graph in Figure 13.3 with $3m = 6$ vertices and $k = m = 2$. Each clause triangle enforces that at most one literal per clause is picked; the dashed inconsistency edges forbid picking x_1 from C_1 together with $\neg x_1$ from C_2 , or $\neg x_3$ from C_1 together with x_3 from C_2 . Picking x_1 from C_1 and x_2 from C_2 (shaded) yields an independent set of size 2, and reading off the corresponding literals gives the satisfying assignment $x_1 = \text{true}, x_2 = \text{true}$ (x_3 is unconstrained).

With Independent Set now established as NP-complete, the duality between Independent Set and Vertex Cover (Theorem 13.6) immediately gives the NP-completeness of Vertex Cover as well.

Corollary 13.19 *Vertex Cover is NP-complete.*

Proof: Vertex Cover is in NP: a candidate cover $C \subseteq V$ with $|C| \leq k$ is a certificate, verified in $O(|E|)$ time by checking that every edge is incident to some vertex in C . For NP-hardness, reduce Independent

Set to Vertex Cover via Theorem 13.6: given an Independent Set instance (G, k) with $|V| = n$, output the Vertex Cover instance $(G, n - k)$. By the theorem, G has an independent set of size $\geq k$ iff G has a vertex cover of size $\leq n - k$. Combined with Theorem 13.17 (Independent Set is NP-complete), Vertex Cover is NP-hard. ■

13.4 More NP-Complete Problems

In this section we continue building the catalogue of NP-complete problems by proving NP-completeness of Clique and Subset Sum. For each problem we start from a problem already known to be NP-complete and give a polynomial-time reduction to the new problem.

Definition 13.20 (Clique) *The Clique problem asks: Given a graph $G = (V, E)$ and an integer k , does there exist a subset $Q \subseteq V$ of at least k vertices such that every pair of distinct vertices in Q is adjacent (i.e., $\forall u, v \in Q$ with $u \neq v$, $\{u, v\} \in E$)? Such a set Q is called a clique.*

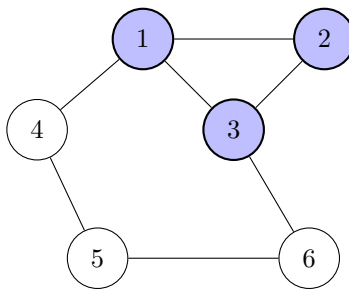


Figure 13.4: The same graph as in Figure 13.1, with the clique $Q = \{1, 2, 3\}$ shaded blue.

Example 13.21 In Figure 13.4, the set $Q = \{1, 2, 3\}$ is a clique of size 3: all three pairs $\{1, 2\}$, $\{1, 3\}$, $\{2, 3\}$ are edges. No clique of size 4 exists: any vertex outside the triangle is 4, 5, or 6, each of which has at most two neighbours, so no four vertices can be pairwise adjacent. Note that this clique is distinct from the vertex cover $C = \{1, 3, 5\}$ identified in Example 13.7; the two sets share only the vertices 1 and 3.

We now prove that Clique is NP-complete, using a polynomial-time reduction $3\text{-SAT} \leq_P \text{Clique}$.

Theorem 13.22 *Clique is NP-complete.*

Proof: Clique is in NP: a candidate clique $Q \subseteq V$ with $|Q| \geq k$ is a certificate, and verifying that every pair of vertices in Q is joined by an edge takes $O(|Q|^2)$ time.

To establish NP-hardness we reduce 3-SAT to Clique. Given a 3-SAT formula ϕ with n variables x_1, \dots, x_n and m clauses C_1, \dots, C_m — where each clause $C_j = (l_{j1} \vee l_{j2} \vee l_{j3})$ and l_{jk} is a literal — we construct an instance $(G = (V, E), k)$ of Clique:

1. **Vertices:** For each literal l_{jk} in clause C_j , create a vertex v_{jk} . Thus $V = \{v_{jk} \mid j = 1, \dots, m, k = 1, 2, 3\}$, and $|V| = 3m$.
2. **Edges:** Add an edge $\{v_{jk}, v_{j'k'}\}$ (for $j \neq j'$) iff the literals l_{jk} and $l_{j'k'}$ are *consistent*, i.e., they are not of the form x_i and $\neg x_i$. No edges are placed between vertices of the same clause.
3. **Parameter:** $k = m$.

The construction runs in $O(m^2)$ time. We claim that ϕ is satisfiable iff G has a clique of size at least m .

(\Rightarrow) Suppose ϕ has a satisfying assignment τ . In each clause C_j pick one literal l_{jk} that is true under τ , and let $Q = \{v_{jk} : \text{the picked vertex of } C_j\}$. Then $|Q| = m$, vertices of Q come from distinct clauses, and every pair of picked literals is consistent under τ (both evaluate to true), so every pair of vertices in Q is connected by an edge. Hence Q is a clique of size m .

(\Leftarrow) Suppose G has a clique Q with $|Q| \geq m$. Since vertices of the same clause are never adjacent, Q contains at most one vertex per clause; with $|Q| \geq m$ we have $|Q| = m$ and exactly one vertex per clause. Read off the literals: if $v_{jk} \in Q$ corresponds to $l_{jk} = x_i$, set $x_i = \text{true}$; if $l_{jk} = \neg x_i$, set $x_i = \text{false}$. No conflict arises: a conflict would require two vertices $v_{jk}, v_{j'k'} \in Q$ labelled by x_i and $\neg x_i$, but then they would be non-adjacent in G , contradicting that Q is a clique. The resulting assignment makes some literal true in each clause, so ϕ is satisfied.

Hence $\phi \in 3\text{-SAT} \Leftrightarrow (G, m) \in \text{Clique}$, and Clique is NP-hard. Combined with $\text{Clique} \in \text{NP}$, the claim follows. ■

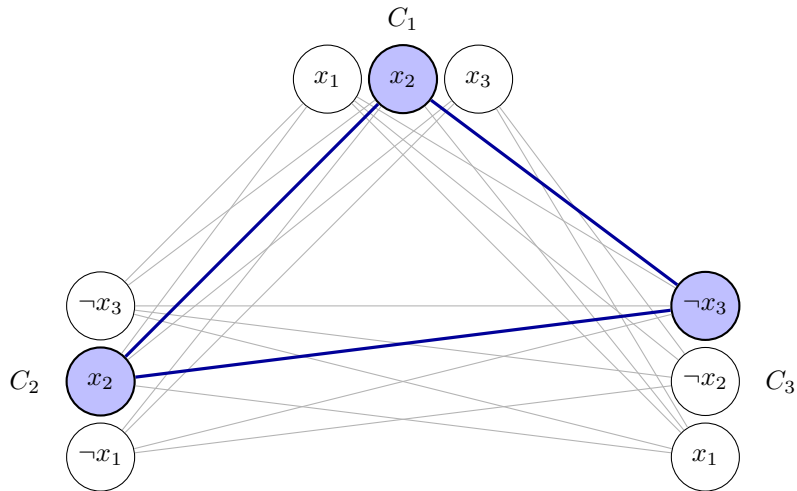


Figure 13.5: Reduction of the 3-SAT instance $\phi = (x_1 \vee x_2 \vee x_3) \wedge (\neg x_1 \vee x_2 \vee \neg x_3) \wedge (x_1 \vee \neg x_2 \vee \neg x_3)$ to Clique. No edges exist within the same clause gadget; an edge joins two vertices from different clauses iff their literals are non-contradictory (grey). The shaded vertices $\{x_2 \text{ in } C_1, x_2 \text{ in } C_2, \neg x_3 \text{ in } C_3\}$ together with the bold blue edges form a clique of size $m = 3$, witnessing that ϕ is satisfiable.

Example 13.23 [3-SAT to Clique] Consider the 3-SAT instance

$$\phi = (x_1 \vee x_2 \vee x_3) \wedge (\neg x_1 \vee x_2 \vee \neg x_3) \wedge (x_1 \vee \neg x_2 \vee \neg x_3),$$

with $m = 3$ clauses. The reduction produces the graph in Figure 13.5 on $3m = 9$ vertices, with clique threshold $k = m = 3$. Note the complement structure compared with the Independent Set reduction (Figure 13.3): here a clause gadget has *no* internal edges, while inter-clause edges join non-contradictory literals. The shaded vertices $\{x_2 \text{ in } C_1, x_2 \text{ in } C_2, \neg x_3 \text{ in } C_3\}$ form a clique of size $k = 3$ (bold blue edges). Reading off the corresponding literals gives the satisfying assignment $x_2 = \text{true}$, $x_3 = \text{false}$ (with x_1 unconstrained).

Definition 13.24 (Subset Sum) *The Subset Sum problem asks whether, given a set of positive integers $S = \{a_1, \dots, a_n\}$ and a target t , there exists a subset $S' \subseteq S$ summing to t .*

Definition 13.25 (3D-Matching) *The 3D-Matching (3DM) problem is defined as follows. Given three disjoint sets $X = \{x_1, \dots, x_n\}$, $Y = \{y_1, \dots, y_n\}$, $Z = \{z_1, \dots, z_n\}$ and a set of triples $T \subseteq X \times Y \times Z$, does there exist a subset $M \subseteq T$ of size n such that every element of $X \cup Y \cup Z$ appears in exactly one triple of M ?*

3D-Matching is one of Karp’s classical NP-complete problems [Kar72]; the reduction $3\text{-SAT} \leq_P 3\text{DM}$ uses “variable gadgets” (one for each x_i , with $2 \cdot \text{occ}(x_i)$ matched triples acting as a truth flag) and “clause gadgets” (one triple per literal, sharing core elements that force exactly one literal per clause to be selected), plus padding triples to balance the universe sizes. We take 3DM’s NP-completeness for granted in what follows and use it to prove NP-completeness of Subset Sum. (An alternative direct route, $\text{Vertex Cover} \leq_P \text{Subset Sum}$, is left as an exercise.)

Theorem 13.26 *Subset Sum is NP-complete.*

Proof: Subset Sum is in NP: a subset $S' \subseteq S$ is a certificate, and checking $\sum_{a \in S'} a = t$ takes time polynomial in the size of the input.

To establish NP-hardness we give a polynomial-time reduction $3\text{D-Matching} \leq_P \text{Subset Sum}$. Given a 3DM instance with sets $X = \{x_1, \dots, x_n\}$, $Y = \{y_1, \dots, y_n\}$, $Z = \{z_1, \dots, z_n\}$ and triples $T = \{t_1, \dots, t_m\} \subseteq X \times Y \times Z$, we construct a Subset Sum instance (S, t) with $3n$ digits in base $b = m + 1$ to ensure carry-free addition:

- **Elements:** the $3n$ elements $x_1, \dots, x_n, y_1, \dots, y_n, z_1, \dots, z_n$ are assigned to digit positions $3n, 3n - 1, \dots, 1$ respectively.
- **Base:** use $b = m + 1$. Since each element appears in at most m triples, the sum in each digit position is at most $m < b$, so additions produce no carries.
- **Numbers:** for each triple $t_j = (x_{j_1}, y_{j_2}, z_{j_3})$, create

$$a_j = b^{3n-j_1} + b^{2n-j_2} + b^{n-j_3},$$

so a_j is 1 in the three digit positions of its three elements and 0 elsewhere.

- **Target:**

$$t = \sum_{k=1}^n (b^{3n-k} + b^{2n-k} + b^{n-k}),$$

which has a 1 in each of the $3n$ digit positions.

- **Set S :** $S = \{a_1, \dots, a_m\}$.

The construction is clearly polynomial time: each a_j and t have $O(n \log m)$ bits.

(\Rightarrow) If T contains a perfect matching $M \subseteq T$ of n triples covering each element exactly once, then $S' = \{a_j : t_j \in M\}$ sums to t : each digit position receives exactly one 1.

(\Leftarrow) Conversely, if $S' \subseteq S$ sums to t , then because addition is carry-free in base b and t has a 1 in each of the $3n$ positions, each digit position of S' must sum to exactly 1. Each a_j contributes three 1s, so $|S'| = n$, and each element is covered exactly once: the corresponding triples form a perfect matching.

Hence $(X, Y, Z, T) \in 3DM \Leftrightarrow (S, t) \in \text{Subset Sum}$, so Subset Sum is NP-hard. Combined with Subset Sum \in NP, the claim follows. ■

Example 13.27 [3D-Matching to Subset Sum] Consider the 3DM instance with $n = 2$, so $X = \{x_1, x_2\}$, $Y = \{y_1, y_2\}$, $Z = \{z_1, z_2\}$, and $m = 3$ triples:

$$t_1 = (x_1, y_1, z_1), \quad t_2 = (x_2, y_2, z_2), \quad t_3 = (x_1, y_2, z_1).$$

The unique perfect matching is $M = \{t_1, t_2\}$: t_1 covers x_1, y_1, z_1 and t_2 covers x_2, y_2, z_2 , so every element appears in exactly one chosen triple. (The pair $\{t_1, t_3\}$ fails because x_1 is covered twice; similarly for $\{t_2, t_3\}$ with y_2 .)

The reduction uses base $b = m + 1 = 4$ and $3n = 6$ digit positions, one per element. Writing numbers in base 4 with six digit positions (most significant digit leftmost corresponds to x_1 , least significant to z_2):

triple	a_j	base-4 digits ($x_1 x_2 y_1 y_2 z_1 z_2$)
$t_1 = (x_1, y_1, z_1)$	$4^5 + 4^3 + 4^1 = 1092$	1 0 1 0 1 0
$t_2 = (x_2, y_2, z_2)$	$4^4 + 4^2 + 4^0 = 273$	0 1 0 1 0 1
$t_3 = (x_1, y_2, z_1)$	$4^5 + 4^2 + 4^1 = 1044$	1 0 0 1 1 0

The target $t = 4^5 + 4^4 + 4^3 + 4^2 + 4^1 + 4^0 = 1365$ has the base-4 digit string 111111, one 1 in every column. The Subset Sum instance is $(S, t) = (\{1092, 273, 1044\}, 1365)$.

Observe that $a_1 + a_2 = 1092 + 273 = 1365 = t$: the corresponding base-4 addition $101010 + 010101 = 111111$ is carry-free and each column sums to exactly 1, matching the fact that $\{t_1, t_2\}$ covers every element exactly once. No other subset of S sums to t : for example, $a_1 + a_3$ produces a 2 in the x_1 column (since t_1 and t_3 both use x_1), so the resulting base-4 string is not 111111.

Problem	Definition	Reduction From
3-Colorability	Graph G 3-colorable?	SAT
3D-Matching	Perfect matching in $X \times Y \times Z$?	3-SAT
Knapsack	Items fit W , value $\geq V$?	Subset Sum
Hamiltonian Path	Path visits all vertices once?	3-SAT
Hamiltonian Cycle	Cycle visits all vertices once?	Hamiltonian Path
Traveling Salesman	Tour with weight $\leq B$?	Hamiltonian Cycle

Figure 13.6: NP-Complete Problems and Reductions

Two further reductions in the chain Cook–Levin \rightarrow SAT \rightarrow 3-SAT \rightarrow Hamiltonian Path \rightarrow Hamiltonian Cycle \rightarrow TSP advertised by Figure 13.8 are short enough to give in full here. We take Hamiltonian Path’s NP-completeness as established (the 3-SAT \rightarrow HP reduction with variable-and-clause gadgets is given in, e.g., Sipser [Sip13]).

Theorem 13.28 *Hamiltonian Cycle is NP-complete.*

Proof: HC is in NP: a permutation of the vertices is a certificate, verified in $O(n)$ time. For NP-hardness we reduce HP \leq_P HC. Given an HP instance $G = (V, E)$, build $G' = (V \cup \{v^*\}, E \cup \{\{v^*, u\} : u \in V\})$: a fresh vertex v^* adjacent to every original vertex. The construction is $O(n)$. If G has a Hamiltonian path u_1, u_2, \dots, u_n , then $u_1, u_2, \dots, u_n, v^*, u_1$ is a Hamiltonian cycle in G' . Conversely, removing v^* from any Hamiltonian cycle of G' leaves a Hamiltonian path of G between the two original neighbours of v^* in the cycle. ■

Theorem 13.29 *The decision version of the Traveling Salesman Problem is NP-complete.*

Proof: TSP is in NP: a permutation of the vertices certifies a tour, verified in $O(n)$ time. For NP-hardness we reduce HC \leq_P TSP. Given an HC instance $G = (V, E)$ with $n = |V|$, build the complete graph $G' = (V, V \times V)$ with edge weights $w(u, v) = 1$ if $\{u, v\} \in E$ and $w(u, v) = 2$ otherwise; set the threshold $B = n$. The construction is $O(n^2)$. G' admits a TSP tour of weight $\leq n$ iff every edge of the tour has weight 1, which is equivalent to the tour using only edges of G , i.e., a Hamiltonian cycle of G . ■

We now list further problems which have been shown to be NP-complete. Table 13.6 summarizes six NP-complete decision problems: 3-Colorability, 3D-Matching, Knapsack, Hamiltonian Path, Hamiltonian Cycle, and Traveling Salesman. For each problem, we provide its definition and identify a known NP-complete problem that can be polynomially reduced to it. The Hamiltonian Cycle and TSP reductions are made explicit in Theorems 13.28 and 13.29 above; the others are sketched in the references.

1. 3-Colorability:

- *Definition:* Given a graph $G = (V, E)$, can the vertices be colored with three colors such that no adjacent vertices share the same color?
- *Reduction From:* SAT. Construct a graph with gadgets for variables (true/false colors) and clauses (ensuring at least one true literal), where a 3-coloring encodes a satisfying assignment.

2. 3D-Matching:

- *Definition:* Given sets X, Y, Z , each of size n , and triples $T \subseteq X \times Y \times Z$, is there a subset of n triples covering each element exactly once?
- *Reduction From:* 3-SAT. Create variable gadgets (true/false triples) and clause gadgets (triples for true literals), where a matching corresponds to a satisfying assignment.

3. Knapsack:

- *Definition:* Given items with weights w_i , values v_i , capacity W , and target V , is there a subset with weight $\leq W$, value $\geq V$?
- *Reduction From:* Subset Sum. Set $v_i = w_i$, $W = t$, $V = t$, making Knapsack equivalent to Subset Sum.

4. Hamiltonian Path:

- *Definition:* Given a graph $G = (V, E)$, is there a path visiting each vertex exactly once?
- *Reduction From:* 3-SAT. Build a graph with variable paths (true/false) and clause nodes, where a Hamiltonian path encodes a satisfying assignment.

5. Hamiltonian Cycle:

- *Definition:* Given a graph $G = (V, E)$, is there a cycle visiting each vertex exactly once?
- *Reduction From:* Hamiltonian Path. Add a vertex connected to all others, where a Hamiltonian cycle implies a Hamiltonian path in the original graph.

6. Traveling Salesman:

- *Definition:* Given a complete graph with edge weights and bound B , is there a tour of weight $\leq B$?
- *Reduction From:* Hamiltonian Cycle. Set weights: 1 for original edges, 2 otherwise, $B = n$. A tour of weight n is a Hamiltonian cycle.

13.5 co-NP Class of Problems

In computational complexity theory, the class co-NP plays a crucial role alongside NP in understanding the structure of decision problems.

Definition 13.30 (co-NP) *The complexity class co-NP consists of all decision problems whose complements are in NP. That is, $L \in \text{co-NP}$ iff $\bar{L} \in \text{NP}$.*

Equivalently, a problem L is in co-NP if there exists a polynomial-time verifiable certificate for *no* instances (i.e., instances $x \notin L$). This means there is a polynomial-time Turing machine M and a polynomial p such that for every input $x \notin L$, there exists a certificate c of length at most $p(|x|)$ where $M(x, c) = 1$, and for every $x \in L$, no such certificate exists.

In contrast, a problem L is in NP if there exists a polynomial-time verifiable certificate for *yes* instances ($x \in L$). Thus, co-NP is the class of problems where the evidence for rejection is efficiently verifiable, while NP focuses on evidence for acceptance.

The relationship between co-NP and NP is symmetric due to their complementary definitions:

- If $L \in \text{NP}$, then $\bar{L} \in \text{co-NP}$.
- If $L \in \text{co-NP}$, then $\bar{L} \in \text{NP}$.

This symmetry implies that NP and co-NP are *dual* classes. Notably:

- $\text{P} \subseteq \text{NP} \cap \text{co-NP}$, since if a problem is in P, both its yes and no instances can be decided in polynomial time, providing trivial certificates.
- $\text{NP} \cap \text{co-NP}$ contains problems like Integer Factorization (deciding if a number n has a factor d with $1 < d < n$), where both existence (NP) and non-existence (co-NP) of factors are verifiable with certificates (a factor or a primality certificate).

A central question is whether $\text{NP} = \text{co-NP}$. If $\text{NP} = \text{co-NP}$, then for every NP problem, its complement would also have efficiently verifiable yes certificates, implying a symmetry in verification. However, it is widely believed that $\text{NP} \neq \text{co-NP}$, as their equality would collapse the polynomial hierarchy and have profound implications for problems like SAT and UNSAT.

The following figure illustrates the relationship between NP, co-NP, and P. P is explicitly drawn as a distinct region within the intersection of NP and co-NP, representing problems solvable in polynomial time. NP and co-NP are dual classes, where a problem is in co-NP if its complement is in NP. Their intersection, $\text{NP} \cap \text{co-NP}$, contains P and problems like Integer Factorization, not known to be in P.

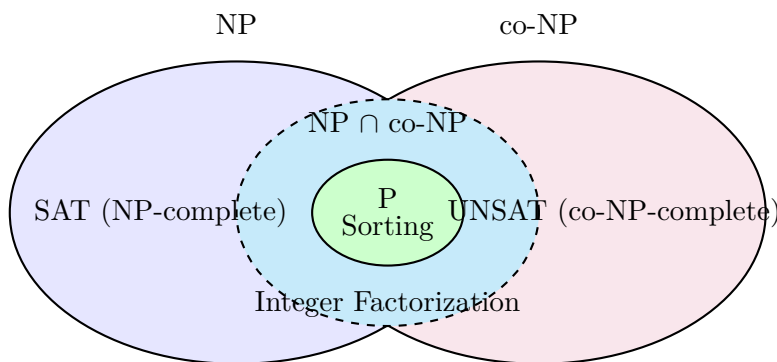


Figure 13.7: Relationship between NP, co-NP, and P. P is drawn within $\text{NP} \cap \text{coNP}$, which also includes problems like Integer Factorization. SAT is the canonical NP-complete problem; its complement UNSAT is co-NP-complete. It is unknown whether $\text{NP} = \text{coNP}$.

We present several examples of problems in co-NP, focusing on their complements in NP and their verification mechanisms.

Definition 13.31 (Tautology) *The Tautology problem asks: Given a Boolean formula ϕ in disjunctive normal form (DNF), is ϕ true for all possible truth assignments (i.e., is ϕ a tautology)?*

Definition 13.32 (UNSAT) *The Unsatisfiability (UNSAT) problem asks: Given a Boolean formula ϕ in conjunctive normal form (CNF), is ϕ unsatisfiable (i.e., no truth assignment makes ϕ true)?*

Definition 13.33 (Graph Non-Isomorphism) *The Non-Isomorphism problem for graphs asks: Given two graphs $G_1 = (V_1, E_1)$ and $G_2 = (V_2, E_2)$, are G_1 and G_2 not isomorphic (i.e., there exists no bijection $f : V_1 \rightarrow V_2$ preserving edges)?*

Theorem 13.34 *Tautology, UNSAT, and Graph Non-Isomorphism are all in co-NP.*

Proof: For each problem, the complement is in NP:

- The complement of Tautology is: Is ϕ not a tautology? A no instance (non-tautology) has a certificate: a truth assignment making ϕ false, verifiable in polynomial time by evaluating ϕ . Thus, Tautology is in co-NP.
- The complement of UNSAT is SAT, which is in NP: a yes instance of SAT has a certificate (a satisfying assignment) verifiable in polynomial time. Thus, UNSAT is in co-NP.
- The complement of Non-Isomorphism is Graph Isomorphism, which is in NP: a yes instance has a certificate (a bijection f) verifiable in polynomial time by checking edge preservation. Thus, Non-Isomorphism is in co-NP.

■

A decision problem L is *co-NP-complete* if $L \in \mathbf{coNP}$ and every $L' \in \mathbf{coNP}$ reduces to L in polynomial time. UNSAT and Tautology are both co-NP-complete: their complements (SAT and DNF non-tautology) are NP-complete, and a polynomial-time reduction from any NP-complete problem L' to SAT extends, after complementation, to a polynomial-time reduction from $\overline{L'}$ to UNSAT. It is not known if Non-Isomorphism is co-NP-complete.

Several open problems in complexity theory involve co-NP, reflecting its central role in understanding computational limits. We highlight key questions:

- Does $\mathbf{NP} = \mathbf{co-NP}$? The most significant open problem is whether $\mathbf{NP} = \mathbf{co-NP}$. If $\mathbf{NP} = \mathbf{co-NP}$, then for every NP-complete problem like SAT, its complement (UNSAT) would also be in NP, implying efficient certificates for unsatisfiability. This would collapse the polynomial hierarchy, as the hierarchy relies on the distinction between NP and co-NP. Most researchers believe $\mathbf{NP} \neq \mathbf{co-NP}$ due to the intuitive asymmetry: verifying satisfiability (NP) seems easier than verifying unsatisfiability (co-NP).
- Which problems lie in $\mathbf{NP} \cap \mathbf{co-NP}$? Known examples include Primality and Integer Factorization, but identifying others (e.g., Graph Isomorphism) is open. It is conjectured that $\mathbf{NP} \cap \mathbf{co-NP} \neq \mathbf{NP}$ and $\mathbf{NP} \cap \mathbf{co-NP} \neq \mathbf{co-NP}$, but the boundaries are unclear.

13.6 Coping with NP-hardness

A proof that a problem is NP-complete is rarely the end of the engineering conversation; it is the start of one. Practitioners use a five-strategy menu, all of which are explored elsewhere in the literature and several of which we develop in the rest of this book:

- **Approximation algorithms.** Trade optimality for polynomial running time. Chapter 14 develops this strategy in detail.
- **Parameterized / fixed-parameter tractable algorithms.** Identify a problem-specific parameter k (cover size, treewidth, solution depth) and design algorithms whose exponential cost is confined to k rather than the input size. The canonical example is Vertex Cover in $O(2^k \cdot (n + m))$ time, which is practical whenever the optimal cover is small.
- **Industrial SAT and ILP solvers.** Modern conflict-driven SAT solvers and branch-and-cut ILP solvers routinely dispatch instances with millions of variables, even though the worst-case problems are NP-hard. Encoding a target NP-complete problem as SAT or ILP and handing the result to such a solver is often the most cost-effective practical answer.
- **Special-case polynomial algorithms.** Many NP-complete problems become polynomial when the input is restricted: 2-SAT is in **P**, planar graph 3-coloring admits a $2^{O(\sqrt{n})}$ algorithm, and Vertex Cover on bipartite graphs is in **P** via König's theorem (Chapter 12). Recognizing the special case can save an exponential factor.
- **Heuristics and randomization.** When all else fails, local-search heuristics (simulated annealing, genetic algorithms, tabu search) and randomized rounding give solutions with no worst-case guarantees but often excellent empirical behaviour.

Chapter 14 treats the first strategy in depth; the others are pursued in the references at the end of the chapter.

13.7 Summary

This chapter introduced NP-completeness and the theory of computational intractability, covering the complexity classes **P**, **NP**, and **coNP**, and polynomial-time reductions. Approximation algorithms, the most prominent constructive response to NP-hardness, are developed in Chapter 14.

Figure 13.8 summarizes the chain of reductions established in this chapter and the catalogue. Each arrow $A \rightarrow B$ denotes a polynomial-time reduction $A \leq_P B$ used to transfer NP-hardness from A to B .

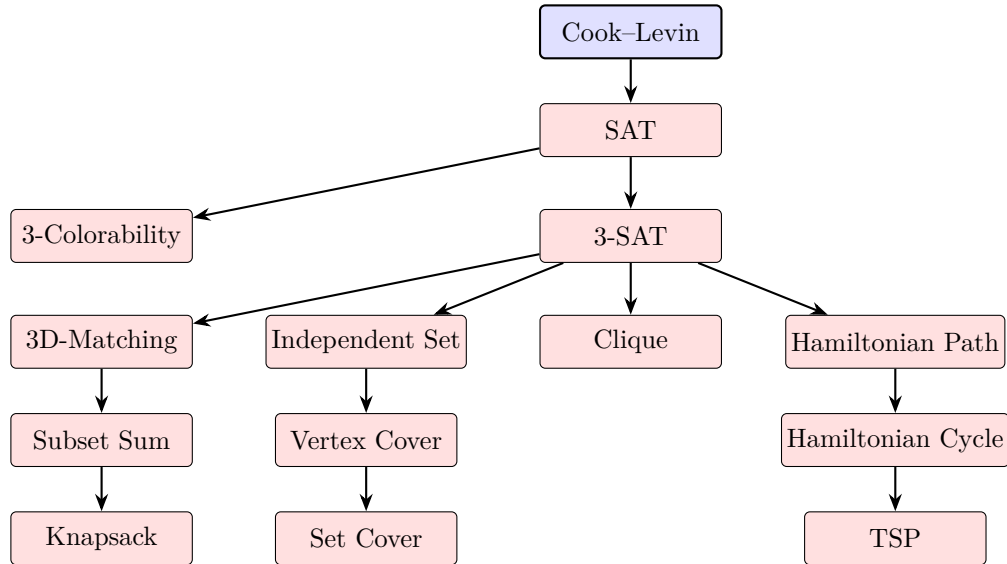


Figure 13.8: Reductions established in this chapter (and used in the catalogue). An arrow $A \rightarrow B$ means $A \leq_P B$, transferring NP-hardness from A to B . Cook–Levin (blue) seeds the chain by giving SAT’s NP-hardness from first principles; every other problem inherits NP-hardness through one of these reductions.

Problem/Topic	Result	Key Technique
SAT	NP-complete	Cook–Levin Theorem
3-SAT	NP-complete	Reduction from SAT
Clique	NP-complete	Reduction from 3-SAT
Independent Set	NP-complete	Reduction from 3-SAT
Vertex Cover	NP-complete	Reduction from Independent Set (duality)
Subset Sum	NP-complete	Reduction from 3D-Matching

13.8 Problems

1. You are given a graph $G = (V, E)$ with non-negative lengths $w_e \geq 0$ for all edges $e \in E$. You are also given a source node s and destination node t and an integer K . Show that deciding whether there is an s - t path of length exactly K is NP-complete.
2. Consider the *Job Scheduling with Deadlines* problem: Given n jobs with processing times p_i , deadlines d_i , and profits r_i , and a single processor, select a subset of jobs to maximize total profit such that each selected job completes by its deadline (job i takes p_i time units, must finish by d_i , and jobs are non-preemptive). Prove this problem is NP-complete.
3. (*Partition.*) Given a multiset of positive integers $A = \{a_1, \dots, a_n\}$, decide whether A can be partitioned into two subsets A_1, A_2 with $\sum_{a \in A_1} a = \sum_{a \in A_2} a$. Show that Partition is NP-complete. *Hint:* Partition is in NP (a candidate A_1 is a polynomial-time-verifiable certificate). For NP-hardness, reduce from Subset Sum: given (S, t) with $\sigma = \sum_{a \in S} a$, add one new element $|2t - \sigma|$ (or two padding elements) so that the augmented multiset has a balanced partition iff the original Subset Sum has a solution.

4. (*Knapsack* (decision version).) Given items with weights w_1, \dots, w_n , values v_1, \dots, v_n , capacity W , and threshold V , decide whether there is a subset $I \subseteq \{1, \dots, n\}$ with $\sum_{i \in I} w_i \leq W$ and $\sum_{i \in I} v_i \geq V$. Show that Knapsack is NP-complete.
Hint: Reduce from Subset Sum: given (S, t) , set $w_i = v_i = a_i$, $W = V = t$. A subset realises weight $\leq t$ and value $\geq t$ iff it sums to exactly t .
5. (*Hamiltonian Cycle*.) Given an undirected graph G , decide whether G has a Hamiltonian cycle (a cycle visiting every vertex exactly once). Show that Hamiltonian Cycle is NP-complete.
Hint: In NP — the cycle itself is the certificate. For NP-hardness, reduce from Hamiltonian Path: attach a new vertex u adjacent to every vertex of G ; G has a Hamiltonian path iff the new graph has a Hamiltonian cycle through u .
6. (*Travelling Salesman* (decision).) Given a complete weighted graph G and a bound B , decide whether G has a tour visiting every vertex exactly once with total weight at most B . Show that TSP is NP-complete.
Hint: Reduce from Hamiltonian Cycle. Given $G = (V, E)$, build a complete graph G' on V with edge weight $w(e) = 0$ if $e \in E$ and $w(e) = 1$ if $e \notin E$; set $B = 0$. A Hamiltonian cycle in G corresponds to a tour in G' of weight 0.
7. (*3-Colorability*.) Given an undirected graph G , decide whether the vertices of G can be coloured with three colours so that no edge has both endpoints of the same colour. Show that 3-Colorability is NP-complete.
Hint: Reduce from 3-SAT. Introduce three anchor vertices T, F, N forming a triangle (forcing them to take the three colour classes). For each variable x_i , add two vertices $x_i, \neg x_i$ joined by an edge and each joined to N (forcing one to colour T , the other to colour F). For each clause, add a six-vertex *OR-gadget* that is 3-colourable iff at least one of its three input literals is coloured T .
8. (*Dominating Set*.) Given a graph $G = (V, E)$ and an integer k , decide whether there exists $D \subseteq V$ with $|D| \leq k$ such that every vertex not in D has at least one neighbour in D . Show that Dominating Set is NP-complete.
Hint: Reduce from Vertex Cover. Given (G, k) , construct G' from G by subdividing each edge $\{u, v\}$ with a new vertex w_{uv} and adding the triangle $\{u, v, w_{uv}\}$; then G has a vertex cover of size $\leq k$ iff G' has a dominating set of size $\leq k$.
9. (*Max-Cut*.) Given an undirected graph G and an integer k , decide whether V can be partitioned into two sets A, B so that the number of edges with one endpoint in A and the other in B is at least k . Show that Max-Cut is NP-complete.
Hint: Max-Cut is in NP (a candidate bipartition is a certificate). For NP-hardness, reduce from Not-All-Equal 3-SAT (NAE-3-SAT), which is itself NP-complete by a reduction from 3-SAT. Each NAE clause produces a small graph gadget whose edges are cut precisely when the clause is “not-all-equal” satisfied.
10. (*Bin Packing* (decision)). Given n items of sizes $s_1, \dots, s_n \in (0, 1]$ and an integer k , decide whether the items can be packed into k unit-capacity bins. Show that Bin Packing is NP-complete (even for $k = 2$).
Hint: Reduce from Partition. Given a multiset A of positive integers with sum $2t$, scale each a_i to $a_i/(2t) \in (0, 1]$; the scaled items fit into $k = 2$ unit bins iff A admits a balanced partition.

11. (*Vertex Cover* \leq_P *Subset Sum*.) Give a polynomial-time reduction from Vertex Cover to Subset Sum. *Hint:* encode each vertex as a digit position and each edge as a number whose digits flag its two endpoints; choose a base larger than the maximum vertex degree to keep additions carry-free. This closes the alternative route to Subset Sum's NP-hardness mentioned in the chapter.
12. (*Vertex Cover is FPT*.) Give a $O(2^k \cdot (n+m))$ -time algorithm for deciding whether a graph $G = (V, E)$ has a vertex cover of size $\leq k$. *Hint:* pick any uncovered edge $\{u, v\}$ and branch on which of u, v is in the cover, decreasing k by 1 in each branch.
13. (*2-SAT is in P*.) Show that 2-SAT can be decided in $O(n+m)$ time by building the implication graph (literals as vertices, each clause $(\ell_1 \vee \ell_2)$ contributing edges $\bar{\ell}_1 \rightarrow \ell_2$ and $\bar{\ell}_2 \rightarrow \ell_1$) and computing strongly connected components: the formula is satisfiable iff no variable lies in the same SCC as its negation.

13.9 Bibliographic Remarks

The Cook-Levin Theorem (Theorem 13.1), independently established by Cook [Coo71] and Levin [Lev73], marks SAT as the first NP-complete problem, showing that every NP problem can be reduced to SAT in polynomial time. This result laid the groundwork for identifying other NP-complete problems, as detailed in the chapter's reductions (e.g., $\text{SAT} \leq_P \text{3-SAT}$, $\text{3-SAT} \leq_P \text{Independent Set}$, $\text{3-SAT} \leq_P \text{Clique}$). Karp [Kar72] significantly expanded this landscape by proving NP-completeness for 21 fundamental problems, including Clique, Vertex Cover, and Hamiltonian Path, many of which are covered in the chapter's Figure 13.2. For a comprehensive treatment of NP-completeness and its reductions, Garey and Johnson [GJ79] remains a definitive reference.

Polynomial-time reductions, as illustrated in Sections 13.2 and 13.3, are central to establishing the hardness of problems like Independent Set, Vertex Cover, and 3-SAT. These reductions exploit structural relationships, such as the complementary nature of Independent Set and Vertex Cover, to demonstrate computational equivalence. The chapter's treatment of co-NP (Section 13.4) introduces the dual class of problems with efficiently verifiable no certificates, with examples like UNSAT and Tautology. Papadimitriou [Pap94] offers an authoritative resource on complexity classes, including NP, co-NP, and their intersections.

Chapter 14

Approximation Algorithms

14.1 Introduction

A proof that an optimization problem is NP-hard is rarely the end of the engineering conversation: it is the start of one. *Approximation algorithms* respond to NP-hardness by trading optimality for polynomial running time. Instead of insisting on the exact optimum, an approximation algorithm computes, in polynomial time, a feasible solution whose cost is provably within a guaranteed factor of the optimum.

This chapter develops the basic theory of approximation algorithms and works through several canonical examples lifted from Chapter 13. We assume the reader is familiar with the NP-completeness of Vertex Cover, Set Cover, Knapsack, and Traveling Salesman; for these problems the goal of this chapter is constructive: we do not just prove they are hard, we hand the practitioner an algorithm that returns a solution with a guarantee.

The chapter is organized as follows. Section 14.2 defines the approximation ratio formally and distinguishes minimization from maximization variants. Section 14.3 gives a 2-approximation for minimum Vertex Cover via maximal matching. Section 14.4 gives the greedy H_n -approximation for Set Cover. Section 14.5 introduces approximation schemes (PTAS / FPTAS) and sketches the FPTAS for Knapsack. Section 14.6 casts the Vertex Cover 2-approximation as a parallel LLP algorithm using lexicographic edge order, and gives parallel LLP variants of Set Cover and Knapsack. Section 14.7 surveys hardness-of-approximation results, including Håstad's tight bound for Max-3-SAT and the $(1 - \alpha) \ln n$ lower bound for Set Cover.

14.2 The Approximation Ratio

Definition 14.1 (Approximation Ratio) Let Π be a minimization optimization problem and A a polynomial-time algorithm that, on every instance I , outputs a feasible solution of cost $A(I)$. Let $OPT(I)$ denote the cost of an optimal solution to I . We say A is an α -approximation algorithm for Π (with $\alpha \geq 1$) if for every instance I ,

$$A(I) \leq \alpha \cdot OPT(I).$$

The constant α is called the approximation ratio of A . For a maximization problem, the condition is $A(I) \geq \frac{1}{\alpha} \cdot OPT(I)$.

The smaller α is, the stronger the guarantee. An exact polynomial-time algorithm is a 1-approximation; for an NP-hard problem we expect $\alpha > 1$. Some problems admit constant-factor approximations independent of the input size; others admit only $O(\log n)$ -approximations; others admit no constant-factor approximation at all unless $P = NP$. The question of the best achievable α is itself a deep area of complexity theory, the so-called *hardness of approximation*, treated briefly in Section 14.7.

14.3 Vertex Cover

Recall (Chapter 13) that the minimum Vertex Cover problem asks, given an undirected graph $G = (V, E)$, for the smallest subset $C \subseteq V$ such that every edge of G has at least one endpoint in C . The decision version is NP-complete, hence the optimization version is NP-hard. We give a clean 2-approximation.

Algorithm ApproxVertexCover: ApproxVertexCover(G)

Input: Graph $G = (V, E)$

Output: Vertex cover C

```

1  $C := \emptyset$  ;
2  $E' := E$  ;
3 while  $E' \neq \emptyset$  do
4   | Pick any edge  $(u, v) \in E'$  ;
5   |  $C := C \cup \{u, v\}$  ;
6   | Remove all edges incident to  $u$  or  $v$  from  $E'$  ;
7 end
8 return  $C$ 
```

Time: $O(m)$.

Theorem 14.2 *ApproxVertexCover is a 2-approximation algorithm for the minimum Vertex Cover problem.*

Proof: Let F be the set of edges picked by the algorithm in successive iterations. Because the algorithm removes every edge incident to the endpoints of a picked edge before the next iteration, no two edges of F share an endpoint; i.e., F is a matching. The algorithm outputs C consisting of both endpoints of each edge in F , so $|C| = 2|F|$.

Every vertex cover C^* must contain at least one endpoint of each edge in F , and since the edges of F are pairwise disjoint, $|C^*| \geq |F|$. In particular, $OPT \geq |F|$. Therefore $|C| = 2|F| \leq 2 \cdot OPT$, proving the

2-approximation guarantee. ■

Tightness. The factor 2 is tight for this algorithm: on a complete bipartite graph $K_{n,n}$, the algorithm picks all $2n$ vertices, while the optimum is n . More refined approximations (LP-rounding, semi-definite programming) shave the constant slightly, but no polynomial algorithm is known to achieve a $(2 - \epsilon)$ -approximation under the Unique Games Conjecture.

Example 14.3 [ApproxVertexCover on a small graph] Take $V = \{1, 2, 3, 4, 5\}$ with $E = \{(1, 2), (1, 3), (2, 4), (3, 4), (3, 5)\}$. The minimum vertex cover has size 3 (e.g., $\{1, 4, 5\}$ does not cover all edges; $\{2, 3, 4\}$ does).

Iter 1. $E' = E$. Pick the first edge $(1, 2)$; add $\{1, 2\}$ to C . Remove every edge incident to 1 or 2: $(1, 2), (1, 3), (2, 4)$ are removed. $E' = \{(3, 4), (3, 5)\}$.

Iter 2. Pick $(3, 4)$; add $\{3, 4\}$ to C . Remove $(3, 4)$ and $(4, 5)$ (incident to 4). $E' = \emptyset$.

The output $C = \{1, 2, 3, 4\}$ has size 4. The matching $F = \{(1, 2), (3, 4)\}$ has size 2, so $|C| = 2|F| = 4 \leq 2 \cdot OPT = 6$. The algorithm's ratio on this instance is $4/3$, well within the 2-approximation guarantee.

14.4 Set Cover

Set Cover generalizes Vertex Cover: given a universe U of n elements and a collection \mathcal{S} of subsets of U , find the smallest sub-collection $\mathcal{C} \subseteq \mathcal{S}$ whose union is U . (Vertex Cover is the special case in which each element of U corresponds to an edge and each set in \mathcal{S} to the incident edges of a vertex.) Set Cover admits a clean greedy approximation: repeatedly pick the set in \mathcal{S} that covers the most uncovered elements, until every element is covered.

Algorithm ApproxSetCover: ApproxSetCover(U, \mathcal{S})

Input: Universe U , collection \mathcal{S} of subsets of U

Output: A sub-collection $\mathcal{C} \subseteq \mathcal{S}$ covering U

```

1  $\mathcal{C} := \emptyset; R := U;$ 
2 while  $R \neq \emptyset$  do
3   | Pick  $S \in \mathcal{S}$  maximizing  $|S \cap R|;$ 
4   |  $\mathcal{C} := \mathcal{C} \cup \{S\}; R := R \setminus S;$ 
5 end
6 return  $\mathcal{C}$ 

```

Theorem 14.4 *ApproxSetCover returns a cover of size at most $H_n \cdot OPT$, where $H_n = \sum_{k=1}^n 1/k = O(\ln n)$ is the n -th harmonic number and $n = |U|$.*

The proof, due to Chvátal [Chv79], charges each newly covered element a fee of $1/|S \cap R|$ at the moment S is picked; the total fee equals $|\mathcal{C}|$, while a careful averaging argument shows the fee on any optimal set is at most $H_n \cdot 1$, yielding $|\mathcal{C}| \leq H_n \cdot OPT$.

Comparison with Vertex Cover. Specialising `ApproxSetCover` to a `Vertex Cover` instance gives an $O(\log n)$ -approximation, weaker than the 2-approximation of Section 14.3. The 2-approximation exploits the special structure of `Vertex Cover` (each element appears in exactly two sets) that the general greedy ignores. This pattern recurs: domain-specific structure often beats the generic greedy.

Example 14.5 [`ApproxSetCover` on a small instance] Take $U = \{1, 2, 3, 4, 5, 6\}$ and $\mathcal{S} = \{S_1 = \{1, 2, 3\}, S_2 = \{1, 4, 5\}, S_3 = \{2, 4, 6\}, S_4 = \{3, 5, 6\}\}$. The optimal cover is $\{S_2, S_3\}$ with $|OPT| = 2$.

Iter 1. $R = U$. Coverage: $|S_1 \cap R| = 3$, $|S_2 \cap R| = 3$, $|S_3 \cap R| = 3$, $|S_4 \cap R| = 2$. Tied at 3; pick $S_1 = \{1, 2, 3\}$ (any tie-breaker). $\mathcal{C} = \{S_1\}$, $R = \{4, 5, 6\}$.

Iter 2. Coverage: $|S_2 \cap R| = 2$, $|S_3 \cap R| = 2$, $|S_4 \cap R| = 1$. Pick $S_2 = \{1, 4, 5\}$. $\mathcal{C} = \{S_1, S_2\}$, $R = \{6\}$.

Iter 3. Pick either S_3 or S_4 ; pick S_3 . $R = \emptyset$.

The output is $\mathcal{C} = \{S_1, S_2, S_3\}$ of size 3. The bound $H_6 = 1 + 1/2 + \dots + 1/6 \approx 2.45$ gives $|\mathcal{C}| \leq H_6 \cdot OPT \approx 4.9$; we used 3 sets, comfortably under the bound.

14.5 Approximation Schemes: PTAS and FPTAS

A constant-factor approximation is sometimes too coarse. Two finer-grained notions make the ratio a tunable parameter.

Definition 14.6 (PTAS) A polynomial-time approximation scheme (PTAS) for a minimization problem Π is a family of algorithms $\{A_\epsilon\}_{\epsilon > 0}$ such that, for every fixed $\epsilon > 0$, A_ϵ runs in time polynomial in the input size and outputs a $(1 + \epsilon)$ -approximate solution.

Definition 14.7 (FPTAS) A fully polynomial-time approximation scheme (FPTAS) is a PTAS whose running time is polynomial in both the input size and $1/\epsilon$.

The distinction matters in practice: a PTAS may have a running time like $O(n^{1/\epsilon})$, blowing up rapidly as $\epsilon \rightarrow 0$, whereas an FPTAS keeps the dependence on $1/\epsilon$ polynomial.

An FPTAS for Knapsack via value scaling

Recall the Knapsack problem from Chapter 13: given n items with weights w_1, \dots, w_n , profits v_1, \dots, v_n , and a knapsack capacity W , find a subset $S \subseteq [n]$ with $\sum_{i \in S} w_i \leq W$ maximising $\sum_{i \in S} v_i$. Knapsack admits a standard pseudo-polynomial dynamic program in $O(nV)$ time, where $V = \sum_i v_i$. The FPTAS scales the profit values down so that the DP table becomes polynomially-bounded, while losing only a $(1 - \epsilon)$ factor in the objective.

Let $M = \max_i v_i$ be the maximum profit of any single item; clearly $M \leq OPT \leq nM$. Define the scaling factor

$$\mu = \frac{\epsilon M}{n}$$

and replace each profit by its scaled-and-rounded counterpart

$$v'_i = \left\lfloor \frac{v_i}{\mu} \right\rfloor.$$

Now run the standard $O(nV')$ DP on the scaled instance, where $V' = \sum_i v'_i$.

Algorithm FPTAS-Knapsack: Fully polynomial-time approximation scheme for Knapsack via value scaling.

Input: Items (w_i, v_i) for $i \in [n]$, capacity W , accuracy parameter $\epsilon > 0$

Output: Subset $S \subseteq [n]$ with profit at least $(1 - \epsilon) \cdot OPT$

1 $M := \max_i v_i; \quad \mu := \epsilon M/n;$

2 **forall** $i \in [n]$ **do**

3 $v'_i := \lfloor v_i/\mu \rfloor$

4 **end**

5 $S :=$ optimal Knapsack solution for (w_i, v'_i, W) via the standard $O(nV')$ DP;

6 **return** S

Theorem 14.8 *FPTAS-Knapsack returns a feasible solution of profit at least $(1 - \epsilon) \cdot OPT$ in $O(n^3/\epsilon)$ time.*

Proof: Let S^* be an optimal solution of the original instance and S the solution returned by the DP on the scaled instance. For each i we have $\mu v'_i \leq v_i < \mu v'_i + \mu$, so $v_i - \mu \leq \mu v'_i \leq v_i$.

Since S is optimal under the scaled values,

$$\sum_{i \in S} v'_i \geq \sum_{i \in S^*} v'_i.$$

Multiplying by μ and using $v_i \geq \mu v'_i$ on the left and $\mu v'_i \geq v_i - \mu$ on the right,

$$\sum_{i \in S} v_i \geq \mu \sum_{i \in S^*} v'_i \geq \sum_{i \in S^*} (v_i - \mu) = OPT - n\mu = OPT - \epsilon M \geq (1 - \epsilon) \cdot OPT,$$

since $OPT \geq M$.

For the running time, $V' = \sum_i v'_i \leq n \cdot M/\mu = n^2/\epsilon$, so the DP runs in $O(nV') = O(n^3/\epsilon)$. ■

Example 14.9 [FPTAS-Knapsack on a 3-item instance] Take $n = 3$ items with $(w_i, v_i) = (2, 30), (3, 40), (4, 50)$ and capacity $W = 6$. The original DP $O(nV)$ has $V = 120$. The optimum on this instance is $S^* = \{1, 2\}$ with profit 70.

Choose accuracy $\epsilon = 0.2$. Then $M = 50$ and $\mu = \epsilon M/n = 0.2 \cdot 50/3 \approx 3.33$. The scaled values are

$$v'_1 = \lfloor 30/3.33 \rfloor = 9, \quad v'_2 = \lfloor 40/3.33 \rfloor = 12, \quad v'_3 = \lfloor 50/3.33 \rfloor = 15.$$

The scaled DP picks the same set $S = \{1, 2\}$ (since $9 + 12 = 21 > 15$) for capacity 6. Recovering original profits: $\sum_{i \in S} v_i = 30 + 40 = 70$. The bound $(1 - \epsilon) \cdot OPT = 0.8 \cdot 70 = 56$ is comfortably satisfied; in fact this scaling preserved optimality exactly.

The benefit of the scaling: instead of running the DP up to $V = 120$, the scaled DP runs only up to $V' = 9 + 12 + 15 = 36$, a $3.3\times$ reduction. For larger instances the gap grows: at $V = 10^6$ and $\epsilon = 0.2$, $V' \leq n^2/\epsilon$ stays polynomial, while the unscaled DP is pseudo-polynomial.

Lattice-scaling viewpoint. The algorithm has a clean LLP interpretation: the original $O(nV)$ DP runs on a lattice of states indexed by item-prefix and accumulated profit, of size $\Theta(nV)$. Rescaling the values by μ shrinks the profit axis from V down to $V/\mu \leq n^2/\epsilon$, decreasing the lattice size by a factor of μ while perturbing each profit by less than μ . The DP solves the same lattice-linear feasibility predicate on the smaller lattice, and the bound $\sum v_i - n\mu \leq \sum \mu v'_i$ tracks the cumulative loss across the n items. This is the FPTAS template in its purest form: shrink the lattice, pay a controlled error, recover a polynomial-time scheme.

Many geometric and scheduling problems admit a PTAS but not an FPTAS unless $P = NP$; see [DPW10] for a comprehensive treatment of the design space.

14.6 An LLP Perspective on Approximation

The approximation algorithms presented so far are sequential. Several of them admit clean parallel formulations within the LLP framework, retaining the same approximation guarantee while operating on the lattice of vertex (or edge) inclusion vectors. This section gives the LLP perspective for Vertex Cover; later sections of this chapter will return to Set Cover and Knapsack.

Why minimum Vertex Cover is not lattice-linear

Recall that an LLP algorithm computes the minimum element of a meet-semilattice of feasible states. A natural attempt is to take the state space to be vectors $G \in \{0, 1\}^n$ where $G[v] = 1$ iff v is in the cover, and the predicate $B(G) =$ “every edge has at least one endpoint with $G[v] = 1$ ”. The set of vertex covers is closed under join (componentwise OR) but *not* under meet: $G_1 = (1, 0, 1, 0)$ and $G_2 = (0, 1, 0, 1)$ may both be vertex covers of a 4-cycle, yet their meet $(0, 0, 0, 0)$ is not. So minimum Vertex Cover is not directly an LLP problem.

Lexically-First Vertex Cover

A small workaround restores the LLP structure. Order the edges lexicographically (first by smaller endpoint, breaking ties by the larger endpoint), so $(1, 3) < (1, 5) < (2, 3) < \dots$. Call an edge (u, v) *lex-minimal at G* if it is the lex-smallest among all currently-uncovered edges sharing an endpoint with (u, v) . The algorithm is: in parallel, select every lex-minimal edge and add both endpoints to the cover;

repeat until no edge remains uncovered.

Algorithm LLP-LexicallyFirstVertexCover: A parallel 2-approximation for Vertex Cover via lex-minimal edges.

Input: Graph $G_{\text{in}} = (V, E)$ with vertices labelled $1, \dots, n$

Output: $G[1..n] \in \{0, 1\}$: a vertex cover

- 1 G : array[1.. n] of $\{0, 1\}$ initially $\forall v : G[v] = 0$;
 - 2 $\text{uncovered}(u, v) \equiv (u, v) \in E \wedge G[u] = 0 \wedge G[v] = 0$;
 - 3 $\text{lexmin}(u, v) \equiv \text{uncovered}(u, v) \wedge \forall (x, y) \in E : (\text{uncovered}(x, y) \wedge \{u, v\} \cap \{x, y\} \neq \emptyset) \Rightarrow (u, v) \leq_{\text{lex}} (x, y)$;
 - 4 **forbidden**(v): $\exists (u, v) \in E : \text{lexmin}(u, v)$;
 - 5 **advance**(v): $G[v] := 1$;
 - 6 **return** $\{v : G[v] = 1\}$
-

The selected edges form a matching. The key structural lemma is that, in any single round, the set of lex-minimal edges is pairwise non-adjacent.

Lemma 14.10 *At any state G , no two lex-minimal edges share an endpoint.*

Proof: Suppose (u, v) and (v, w) are both lex-minimal at G . Each is uncovered; each shares the endpoint v with the other; hence each is among the neighbors of the other. Lex-minimality of (u, v) gives $(u, v) \leq_{\text{lex}} (v, w)$, and lex-minimality of (v, w) gives $(v, w) \leq_{\text{lex}} (u, v)$. So $(u, v) = (v, w)$, contradicting their distinctness. ■

Lemma 14.10 reduces the LLP-LexicallyFirstVertexCover algorithm, in a single round, to picking a matching F of lex-minimal edges and adding all endpoints to the cover — exactly the structure used by the sequential ApproxVertexCover of Section 14.3. The 2-approximation argument carries over verbatim.

Theorem 14.11 *LLP-LexicallyFirstVertexCover is a 2-approximation algorithm for the minimum Vertex Cover problem.*

Proof: Let $F = F_1 \cup F_2 \cup \dots \cup F_r$ be the set of all edges selected as lex-minimal in any round. By Lemma 14.10, each F_i is a matching; by induction on rounds, edges across different F_i are also disjoint — because once a vertex enters the cover at round i , every edge incident to it is covered at round $i + 1$ and cannot be selected later. So F is a matching, and the algorithm outputs $C = \bigcup_{(u,v) \in F} \{u, v\}$ with $|C| = 2|F|$. Every vertex cover C^* contains an endpoint of every edge in F and these endpoints are distinct, so $|C^*| \geq |F|$. Hence $|C| = 2|F| \leq 2 \cdot \text{OPT}$. ■

Termination. The lex-minimal edge of E itself — the globally smallest uncovered edge — is trivially lex-minimal at G , so at least one edge is selected in each round provided E is non-empty. After a round, both endpoints of every selected edge are in the cover, so every edge incident to either endpoint is covered.

Hence at least one edge per round is removed from the uncovered set, and the algorithm terminates in at most $|E|$ rounds. In typical inputs the count of lex-minimal edges in a single round is $\Theta(|E|/\deg_{\text{avg}})$, giving an effective parallel depth of $O(\Delta \log n)$ on graphs of bounded degree.

Comparison with the sequential algorithm. LLP-LexicallyFirstVertexCover and the sequential ApproxVertexCover differ only in the choice of which edge to pick at each step: the sequential version picks an arbitrary uncovered edge; the parallel version picks every lex-minimal one in a single round. Both produce a matching, both pay a factor of at most 2 over the optimum, and both run in polynomial work. The lex-minimal rule is what gives the parallel version a deterministic, well-defined LLP advance condition.

Example 14.12 [LLP-LexicallyFirstVertexCover on a small graph] Take $V = \{1, 2, 3, 4, 5\}$ with edges $E = \{(1, 2), (1, 3), (2, 4), (3, 4), (4, 5)\}$. Sort lexicographically: $(1, 2) < (1, 3) < (2, 4) < (3, 4) < (4, 5)$.

Round 1. The lex-min edge of the whole graph is $(1, 2)$, with neighbors $(1, 3)$ and $(2, 4)$; $(1, 2)$ is smaller than both, so $(1, 2)$ is lex-minimal. Among edges not sharing an endpoint with $(1, 2)$, the next candidate is $(3, 4)$: its neighbors are $(1, 3)$, $(2, 4)$, and $(4, 5)$, all lex-larger than $(3, 4)$, so $(3, 4)$ is also lex-minimal. The two selected edges $(1, 2)$ and $(3, 4)$ are disjoint (Lemma 14.10). Set $G[1] = G[2] = G[3] = G[4] = 1$.

Round 2. Edges $(1, 3)$, $(2, 4)$, $(3, 4)$, $(1, 2)$ are now covered. Only $(4, 5)$ remains uncovered, but $G[4] = 1$ already, so the edge is in fact covered. No edge is uncovered, so no index is forbidden, and the algorithm terminates.

The output is $C = \{1, 2, 3, 4\}$ of size 4, while $OPT = 3$ (e.g. $\{1, 4, ?\}$); the ratio $4/3$ is well within the 2-approximation guarantee.

Lexically-First Set Cover

The lex-min idea generalises directly to Set Cover. The sequential greedy of Section 14.4 picks one set at a time; the parallel LLP analogue picks every set that is locally maximal in coverage and lex-minimal among ties. Two such sets can never share an uncovered element, so they can be added to the cover simultaneously.

Let R denote the set of currently uncovered elements. Two sets $s, s' \in \mathcal{S}$ are *neighbors at G* if $(s \cap R) \cap (s' \cap R) \neq \emptyset$, i.e., they compete for at least one uncovered element. Define

$$\text{lexmaxcov}(s) \equiv (s \cap R \neq \emptyset) \wedge \forall s' \text{ neighbor of } s: (|s \cap R| > |s' \cap R|) \vee (|s \cap R| = |s' \cap R| \wedge s \leq_{\text{lex}} s').$$

A set is selected in a round iff it is *lexmaxcov*.

Algorithm LLP-LexicallyFirstSetCover: A parallel H_n -approximation for Set Cover via lex-first locally-maximal sets.

Input: Universe U , collection $\mathcal{S} = \{s_1, \dots, s_m\}$

Output: $G[1..m] \in \{0, 1\}$: a sub-collection of \mathcal{S} that covers U

1 G : array[1.. m] of $\{0, 1\}$ initially $\forall i : G[i] = 0$;

2 $R(G) = U \setminus \bigcup_{i:G[i]=1} s_i$;

// currently uncovered elements

3 **forbidden**(i): *lexmaxcov*(s_i);

4 **advance**(i): $G[i] := 1$;

5 **return** $\{s_i : G[i] = 1\}$

Selected sets are pairwise disjoint on R . The matching argument from Lemma 14.10 carries over.

Lemma 14.13 *At any state G , no two *lexmaxcov* sets share an uncovered element.*

Proof: Suppose s and s' are both *lexmaxcov* and $(s \cap R) \cap (s' \cap R) \neq \emptyset$. Then they are neighbors. From s 's local optimality, $|s \cap R| > |s' \cap R|$ or ties with $s \leq_{\text{lex}} s'$. From s' 's local optimality, the symmetric inequality. The two strict cases contradict each other, so $|s \cap R| = |s' \cap R|$ and $s \leq_{\text{lex}} s' \leq_{\text{lex}} s$, hence $s = s'$. ■

Approximation guarantee. The parallel version retains the H_n bound of the sequential greedy.

Theorem 14.14 *LLP-LexicallyFirstSetCover returns a cover of size at most $H_n \cdot \text{OPT}$, where H_n is the n -th harmonic number and $n = |U|$.*

Proof: We adapt Chvátal's charging argument. When set s is selected in round t with current uncovered set R_t , charge each newly-covered element $x \in s \cap R_t$ a fee of $1/|s \cap R_t|$. The total fee equals the number of selected sets, i.e., $|\mathcal{C}|$.

Now bound the total fee on any single optimal set S^* . Order the elements of S^* as x_1, x_2, \dots, x_p by the round in which they are first covered: $t_1 \leq t_2 \leq \dots \leq t_p$. At round t_i , x_i is covered by some selected set s containing x_i . Since x_i is also in S^* and uncovered at round t_i , the sets s and S^* share an uncovered element x_i , so S^* is a neighbor of s at round t_i . Local maximality of s then gives $|s \cap R_{t_i}| \geq |S^* \cap R_{t_i}| \geq p - i + 1$ (since x_i, x_{i+1}, \dots, x_p are still uncovered). Hence the fee on x_i is at most $1/(p - i + 1)$, and the total fee on S^* is at most $\sum_{i=1}^p 1/(p - i + 1) = H_p \leq H_n$.

Summing over the OPT optimal sets covers every element of U at least once, so $|\mathcal{C}| = \text{total fee} \leq H_n \cdot \text{OPT}$. ■

Vertex Cover as a special case. Vertex Cover is the special case in which the universe is the edge set E , each set $s_v = \{e : v \in e\}$ corresponds to a vertex v , and every element is in exactly two sets. Specialising LLP-LexicallyFirstSetCover to this case selects, in each round, every *vertex* that is locally

maximal in degree-among-uncovered-edges and lex-minimal among ties — a strictly different selection rule from LLP-LexicallyFirstVertexCover, which selects *edges*. Both yield 2-approximations on Vertex Cover, but via different parallel structures: the edge-selection rule gives a matching of edges; the vertex-selection rule gives an independent set of vertices that all dominate at least one uncovered edge.

Example 14.15 [LLP-LexicallyFirstSetCover on the running instance] Take the same instance as Example 14.5: $U = \{1, \dots, 6\}$ and $\mathcal{S} = \{S_1 = \{1, 2, 3\}, S_2 = \{1, 4, 5\}, S_3 = \{2, 5, 6\}, S_4 = \{3, 6\}\}$. Order the sets by index: $S_1 <_{\text{lex}} S_2 <_{\text{lex}} S_3 <_{\text{lex}} S_4$.

Round 1. $R = U$. Coverage: $|S_1 \cap R| = |S_2 \cap R| = |S_3 \cap R| = 3$, $|S_4 \cap R| = 2$.

- S_1 neighbours S_2 (share element 1), S_3 (share 2), S_4 (share 3). Among neighbours, S_1 ties at 3 with S_2, S_3 and beats S_4 ; ties broken by lex-min, S_1 wins. So S_1 is *lexmaxcov*.
- S_2 neighbours S_1, S_3 . Tied at 3 with S_1 , but $S_1 <_{\text{lex}} S_2$, so S_2 is not *lexmaxcov*.
- S_3 similarly loses to S_1 .
- S_4 has $|S_4 \cap R| = 2 < 3 = |S_1 \cap R|$ and they share element 3, so S_4 is not *lexmaxcov* either.

Only S_1 is selected. Set $G[1] := 1$. $R = \{4, 5, 6\}$.

Round 2. Coverage on $R = \{4, 5, 6\}$: $|S_2 \cap R| = 2$, $|S_3 \cap R| = 2$, $|S_4 \cap R| = 1$. S_2 and S_3 share 5, both at coverage 2; lex-min picks S_2 . S_4 is dominated. So S_2 is selected. $G[2] := 1$. $R = \{6\}$.

Round 3. Only S_3 and S_4 contain 6; both have coverage 1, so both share an uncovered element. By Lemma 14.13 only one is *lexmaxcov* — S_3 wins by lex order. $G[3] := 1$. $R = \emptyset$.

The output is $\mathcal{C} = \{S_1, S_2, S_3\}$ of size 3, identical to Example 14.5. The bound $H_6 \approx 2.45 \cdot \text{OPT} = 4.9$ is satisfied. Note that on this small instance only one set is selected per round, but in larger instances multiple disjoint *lexmaxcov* sets fire in parallel.

LLP-ApproxKnapsack via lattice scaling

The FPTAS-Knapsack of Section 14.5 fits cleanly into the LLP framework. The state is a two-dimensional table $G[i, w]$ for $i \in [0..n]$ and $w \in [0..W]$, where $G[i, w]$ records the maximum profit attainable using items $\{1, \dots, i\}$ with total weight at most w . The lattice-linear feasibility predicate is

$$G[i, w] \geq \max(G[i-1, w-w_i] + v_i, G[i-1, w]) \quad \text{for } w \geq w_i$$

and $G[i, w] \geq G[i-1, w]$ otherwise; the boundary conditions are $G[0, w] = G[i, 0] = 0$. The minimum table satisfying this predicate is exactly the table of optimal Knapsack values, with $G[n, W]$ as the answer.

LLP-ApproxKnapsack pre-processes the values to shrink the lattice along the profit axis before running the standard LLP scheduler. Define $M = \max_i v_i$ and the scaling factor $\mu = \epsilon M/n$; replace each profit v_i

by the integer $v'_i = \lfloor v_i/\mu \rfloor$ and run the LLP advance loop on the scaled instance.

Algorithm LLP-ApproxKnapsack: Parallel LLP FPTAS for Knapsack via value scaling.

Input: Items (w_i, v_i) for $i \in [n]$, capacity W , accuracy $\epsilon > 0$

Output: G : array $[0 \dots n, 0 \dots W]$ of int with $\mu \cdot G[n, W] \geq (1 - \epsilon) \cdot OPT$

1 $M := \max_i v_i$; $\mu := \epsilon M/n$;

2 **forall** $i \in [n]$ **do**

3 | $v'_i := \lfloor v_i/\mu \rfloor$

4 **end**

5 $G[i, w] := 0$ for all $i \in [0..n], w \in [0..W]$;

6 **forbidden** (i, w) : $(w \geq w_i \wedge G[i, w] < \max(G[i - 1, w - w_i] + v'_i, G[i - 1, w]))$
 $\vee (w < w_i \wedge G[i, w] < G[i - 1, w])$;

7 **advance** (i, w) : $G[i, w] := \max(G[i - 1, w - w_i] + v'_i, G[i - 1, w])$ if $w \geq w_i$; $G[i, w] := G[i - 1, w]$ otherwise;

8 **return** G

Approximation guarantee. The bound carries over from Theorem 14.8. The scaled instance satisfies $\mu v'_i \leq v_i < \mu v'_i + \mu$, so for any subset S that LLP-ApproxKnapsack reports as optimal under v' ,

$$\sum_{i \in S} v_i \geq \mu \sum_{i \in S} v'_i \geq \mu \sum_{i \in S^*} v'_i \geq \sum_{i \in S^*} v_i - n\mu = OPT - \epsilon M \geq (1 - \epsilon) \cdot OPT,$$

where S^* is the original optimum and the second inequality is the optimality of S under v' .

Lattice-shrinkage perspective. The pre-processing step is a lattice contraction: each v_i becomes its quantised representative $\lfloor v_i/\mu \rfloor$, so the value axis shrinks from height V to height $V' = O(n^2/\epsilon)$. The forbidden / advance rules are otherwise unchanged — they run on a lattice that is asymptotically polynomially-sized regardless of how large the original profits were. On inputs with V super-polynomial in n (the pseudo-polynomial worst case), this is exponentially smaller than the unscaled version.

Example 14.16 [LLP-ApproxKnapsack on the running instance] Take the same instance as Example 14.9: items $(w_i, v_i) = (2, 30), (3, 40), (4, 50)$ with capacity $W = 6$ and accuracy $\epsilon = 0.2$. The pre-processing computes $M = 50$, $\mu = 0.2 \cdot 50/3 \approx 3.33$, $v'_1 = 9$, $v'_2 = 12$, $v'_3 = 15$.

Initial state. The lattice $G[i, w]$ is initialized to 0 for all $i \in [0..3]$, $w \in [0..6]$.

Forbidden / Advance trace. Each cell (i, w) with $i \geq 1$ is forbidden until it satisfies the lower bound. The advance writes $\max(G[i-1, w-w_i] + v'_i, G[i-1, w])$ when $w \geq w_i$, else $G[i-1, w]$. After the advance loop converges:

$G[i, w]:$	$w=0$	1	2	3	4	5	6
$i = 0$	0	0	0	0	0	0	0
$i = 1$ ($w_1=2, v'_1=9$)	0	0	9	9	9	9	9
$i = 2$ ($w_2=3, v'_2=12$)	0	0	9	12	12	21	21
$i = 3$ ($w_3=4, v'_3=15$)	0	0	9	12	15	21	24

The answer is $G[3, 6] = 21$, achieved by $S = \{1, 2\}$ with scaled profit $9 + 12 = 21$. The corresponding original profit is $30 + 40 = 70$, exactly OPT on this instance. The $(1 - \epsilon) \cdot OPT = 56$ bound is comfortably exceeded.

The pre-scaling shrinks the lattice from $V = 120$ down to $V' = 36$ along the value axis; on this small instance, both DPs run essentially the same number of cells, but as V grows the gap becomes asymptotic.

14.7 Hardness of Approximation

For some optimization problems, even getting close to the optimum is NP-hard:

- **General TSP.** Approximating the minimum-weight tour to within *any* constant factor is NP-hard, by reducing Hamiltonian Cycle to TSP with edge weights 1 for present edges and K for absent edges; choosing K large enough rules out approximation.
- **Metric TSP.** When weights satisfy the triangle inequality, Christofides' algorithm gives a $\frac{3}{2}$ -approximation; recent work tightened this to $\frac{3}{2} - 10^{-36}$.
- **Max-3-SAT.** Approximating the maximum number of simultaneously satisfiable clauses within a factor better than $\frac{7}{8}$ is NP-hard (Håstad's theorem); a random assignment achieves exactly $\frac{7}{8}$, so this hardness is tight.
- **Set Cover.** The H_n ratio above is essentially optimal: approximating Set Cover within $(1 - \alpha) \ln n$ for any $\alpha > 0$ is NP-hard.
- **Vertex Cover.** Approximating within a factor better than $\sqrt{2}$ is known to be NP-hard, and the Unique Games Conjecture would push this to $2 - \epsilon$, matching Section 14.3.

These lower bounds rely on the PCP theorem (probabilistically checkable proofs); the modern theory is treated in Arora–Barak [AB09].

14.8 Summary

This chapter introduced the theory and practice of approximation algorithms.

Problem	Algorithm	Approximation Ratio
Vertex Cover	ApproxVertexCover (matching)	2
Set Cover	ApproxSetCover (greedy)	H_n ($\approx \ln n$)
Knapsack	FPTAS via rounding + DP	$1 + \epsilon$
General TSP	—	no constant factor
Max-3-SAT	Random assignment	$8/7$ (tight)

The general lesson is that NP-hardness is the start of the engineering conversation, not its end. Some problems (Knapsack, Vertex Cover) admit excellent approximations; others (General TSP, Set Cover beyond $\ln n$) provably do not. The boundary is itself a deep area of complexity theory.

14.9 Problems

1. Show that the 2-approximation factor of ApproxVertexCover is tight by constructing a family of graphs on which the algorithm returns a cover of size exactly $2 \cdot OPT$.
2. Suppose the input to Set Cover guarantees that every element of U appears in at most f sets. Show that ApproxVertexCover-style argument gives an f -approximation. In particular, when $f = 2$ the problem reduces to Vertex Cover and we recover the 2-approximation.
3. Prove that the H_n analysis of ApproxSetCover is tight: exhibit a family of instances on which the greedy returns a cover of size approximately $H_n \cdot OPT$.
4. The PTAS for the Euclidean TSP runs in time roughly $n^{O(1/\epsilon)}$. Why is this only a PTAS and not an FPTAS? What does Garey–Johnson tell us about FPTAS-existence for strongly NP-hard problems?
5. Show that any maximization problem Π admits a polynomial-time approximation iff its decision version is in NP. (This is essentially trivial; the point is to internalize the relationship between optimization and decision.)
6. The Steiner Tree problem asks for the minimum-weight tree spanning a designated subset R of the vertices in an edge-weighted graph. Give a 2-approximation by reducing to MST on the metric closure restricted to R .

14.10 Bibliographic Remarks

The ApproxVertexCover algorithm and its 2-approximation guarantee are folklore; the matching-based proof appears already in Gavril (unpublished, ca. 1974) and Bar-Yehuda–Even [BYE85]. The greedy Set Cover analysis is due to Chvátal [Chv79] (and independently Johnson [Joh74] and Lovász [Lov75]). The FPTAS for Knapsack was given by Ibarra and Kim [IK75b].

The hardness-of-approximation results rely on the PCP theorem of Arora, Lund, Motwani, Sudan, and Szegedy [ALM⁺98]; Håstad’s tight $\frac{7}{8}$ bound for Max-3-SAT appears in [Hås01]. Vazirani’s monograph [Vaz03] is the standard reference for the design and analysis of approximation algorithms; Williamson and Shmoys [WS11] give a complementary modern treatment with strong emphasis on LP-based methods.

Chapter 15

The Housing Allocation Problem

15.1 Introduction

Imagine a residential complex of n tenants, each currently living in one of n apartments and each holding strong opinions about which of the other apartments they would prefer. Or imagine n research interns assigned to lab benches, n teams sharing n time-slots on a piece of expensive equipment, or n students rotating through n projects — in each setting, every participant arrives with a property they consider their own and a private ranking of every other participant’s property. The question is whether and how the participants can *swap among themselves* to reach an allocation that no group can collectively beat by trading inside the group. This is the *housing allocation problem*. Its appeal is that it is purely cooperative: there is no money, no central planner choosing winners and losers, and no participant ends up worse off than when they started. It captures the structure of one-sided matching with initial endowments cleanly enough that the same algorithm reappears in kidney-exchange markets, on-campus housing assignment, and the allocation of compute slots in shared HPC clusters.

The housing market problem proposed by Shapley and Scarf [SS74] is a matching problem with one-sided preferences. There are n agents and n houses. Each agent a_i initially owns a house h_i for $i \in \{1, \dots, n\}$ and has a completely ranked list of houses. There are variations of this problem when the agents do not own any house initially. In this book, we focus on the version with the initial endowment of houses for agents. The list of agent preferences is given by $pref[i][k]$ which specifies the k^{th} preference of the agent i . Thus, $pref[i][1] = j$ means that a_i prefers h_j as his top choice. The goal is to come up with an optimal house allocation such that each agent has a house, and no subset of agents can improve the satisfaction of agents in this subset by exchanging houses within the subset. It can be shown that there is a unique such matching called the *core* for any housing market. The standard algorithm for this problem is Gale’s Top Trading Cycle Algorithm (TTC), which takes $O(n^2)$ time. This algorithm is optimal in terms of time complexity, since the input size is $O(n^2)$.

This chapter is organized as follows. Section 15.2 describes Gale’s Top Trading Cycle Algorithm. Section 15.3 applies the LLP method to the housing market problem.

15.2 Gale’s Top Trading Cycle Algorithm

Consider the housing market instance shown in Fig. 15.1. There are four agents a_1, a_2, a_3 and a_4 . Initially, the agent a_i holds the house h_i . The preferences of the agents are shown in Fig. 15.1.

$a_1 : h_2, h_3, h_1, h_4$	$a_1 : h_1$	$a_1 : h_2$
$a_2 : h_1, h_4, h_2, h_3$	$a_2 : h_2$	$a_2 : h_1$
$a_3 : h_1, h_2, h_4, h_3$	$a_3 : h_3$	$a_3 : h_4$
$a_4 : h_2, h_1, h_3, h_4$	$a_4 : h_4$	$a_4 : h_3$
(i) Agents’ Preferences	(ii) Initial Allocation	(iii) Matching returned by TTC

Figure 15.1: Housing Market and the Matching returned by the Top Trading Cycle Algorithm

The Top Trading Cycle (TTC) algorithm attributed to Gale by Shapley and Scarf [SS74] works in stages. At each stage, it has the following steps:

Step 1. We construct the *top choice* directed graph $G_t = (A, E)$ on the set of agents A as follows. We add a directed edge from agent $a_i \in A$ to agent $a_j \in A$ if a_j holds the current top house of a_i . Fig. 15.2 shows the directed graph in the first stage.

Step 2. Since each node has exactly one outgoing edge in G_t , there is at least one cycle in the graph (possibly a self-loop). All cycles are node disjoint. We find all the cycles in the top trading graph and implement the trade indicated by the cycles, that is, each agent that is in any cycle gets its current top house.

Step 3. Remove all agents who get their current top houses, and remove all houses which are assigned to some agent from the preference list of remaining agents.

The above steps are repeated until each agent is assigned a house. At each stage, at least one agent is assigned a final house, so there are at most n stages. Within a stage, building G_t , finding all cycles in it, and updating preference lists all take $O(n)$ time on the unassigned agents (each $wish[i]$ pointer only moves forward across stages, so its total movement across the run of the algorithm is $O(n)$ per agent). The overall running time is therefore $O(n^2)$, which is optimal in the input size.

Beyond its running time, the TTC mechanism has three properties that make it the algorithm of choice for housing-market problems in practice. (i) *Individual rationality*: every agent ends up with a house at least as preferred as their endowment h_i (in the worst case, an agent’s own house remains its top choice and forms a self-loop in G_t). (ii) *Pareto-optimality*: no allocation makes some agent strictly better off without making another strictly worse off. (iii) *Strategy-proofness*: a self-interested agent has no incentive to misreport preferences — truthful reporting is a dominant strategy [Rot82]. The combination of (i)–(iii) is rare among matching mechanisms; it is the reason TTC reappears in kidney exchange, school choice, and dormitory-allocation systems (see Section 15.6).

Algorithm **TTC** states the procedure formally.

Algorithm TTC: Gale's Top Trading Cycle Algorithm

Input: $pref[i][k]$: the k^{th} choice of agent a_i , for $i, k \in [1..n]$. By renumbering houses, agent a_i initially owns house h_i .

Output: $house[1..n]$: $house[i]$ is the house finally assigned to agent a_i .

```

1  $fixed[1..n]$ : array of boolean, initially  $\forall i : fixed[i] := false$ ;
2  $wish[1..n]$ : array of int, initially  $\forall i : wish[i] := pref[i][1]$ ; // current top remaining choice
3 while  $\exists i : \neg fixed[i]$  do
    // Step 1: build the top-choice graph  $G_t$  on unassigned agents
4   forall  $i : \neg fixed[i]$  do
5     | advance  $wish[i]$  down  $pref[i]$  until  $wish[i]$  is held by some unassigned agent  $a_j$ ;
6     | add edge  $a_i \rightarrow a_j$  to  $G_t$ ;
7   end
    // Step 2: every node has out-degree 1, so  $G_t$  contains at least one cycle;
    //           cycles are node-disjoint
8    $C :=$  set of agents lying on some cycle of  $G_t$ ;
    // Step 3: implement the trades indicated by the cycles and remove the
    //           participants
9   forall  $i \in C$  do
10    |  $house[i] := wish[i]$ ;
11    |  $fixed[i] := true$ ;
12  end
13 end
14 return  $house$ ;

```

Example 15.1 [TTC on the running instance] Consider the four-agent instance of Fig. 15.1. Each a_i initially owns h_i and the preference lists are $a_1 : h_2, h_3, h_1, h_4$; $a_2 : h_1, h_4, h_2, h_3$; $a_3 : h_1, h_2, h_4, h_3$; $a_4 : h_2, h_1, h_3, h_4$.

Stage 1. Top-choice graph (Fig. 15.2): $a_1 \rightarrow a_2$, $a_2 \rightarrow a_1$, $a_3 \rightarrow a_1$, $a_4 \rightarrow a_2$. The unique cycle is $\{a_1, a_2\}$, so a_1 takes h_2 and a_2 takes h_1 . Both are fixed and removed.

Stage 2. Only a_3 and a_4 remain, with houses h_3 and h_4 unallocated. Top-choice now: $a_3 \rightarrow a_4$ (top remaining is h_4), $a_4 \rightarrow a_3$ (top remaining is h_3). The cycle $\{a_3, a_4\}$ assigns $a_3 \mapsto h_4$ and $a_4 \mapsto h_3$.

The final core allocation is $a_1:h_2$, $a_2:h_1$, $a_3:h_4$, $a_4:h_3$.

15.3 An LLP Algorithm for the Housing Market Problem

There are n agents and n houses. Each agent proposes houses in the decreasing order of preferences. These proposals are considered events executed by n agents. Thus, we have n events per agent. Each event is labeled as (i, h, k) , which corresponds to the agent i proposing to the house h as his choice number k .

The global state corresponds to the number of proposals made by each of the agents. Let $G[i]$ be the number of proposals made by the agent i . We will assume that in the initial state, every agent has made

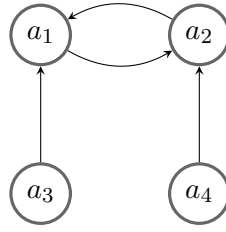


Figure 15.2: The top choice graph at the first stage.

his first proposal. Thus, the initial global state $G = [1, 1, \dots, 1]$. We extend the notation of indexing to subsets $J \subseteq [n]$ such that $G[J]$ corresponds to the subvector given by the indices in J .

We now model the possibility of reallocation of houses based on any global state. Recall that $pref[i][k]$ specifies the k^{th} preference of the agent a_i . Let $wish(G, i)$ denote the house proposed by a_i in the global state G , that is,

$$wish(G, i) = pref[i][G[i]]$$

A global state G satisfies *matching* if every agent proposes a different house, that is,

$$matching(G) \equiv \forall i, j : i \neq j : wish(G, i) \neq wish(G, j).$$

We generalize *matching* to refer to a subset of agents rather than the entire set.

Definition 15.2 (submatching) Let $J \subseteq [n]$. Then *submatching*(G, J) iff $wish(G, J)$ is a permutation of the indices in J .

Intuitively, if *submatching*(G, J) holds, then all agents in J can exchange houses within the subset J . For any G , it is easy to show that

Lemma 15.3 For all G , there always exists a nonempty J such that *submatching*(G, J).

Proof: Given any G , we can create a directed graph as follows. The set of vertices is agents, and there is an edge from i to j if $wish(G, i) = j$. There is exactly one outgoing edge from any vertex in $[n]$ to $[n]$ in this graph. This implies that there is at least one cycle in this graph (possibly a self-loop). The indices of agents in the cycle give us such a subset J . ■

We now show that

Lemma 15.4 *submatching*(G, J_1) and *submatching*(G, J_2) implies *submatching*($G, J_1 \cup J_2$).

Proof: Any index $i \in J_1 \cup J_2$ is mapped to J_1 if $i \in J_1$ and J_2 , otherwise. ■

Hence, there exists the biggest submatching in G . Note that *matching*(G) is equivalent to *submatching*($G, [n]$).

Definition 15.5 (Feasible Global State) *A global state G is feasible for the housing market problem iff it is a matching and for all global states $F < G$, there does not exist any submatching which is better in F than in G . Note that if there exists a submatching J that is better in F than in G , then agents in J can improve their allocation by just exchanging houses within the subset J . Formally, let*

$$B_{\text{housing}}(G) \equiv \text{matching}(G) \wedge (\forall F < G : \forall J \subseteq [n] : \text{submatching}(F, J) \Rightarrow F[J] = G[J]).$$

We show that $B_{\text{housing}}(G)$ is a lattice-linear predicate. This result will let us use the lattice-linear predicate detection algorithm for the housing market problem.

Theorem 15.6 *The predicate $B_{\text{housing}}(G)$ is lattice-linear.*

Proof: Suppose that $\neg B_{\text{housing}}(G)$. This implies that either G is not a matching or it is a matching, but there exists a smaller global state F that has a submatching better than G .

First, consider the case when G is not a matching. Let J be the largest set such that $\text{submatching}(G, J)$. Consider any index $i \notin J$ such that $\text{wish}(G, i) \in J$. We claim $\text{forbidden}(G, i, B_{\text{housing}})$. Let H be any global state greater than G such that $G[i] = H[i]$. We consider two cases.

Case 1: $H[J] > G[J]$.

Then, from the second conjunct of B_{housing} , we know that $\neg B_{\text{housing}}(H)$ because $\text{submatching}(G, J)$ and $H[J] \neq G[J]$.

Case 2: $H[J] = G[J]$.

Since $\text{wish}(H, i) = \text{wish}(G, i)$, $\text{wish}(G, i) \in J$, and $G[J] = H[J]$, we get that H is not a matching because the house given by $\text{wish}(G, i)$ is also in the wish list of some agent in J .

Now consider the case where G is a matching, but $\neg B_{\text{housing}}(G)$. This implies,

$$\exists F < G : \exists J \subseteq [n] : \text{submatching}(F, J) \wedge F[J] < G[J].$$

However, the same F will also result in guaranteeing $\neg B_{\text{housing}}(H)$ for any $H \geq G$. ■

It is also easy to see from the proof that if an index is part of a submatching, then it will never become forbidden.

This theorem gives us the algorithm shown in Algorithm [LLP-Housing-Market-Algorithm](#). Let G be the initial global state. Let $S(G)$ be the biggest submatching in G . All agents such that they are not in $S(G)$ and wish a house which is part of $S(G)$ are forbidden and can move on to their next proposal. The algorithm terminates when no agent is forbidden. This algorithm is a parallel version of the top trading cycle (TTC) mechanism attributed to Gale in [SS74].

Algorithm LLP-Housing-Market-Algorithm: An LLP algorithm to find the optimal housing market

Input: $pref[i][k]$: the k^{th} preference of agent i

Output: $G[1..n]$: proposal vector giving the core allocation

```

1  $G$ : array[1.. $n$ ] of int initially 1 ; // every agent starts with the top choice
2  $wish(G, i) = pref[i][G[i]]$ ;
3  $S(G) =$  largest  $J$  such that  $submatching(G, J)$ ;
4 forbidden( $j$ ): ( $j \notin S(G)$ )  $\wedge$  ( $wish(G, j) \in S(G)$ );
5 advance( $j$ ):  $G[j] := G[j] + 1$ ;
```

Example 15.7 [LLP-Housing-Market on the running instance] On the four-agent instance of Example 15.1 we trace the proposal vector G :

$G = [1, 1, 1, 1]$. $wish : a_1 \rightarrow h_2, a_2 \rightarrow h_1, a_3 \rightarrow h_1, a_4 \rightarrow h_2$. The largest submatching is $S(G) = \{1, 2\}$ (the cycle $a_1 \leftrightarrow a_2$). Agents 3 and 4 are not in $S(G)$ and their wishes lie in $S(G)$, so both are forbidden.

Advance: $G[3], G[4] := 2$.

$G = [1, 1, 2, 2]$. $wish : a_3 \rightarrow h_2, a_4 \rightarrow h_1$. $S(G) = \{1, 2\}$ still; agents 3, 4 remain forbidden. Advance again.

$G = [1, 1, 3, 3]$. $wish : a_3 \rightarrow h_4, a_4 \rightarrow h_3$. Now $S(G) = \{1, 2, 3, 4\}$ (two cycles: $a_1 \leftrightarrow a_2$ and $a_3 \leftrightarrow a_4$). No agent is forbidden; the algorithm terminates. Reading off $pref[i][G[i]]$ gives the same core allocation $a_1:h_2, a_2:h_1, a_3:h_4, a_4:h_3$ as Example 15.1.

We now show that

Theorem 15.8 *There exists at least one feasible global state G such that $B_{housing}(G)$.*

Proof: Every agent has his own house on the list of preferences. If he ever makes a proposal to his own house, he forms a submatching. That particular event is never forbidden because it is part of a submatching. Hence, the lattice-linear predicate detection algorithm will never mark that event as forbidden. Since such an event exists for all processes, we are guaranteed to never go beyond this global state. ■

The above proof also shows that agents can never be worse off participating in the algorithm. Each agent will either get his own house back or get a house that he prefers to his own house.

We now show that the feasible state is in fact unique — justifying the claim of *the* core in Section 15.2.

Corollary 15.9 *The feasible global state G satisfying $B_{housing}(G)$ is unique.*

Proof: Suppose, for contradiction, that two distinct global states U and V both satisfy $B_{housing}$. Let $W = U \sqcap V$ be their componentwise meet, so $W \leq U$ and $W \leq V$ with $W \neq U$ or $W \neq V$. Because $B_{housing}$ is lattice-linear (Theorem on $B_{housing}$ in this section), it is closed under meet, and hence W also satisfies $B_{housing}$ — in particular $matching(W)$ holds. But then $W < U$ is a global state strictly smaller than U in which every component still proposes a matched house; the second conjunct of $B_{housing}(U)$

requires $U[J] = W[J]$ for every $J \subseteq [n]$ with $submatching(W, J)$, taking $J = [n]$ this forces $U = W$. By the same argument $V = W$, contradicting $U \neq V$. ■

15.4 Summary

This chapter presented algorithms for the housing market problem, which finds the unique core allocation where no subset of agents can improve by trading among themselves.

Problem	Algorithm	Time Complexity
Housing Market	Gale's Top Trading Cycle	$O(n^2)$
Housing Market	LLP Top Trading Cycle	$O(n^2)$

15.5 Problems

1. Consider a housing market with four agents, $\{A_1, A_2, A_3, A_4\}$, who own houses $\{h_1, h_2, h_3, h_4\}$, where A_i owns h_i . Preferences are:

Agent	Preference List
A_1	$h_2 \succ h_4 \succ h_1 \succ h_3$
A_2	$h_1 \succ h_3 \succ h_2 \succ h_4$
A_3	$h_3 \succ h_1 \succ h_4 \succ h_2$
A_4	$h_3 \succ h_2 \succ h_4 \succ h_1$

Table 15.1: Preferences for the agents.

Apply the TTC algorithm to find the final allocation.

2. Is the outcome of the TTC algorithm unique for a given set of truthful preferences and initial endowments? Justify your answer.
3. Give an algorithm to determine whether the given matching is the core of the housing market.
4. Prove that the allocation returned by the TTC algorithm is Pareto optimal.
5. Prove or disprove: The Top Trading Cycle algorithm is strategyproof, i.e., no agent can benefit by misreporting their preferences.
6. (**Serial Dictatorship.**) In the *Serial Dictatorship* mechanism with a fixed priority order $\pi = (a_{\pi(1)}, \dots, a_{\pi(n)})$, agents are processed one at a time; each agent chooses her most preferred house among the houses still available. Prove that the resulting allocation is Pareto efficient and strategyproof.
7. (**YRMH-IGYT with Existing Tenants.**) Consider a house-allocation problem with a mix of existing tenants (who own their current house) and newcomers (who do not own any house), plus some vacant houses. The *You Request My House, I Get Your Turn* (YRMH-IGYT) mechanism, given a priority order, works as follows: for each agent in order, either she is assigned her top available house (if that house is vacant or unoccupied by a still-unassigned tenant), or a *request* is made to

the current owner who then jumps to the front of the queue. Show that YRMH-IGYT respects the individual rationality of existing tenants (no tenant gets a house she likes strictly less than her endowment).

8. **(Sensitivity to Priority Order in TTC.)** In the TTC mechanism for housing markets, the initial endowment is fixed, so the outcome is independent of how we pick among multiple disjoint cycles in one step. Show, however, that for *Serial Dictatorship* (no initial endowments), different priority orders can give Pareto-incomparable allocations. Provide a small example.
9. **(Individual Rationality of TTC.)** Prove that the allocation μ returned by the TTC algorithm is *individually rational*, i.e., every agent weakly prefers $\mu(i)$ to her endowed house h_i .
10. **(Uniqueness of the Core.)** Prove that in any housing market (with strict preferences), the TTC allocation is the *unique* core allocation. That is, no matching other than the TTC outcome is in the core.

15.6 Bibliographic Remarks

The housing market problem has been studied by many researchers [SS74, HZ79, Zho90, AS98, AS99, RP77, Rot82, Dav13]. Possible applications of the housing market problem include assigning virtual machines to servers in cloud computers, allocating graduates to trainee positions, professors to offices, and students to roommates. This problem has also been studied by Zheng and Garg [ZG19] where it is shown that the problem of verifying that a matching is a core is in NC, but the problem of computing the core is CC-hard. The class CC (Comparator Circuits) is the complexity class containing decision problems that can be solved by comparator circuits of polynomial size. The paper [ZG19] also provides a *distributed* message passing algorithm to find the core with $O(n^2)$ messages. A algorithm is described in [Gar21].

The mechanism-design properties of TTC — individual rationality, Pareto-optimality, and strategy-proofness (Roth [Rot82]) — have made it the basis of several deployed allocation systems. Roth, Sönmez, and Ünver [RSU04] adapted TTC to *kidney-paired donation*, where incompatible donor–recipient pairs swap kidneys to enable transplants that would otherwise be impossible; the mechanism is now used by national kidney-exchange registries. Abdulkadirouğlu and Sönmez [AS99] extended TTC to *college dormitory allocation*, where some students hold endowments (returning residents) while others do not (incoming first-years). Pathak and Sönmez [PS08] analyse the role of TTC in the *school-choice* debate around the Boston mechanism, the version of which (TTC vs. Deferred Acceptance) shaped the redesign of public school assignment in Boston, New Orleans, and elsewhere.

Chapter 16

The Assignment Problem

16.1 Introduction

The assignment problem arises whenever a set of indivisible resources must be matched one-to-one with a set of agents who value them differently. Sponsored-search ad slots are auctioned to advertisers; donor kidneys must be matched with compatible recipients; ride-hailing platforms dispatch drivers to riders; faculty are assigned to courses; and GPU jobs are mapped onto cluster nodes. In each case, every (resource, agent) pair carries a numeric value, and the goal is to choose a one-to-one assignment that maximizes the total.

In this chapter, we consider the assignment problem in which we seek an assignment of items to bidders such that the total payoff is maximized. We view items and bidders as nodes in a bipartite graph. The weight $v_{b,i}$ of an edge between the bidder b and the item i is the utility of assigning the item i to the bidder b . The assignment problem then corresponds to finding the maximum weight bipartite matching.

The assignment problem has a rich history. The most famous algorithm for this problem is due to the American scientist Kuhn, who called it the Hungarian algorithm because it is inspired by the earlier works of two Hungarian mathematicians, Konig and Egervary. Another American mathematician, Munkres, observed that the algorithm is strongly polynomial, and the algorithm is also known as the Kuhn-Munkres algorithm. It was later discovered that the problem had previously been solved by Jacobi in the 19th century and was published in Latin in the year 1890.

This chapter is organized as follows. Section 16.2 formulates the assignment problem as a maximum weight bipartite matching. Section 16.3 describes the LLP-based algorithm for finding the minimum market clearing price vector and illustrates it on a small worked example. Section 16.4 generalizes the market clearing price problem to incorporate lattice-linear constraints.

16.2 Problem Formulation

Let I be a set of n indivisible items and U , a set of n bidders. Each item $i \in I$ receives a valuation $v_{b,i}$ from each bidder $b \in U$. The valuation of any item i is an integer between 0 and T . Each item i has a price $G[i]$ which is also an integer between 0 and T .

We will assume that the number of items is equal to the number of bidders. If one of the sets is smaller, say there are fewer items than bidders, then we can add virtual items such that all bidders give a valuation

of zero to those items. We also note that we are considering the maximum weight bipartite matching. By subtracting the valuation from the maximum valuation for each edge and calling it the cost of that edge, we can transform this problem to minimum-cost bipartite matching.

It is important to note that it is the relative valuation of the items that is important, not the absolute values. Suppose that there are three items and a bidder provides a valuation of $(10, 8, 4)$ for these items, then the assigned matching will not change if his valuation is $(6, 4, 0)$ instead. The weight of the matching found for the valuation $(6, 4, 0)$ would be exactly less by 4 because in each assignment a bidder is assigned exactly one item. Hence, one can assume that there is at least one item for each bidder that is valued as 0. Similarly and dually, suppose that an item is valued by three bidders as $(10, 8, 4)$. Then, the assignment will not change if this item is valued as $(6, 4, 0)$ instead. If we view V as a square matrix such that $V[b, i]$ equals $v_{b,i}$, then we can assume that there is a zero in every row by subtracting the minimum value in each row from every entry in the row. Similarly, we can assume that there is at least one zero in every column.

16.3 Market Clearing Price

In this section, we apply the LLP technique to the problem of finding a market clearing price. The set of feasible price vectors is given by

$$\{G \mid 0 \leq G[i] \leq T, 1 \leq i \leq n\}$$

This set of price vectors forms a distributive lattice under the component-wise comparison of the vectors. The minimum is given by the zero vector and the maximum is given by the vector T .

Given a price vector G , we define the bipartite graph $(I, U, E(G))$ as follows. One side of the bipartite graph is the set of items I . The other side of the graph is the set of bidders U . We now add edges between items and bidders as follows.

$$(j, b) \in E(G) \equiv \forall i : (v_{b,j} - G[j]) \geq (v_{b,i} - G[i]).$$

Informally, there is an edge between item j and bidder b if the payoff for the bidder (the bid minus the price) is maximized with that item. Given any set $U' \subseteq U$, let $N(U', G)$ denote all the items that are adjacent to the vertices in U' in the graph $(I, U, E(G))$. A price vector G is a *market clearing price* if the bipartite graph $(I, U, E(G))$ has a perfect matching.

We construct a computation graph (E, \rightarrow) for this problem as follows. There are n processes corresponding to n items in the computation, and each process has T events. If the process i has executed k events, then $G[i] = k$, so the global state G can be viewed as the price vector G with $G[i] = k$. A price vector G is a *market clearing price*, denoted by $B_{\text{clearingPrice}}(G)$, if the bipartite graph $(I, U, E(G))$ has a perfect matching. The unconstrained instance has no cross-process edges; we will see the more interesting computation graph in Fig. 16.1 once constraints are introduced in Section 16.4. We now claim that

Lemma 16.1 *The predicate $B_{\text{clearingPrice}}(G)$ is a lattice-linear predicate on the lattice of price vectors.*

Proof: Suppose that the price vectors G and H satisfy $B_{\text{clearingPrice}}$. Let $K = \min(G, H)$. We show that K also satisfies $B_{\text{clearingPrice}}$. Suppose that $(I, U, E(K))$ does not have a perfect matching. This implies that there exists a minimal overdemanded set of items in K . Let an item i be one of the minimal overdemanded items in K . Without loss of generality, assume that $K[i]$ equals $G[i]$ (the argument for the case where $K[i]$ equals $H[i]$ is identical). The item i is not an overdemanded item in G . But this is a

contradiction because for all other items i' , the price of the item i' has either stayed the same or increased in going from K to G .

■

Algorithm ConstrainedMarketClearingPrice: finding the minimum cost assignment vector

Input: $v[b, i]$: int for all b, i
Output: $G[1..n]$: minimum clearing price vector

- 1 **shared var** G : array[1..n] of 0..maxint initially $G[j] := 0$;
- 2 $E = \{(k, b) \mid \forall i : (v[b, k] - G[k]) \geq (v[b, i] - G[i])\}$;
- 3 $demand(U') = \{k \mid \exists b \in U' : (k, b) \in E\}$;
- 4 $overDemanded(J) \equiv \exists U' \subseteq U : (demand(U') = J) \wedge (|J| < |U'|)$;
- 5 **forbidden**(j): \exists minimal J : $OverDemanded(J) \wedge (j \in J)$;
- 6 **advance**(j): $G[j] := G[j] + 1$;

In Fig. [ConstrainedMarketClearingPrice](#), we have used $\alpha(G, j)$ as simply one unit of price. For any item j that is part of a minimal over-demanded set of items, we can increase its price by the minimum amount to ensure that some bidder b can switch to her second most preferred item. We now give an implementation in Algorithm [LLP-Assignment](#) based on this idea.

Algorithm LLP-Assignment: Finding the minimum clearing price vector

Input: $v[b, i]$: valuations for all bidders b and items i
Output: $G[1..n]$: minimum clearing price vector; M : perfect matching

- 1 G : real initially $\forall i : G[i] = 0$;
- 2 **while true do**
- 3 $E := \{(i, b) \mid \forall j : (v_{b,i} - G[i]) \geq (v_{b,j} - G[j])\}$;
- 4 **if** there exists a perfect matching M in E **then return** M ;
- 5 **else**
- 6 $J :=$ minimal OverDemanded set in the bipartite graph with edges $E(G)$;
- 7 **forall** $j \in J$ **do**
- 8 $\delta_j = \min\{(v_{b,j} - G[j]) - \max_{i \neq j} (v_{b,i} - G[i]) \mid (j, b) \in E(G)\}$;
- 9 $G[j] := G[j] + \delta_j$;
- 10 **end**
- 11 **end**
- 12 **end**

Example 16.2 [LLP-Assignment on a 3×3 instance] We trace LLP-Assignment on three buyers $\{b_1, b_2, b_3\}$ and three items $\{h_1, h_2, h_3\}$ with valuation matrix

$$V = \begin{array}{c|ccc} & h_1 & h_2 & h_3 \\ \hline b_1 & 10 & 8 & 6 \\ b_2 & 9 & 7 & 4 \\ b_3 & 8 & 6 & 5 \end{array}$$

At prices $G = (p_1, p_2, p_3)$, buyer b_i demands $D(b_i) = \arg \max_j (v_{ij} - p_j)$; the algorithm raises prices on a minimal overdemanded set until the demands admit a perfect matching.

Step 0. $G = (0, 0, 0)$. All three buyers demand h_1 , so $J = \{h_1\}$ is overdemanded.

Step 1. Raise p_1 . After two unit advances, $G = (2, 0, 0)$ gives utilities

$$\begin{array}{c|ccc} & h_1 & h_2 & h_3 \\ \hline b_1 & 8 & 8 & 6 \\ b_2 & 7 & 7 & 4 \\ b_3 & 6 & 6 & 5 \end{array}$$

Now $D(b_1) = D(b_2) = D(b_3) = \{h_1, h_2\}$, so $J = \{h_1, h_2\}$ is overdemanded.

Step 2. Raise both p_1 and p_2 by 1, giving $G = (3, 1, 0)$. Utilities become

$$\begin{array}{c|ccc} & h_1 & h_2 & h_3 \\ \hline b_1 & 7 & 7 & 6 \\ b_2 & 6 & 6 & 4 \\ b_3 & 5 & 5 & 5 \end{array}$$

Now $D(b_1) = D(b_2) = \{h_1, h_2\}$ and $D(b_3) = \{h_1, h_2, h_3\}$. The demand graph admits the perfect matching $b_1 \rightarrow h_1$, $b_2 \rightarrow h_2$, $b_3 \rightarrow h_3$, and the algorithm returns $G = (3, 1, 0)$ as the minimum market clearing price.

Termination and complexity. The unit-step Algorithm [ConstrainedMarketClearingPrice](#) (`ConstrainedMarketClearingPrice`) raises a price by 1 per advance, so it terminates after at most nT advances and runs in time $O(n^{3.5}T)$ — weakly polynomial, since it depends on the magnitude T of the valuations. Algorithm [LLP-Assignment](#) (`LLP-Assignment`) avoids this dependence by jumping directly to the next critical price: the advance amount δ_j is exactly the slack at which at least one bidder b in the demand of j becomes indifferent to some other item, so each advance adds a new tight edge to the equality graph $E(G)$. The standard Hungarian-method analysis then applies. Define a *phase* as the iterations between two successive increases in the cardinality of a maximum matching in $E(G)$; there are at most n phases. Within a phase, each advance grows the alternating tree rooted at unmatched bidders by at least one new vertex, so at most n advances per phase suffice before an augmenting path is found and the matching size grows. Hence LLP-Assignment terminates in $O(n^2)$ iterations, with each iteration costing $O(n^2)$ work for slack computation and $O(n^{2.5})$ for the matching test, giving $O(n^{4.5})$ overall and $O(n^3)$ with careful incremental bookkeeping (matching the Kuhn–Munkres bound). The complexity is *strongly polynomial*: it depends only on n , not on the size of the valuations. The best known strongly polynomial bound for the assignment problem is

$O(n^3)$ [Mun57, EK72], which LLP-Assignment achieves with the standard incremental implementation.

16.4 Constrained Market Clearing Price Problem

We now generalize the problem of finding a market clearing price to that of finding a constrained market clearing price. For example, constraints on the clearing prices of the form $(G[j] \geq k) \Rightarrow (G[i] \geq k')$ for $1 \leq k, k' \leq C$ are lattice-linear. The constraint says that if item j is priced at least k , then item i must be priced at least k' . The constraint $G[i] \geq G[j]$ is also lattice-linear. Observe that the constraint $G[i] \geq G[j]$ is equivalent to

$$(G[j] \geq 1 \Rightarrow G[i] \geq 1) \wedge (G[j] \geq 2 \Rightarrow G[i] \geq 2) \dots (G[j] \geq C \Rightarrow G[i] \geq C)$$

Similarly, the constraint $(G[i] = G[j])$ can be modeled as $(G[i] \geq G[j]) \wedge (G[j] \geq G[i])$. Also, a constraint of the form $G[j] \geq f(G)$ for monotone f is lattice-linear. Given any set of valuations, and a boolean predicate B that is a conjunction of lattice-linear constraints, a price vector G is a *constrained market clearing price*, denoted by $\text{constrainedClearing}(G)$ iff $\text{clearing}(G) \wedge B(G)$. Since $B(G)$ is lattice-linear, it is sufficient to give an algorithm for $\text{clearing}(G)$. It follows that the set of constrained market clearing price vectors is closed under meets. By applying the lattice-linear predicate detection, we get an algorithm to compute the least constrained market clearing price shown in Fig. [ConstrainedMarketClearingPrice](#). We get a generalization of Demange, Gale and Sotomayor's exact auction mechanism [DGS86] to incorporate lattice-linear constraints on the market clearing price.

We construct a computation graph (E, \rightarrow) for this problem as follows (see Fig. 16.1). For any constraint, $(G[j] \geq k) \Rightarrow (G[i] \geq k')$, we put an edge from the event k' in P_i to the event k in P_j . By putting such edges, we get a computation graph and any price vector that does not satisfy constraints is not a consistent global state.

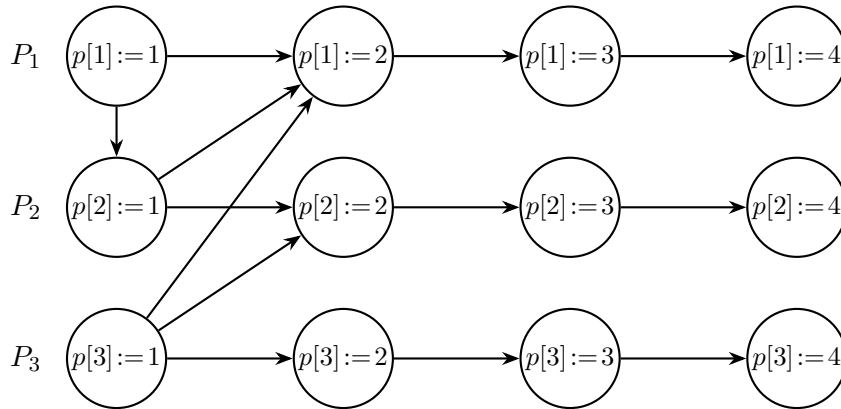


Figure 16.1: The computation graph for a market with three items and three bidders. The valuation and price of any item is a number between 0 and 4. The computation graph models the constraint $B \equiv (p[2] \geq 2 \Rightarrow p[1] \geq 3) \wedge (p[1] \geq 2 \Rightarrow p[3] \geq 1)$

Example 16.3 [Constrained MCP on the computation graph of Fig. 16.1] Take three bidders with valuation vectors $[4, 1, 0]$, $[4, 1, 2]$, $[4, 2, 0]$ and the constraint $B \equiv (p[2] \geq 2 \Rightarrow p[1] \geq 3) \wedge (p[1] \geq 2 \Rightarrow p[3] \geq 1)$ from Fig. 16.1. Starting at $G = [0, 0, 0]$, item 1 is overdemanded so we advance P_1 to $G = [1, 0, 0]$, then again to $G = [2, 0, 0]$. The latter is clearing but violates $(p[1] \geq 2 \Rightarrow p[3] \geq 1)$, so we advance P_3 to $G = [2, 0, 1]$. This satisfies the constraint but item 1 is again overdemanded; advancing P_1 once more gives $G = [3, 0, 1]$, which is the least price vector that is clearing and constraint-feasible. The supporting assignment is $1 \rightarrow b_1, 2 \rightarrow b_3, 3 \rightarrow b_2$.

16.5 Summary

This chapter presented algorithms for the assignment problem, which seeks a maximum weight matching in a bipartite graph of items and bidders. The LLP method yields an ascending-price algorithm that computes the minimum market clearing price vector, generalizing the Gale–Demange–Sotomayor auction mechanism.

Problem/Topic	Algorithm	Time Complexity
Market Clearing Price (unit step)	ConstrainedMarketClearingPrice	$O(n^{3.5} T)$
Market Clearing Price (δ -step)	LLP-Assignment	$O(n^3)$ (strongly polynomial)

Here, n is the number of items (or bidders) and T is the maximum valuation. LLP-Assignment matches the best known strongly polynomial bound for the assignment problem, achieved by the Kuhn–Munkres (Hungarian) and Edmonds–Karp algorithms.

16.6 Problems

1. Show that the set of market clearing price vectors is also closed under the join operation.
2. Give the linear programming formulation and its dual for solving the assignment problem.
3. Suppose that your friend gives you an assignment and claims that the assignment is optimal. It is easy to check that the assignment is proper, i.e., every item is assigned to exactly one bidder. But, how can she quickly convince you of the optimality?
4. Run Algorithm [LLP-Assignment](#) by hand on the 3×3 instance with valuation matrix

$$V = \begin{pmatrix} 5 & 3 & 1 \\ 3 & 5 & 1 \\ 2 & 3 & 4 \end{pmatrix}.$$

Report the sequence of price vectors visited, the final minimum market clearing price, and a supporting perfect matching.

5. Show that there is a unique minimum market clearing price vector. (Hint: use Lemma 16.1.)
6. Suppose the auctioneer imposes a reserve price $r_i \geq 0$ on each item i , meaning that no item may be sold below its reserve. Show how to model this as a constrained market clearing price problem and how Algorithm [ConstrainedMarketClearingPrice](#) should be modified.

7. Suppose all valuations $v_{b,i}$ are increased by the same constant c , giving new valuations $v'_{b,i} = v_{b,i} + c$. How does the optimal assignment change? How does the minimum market clearing price change?

16.7 Bibliographic Remarks

Kuhn's Hungarian method for solving the assignment problem is from [Mun57]. Shapley and Shubik [SS71] introduced the assignment game and connected market-clearing prices to the core of a cooperative game; Demange, Gale, and Sotomayor [DGS86] gave the ascending-price (exact) auction mechanism that the LLP algorithm of this chapter generalizes. Bertsekas's auction algorithm [Ber92] is a closely related ascending-bid scheme with strong empirical performance. The classical Edmonds–Karp paper [EK72] embeds the assignment problem inside minimum-cost flow, which is the natural generalization to non-bipartite supply-and-demand instances and to the transportation problem. The notion of a constrained market clearing price is from [Gar20a]. The monograph by Burkard, Dell'Amico, and Martello [BDM12] is a comprehensive reference for variants of the assignment problem and their algorithms.

Chapter 17

Horn and 2-SAT Satisfiability

17.1 Introduction

The satisfiability problem—given a Boolean formula, determine whether there exists an assignment of truth values to its variables that makes the formula true—is the canonical NP-complete problem. Yet within this vast landscape of intractability lie islands of tractability: special classes of Boolean formulas for which satisfiability can be decided efficiently. Understanding these tractable classes is important for both theory and practice, as many real-world constraint satisfaction problems fall naturally into these classes.

In this chapter, we study two of the most important such classes: *Horn formulas* and *2-SAT formulas*. Horn formulas arise naturally in logic programming, database theory, and artificial intelligence. The Prolog programming language, for instance, is built entirely on Horn clauses: every Prolog rule is a Horn implication, and every Prolog query is answered by finding a satisfying assignment for a conjunction of Horn clauses. In database theory, Horn clauses model integrity constraints and deductive rules.

The 2-SAT problem, where every clause has at most two literals, appears in numerous settings: type checking in programming languages, constraint propagation in planning, and as a subroutine in approximation algorithms for harder optimization problems.

What makes these two classes tractable is fundamentally different. Horn formulas are lattice-linear predicates: the set of satisfying assignments is closed under meet (componentwise AND), and the Lattice Linear Predicate (LLP) framework yields the *least* satisfying assignment. In contrast, 2-SAT formulas are *not* lattice-linear in general, but their structure can be captured by an *implication graph* whose strongly connected components reveal satisfiability in linear time.

Throughout this chapter, we assume that the predicate is given in the *conjunctive normal form* (CNF). A Boolean formula B in conjunctive normal form is simply a conjunction of *clauses*, where each clause is a pure disjunction of *literals*. A literal is a variable in its simple form or in a complemented form. For example, suppose that we have n Boolean variables $\{x_1, x_2, \dots, x_n\}$. Then, the formula $(\neg x_1 \vee \neg x_2 \vee x_3) \wedge (x_1 \vee x_2)$ is in CNF with two clauses. The first clause has three literals $\{\neg x_1, \neg x_2, x_3\}$, and the second clause has two literals $\{x_1, x_2\}$.

The chapter is organized as follows. Section 17.2 defines Horn clauses and presents the LLP-based algorithm for Horn satisfiability, proving lattice-linearity along the way. Section 17.3 gives an alternative proof via arithmetization. Section 17.4 develops the implication-graph approach for 2-SAT. Section 17.5

summarizes the results.

17.2 Horn Satisfiability

A clause is a *Horn clause* if it has at most one positive literal. An example of a Horn clause is $(\neg x_1 \vee \neg x_2 \vee x_3)$. Note that this formula is also equivalent to $(x_1 \wedge x_2) \Rightarrow x_3$. A Horn clause written in this form is also called a *Horn implication* or a *definite clause*. Notice that it has only positive literals on the left-hand side and a single positive literal on the right-hand side. The left-hand side may be empty. Thus, x_3 is also a Horn implication equivalent to $true \Rightarrow x_3$.

Another example of a Horn clause is $(\neg x_1 \vee \neg x_2 \vee \neg x_3)$. This clause does not have any positive literal. It is a *pure negative* clause.

A Horn formula is just a conjunction of Horn clauses. The problem of HornSAT is to determine if the given Horn formula is satisfiable.

Example 17.1 Consider the following Horn formula B with three variables x_1 , x_2 , and x_3 :

$$x_1 \wedge (\neg x_1 \vee x_2) \wedge (\neg x_1 \vee \neg x_2 \vee x_3),$$

or, written as implications, $\top \Rightarrow x_1$, $x_1 \Rightarrow x_2$, and $x_1 \wedge x_2 \Rightarrow x_3$. The unit fact forces $x_1 = \text{true}$; the second clause then forces $x_2 = \text{true}$; the third clause forces $x_3 = \text{true}$. Hence $(1, 1, 1)$ satisfies B , and any satisfying assignment must agree with it on every variable. So $(1, 1, 1)$ is the *least* satisfying assignment in the Boolean lattice of three variables. We will use B as the running example throughout this section.

Example 17.2 The Boolean expression $B = (\neg x_1 \vee x_2) \wedge (\neg x_2 \vee x_3)$ is a Horn formula with two implications: $x_1 \Rightarrow x_2$ and $x_2 \Rightarrow x_3$. Its satisfying assignments are shown in the Boolean lattice of Fig. 17.1. Note that the four satisfying assignments $\{000, 001, 011, 111\}$ form a meet-closed (i.e., closed under componentwise AND) subset of the lattice. The least satisfying assignment is 000.

Example 17.3 Consider the following set of Horn clauses with two variables x_1 and x_2 :

$$(\neg x_1 \vee x_2) \wedge (\neg x_2 \vee x_1) \wedge (\neg x_1 \vee \neg x_2) \wedge x_1$$

This set of Horn clauses is unsatisfiable because there is no assignment of truth values to x_1 and x_2 that can satisfy all the clauses simultaneously. To see why: the unit clause x_1 forces $x_1 = \text{true}$. The first implication $x_1 \Rightarrow x_2$ then forces $x_2 = \text{true}$. But now the pure negative clause $(\neg x_1 \vee \neg x_2)$ is violated.

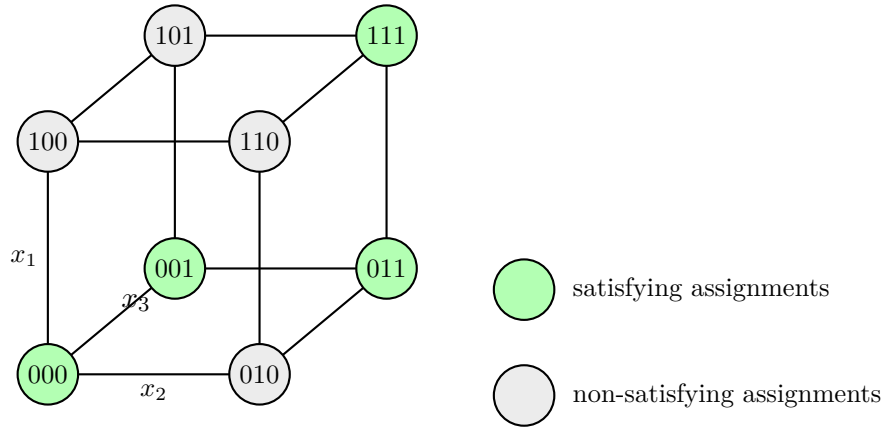


Figure 17.1: Lattice of Boolean assignments on (x_1, x_2, x_3) for the Horn formula $B = (\neg x_1 \vee x_2) \wedge (\neg x_2 \vee x_3)$. The four satisfying assignments form a meet-closed subset $\{000, 001, 011, 111\}$.

Example 17.4 Horn clauses arise naturally in logic programming. Consider a small knowledge base:

<code>parent(alice, bob)</code>	(fact: Alice is Bob’s parent)
<code>parent(bob, carol)</code>	(fact: Bob is Carol’s parent)
<code>parent(X, Y) ∧ parent(Y, Z) ⇒ grandparent(X, Z)</code>	(rule)

The query “Is Alice a grandparent of Carol?” amounts to checking whether `grandparent(alice, carol)` can be derived. This is exactly a Horn satisfiability problem: the facts and rules are Horn clauses, and the query asks whether a certain assignment follows. The Prolog programming language evaluates such queries by forward chaining through the implications.

We now show that Horn formulas are lattice-linear predicates, which enables us to apply the LLP framework to find the least satisfying assignment efficiently.

Lemma 17.5 *Any Horn formula is a lattice-linear predicate.*

Proof: Let $B = B_1 \wedge B_2 \wedge \dots \wedge B_m$ be the Horn formula, where each B_i for $i \in [m]$ is a Horn clause. Since lattice-linear predicates are closed under conjunction, it suffices to show that every Horn clause B_i is lattice-linear. We consider two types of Horn clauses.

First, suppose that B_i is a Horn implication, i.e., $B_i \equiv (x_{i_1} \wedge x_{i_2} \wedge \dots \wedge x_{i_{k-1}} \Rightarrow x_{i_k})$. Consider any Boolean vector G in which B_i is false. This implies that $x_{i_1}, x_{i_2}, \dots, x_{i_{k-1}}$ are true in G but x_{i_k} is false. Consider any H such that $H \geq G$ and $H[i_k] = G[i_k]$. Since $H \geq G$, we have that $(x_{i_1} \wedge x_{i_2} \wedge \dots \wedge x_{i_{k-1}})$ holds in H . Furthermore, since $H[i_k]$ is equal to $G[i_k]$, x_{i_k} is false in H . Hence, B_i is also false in H .

Now suppose that B_i is a pure negative clause, i.e., $B_i \equiv (\neg x_{i_1} \vee \neg x_{i_2} \vee \dots \vee \neg x_{i_k})$. Suppose that B_i is false in G . This implies that all the variables x_{i_1}, \dots, x_{i_k} are true. Consider any Boolean vector $H \geq G$. H must also have all these variables true and, therefore, B_i is false in H (and any index is trivially forbidden).

■

The proof of Lemma 17.5 also shows us the appropriate advancement function. For Horn implications, we must advance on the right-hand-side literal. For pure negative clauses, all Boolean vectors greater than G do not satisfy B . Hence, the algorithm can simply return that “no satisfying vector exists.”

We can now apply the LLP algorithm to find a satisfying assignment for a Horn formula. We assume that the Horn formula uses the variables x_1, \dots, x_n . These n Boolean variables define a Boolean lattice L of size 2^n that consists of Boolean vectors of size n .

Algorithm LLP-HornSAT: An Algorithm to find the least assignment that satisfies B

Input: Horn CNF B with variables x_1, \dots, x_n
Output: G : least satisfying assignment, or null

```

1  $G$ : vector of boolean initially  $\forall i : G[i] = \text{false}$ ;
2 forbidden( $j$ ):  $\exists h$ : all antecedents of  $B_h$  are true, they imply  $x_j$ , and  $x_j$  is false in  $G$ ;
3   advance( $j$ ):  $G[j] := \text{true}$ ;
4 forbidden( $j$ ):  $\exists h$ : a negative clause  $B_h$  is false, and  $x_j$  is a part of  $B_h$ ;
5   advance return null ; // no satisfying assignment exists
6 return  $G$  ; // the least assignment that satisfies  $B$ 
```

We leave an efficient implementation of Algorithm LLP-HornSAT as an exercise.

Observe that we also get the following consequence of Lemma 17.5 from its lattice-linearity.

Corollary 17.6 *If G and H satisfy the Horn formula, then $G \sqcap H$ also satisfies that formula.*

A direct proof of this fact is left as an exercise. Although the set of assignments satisfying a Horn formula is closed under meets, it is not closed under joins. For example, consider the Horn formula $B \equiv (\neg x_1 \vee \neg x_2)$. The bit vectors $G = [1, 0]$ and $H = [0, 1]$ satisfy B but their join $[1, 1]$ does not satisfy B .

Example 17.7 Let us trace LLP-HornSAT on the Horn formula B from Example 17.1, written as the three implications

$$C_1 : \top \Rightarrow x_1, \quad C_2 : x_1 \Rightarrow x_2, \quad C_3 : x_1 \wedge x_2 \Rightarrow x_3.$$

Start with $G = (0, 0, 0)$.

Step 0. C_1 has empty antecedent and consequent x_1 , which is false. So C_1 is violated and index 1 is forbidden. Advance: $G[1] := \text{true}$, giving $G = (1, 0, 0)$.

Step 1. C_1 now holds. C_2 's antecedent x_1 is true but the consequent x_2 is false, so C_2 is violated and index 2 is forbidden. Advance: $G[2] := \text{true}$, giving $G = (1, 1, 0)$.

Step 2. C_1, C_2 hold. C_3 's antecedents $x_1 \wedge x_2$ are both true but the consequent x_3 is false, so C_3 is violated and index 3 is forbidden. Advance: $G[3] := \text{true}$, giving $G = (1, 1, 1)$.

Step 3. All three clauses are satisfied; no index is forbidden. The algorithm returns $G = (1, 1, 1)$ as the least satisfying assignment, agreeing with Example 17.1. The trace illustrates how the LLP advance rule realizes the standard *forward chaining* of Horn solvers: each violated implication fires its consequent, possibly enabling further implications in the next iteration.

Example 17.8 Now augment B with the pure negative clause $C_4 \equiv (\neg x_2 \vee \neg x_3)$:

$$B' = B \wedge (\neg x_2 \vee \neg x_3).$$

The trace proceeds through Steps 0–2 of Example 17.7 unchanged, reaching $G = (1, 1, 1)$. At this state the pure negative clause C_4 becomes false because both x_2 and x_3 are true. The advance rule for negative clauses fires the unsatisfiability return, so LLP-HornSAT returns *null*: B' has no satisfying assignment. This matches the lemma that pure negative clauses, once violated, can never be re-satisfied by further advances.

The LLP-HornSAT algorithm with m clauses and n variables can be implemented in $O(m + n)$ time.

Dual-Horn formulas. A CNF formula is *dual-Horn* if every clause has at most one *negative* literal. Dual-Horn formulas are the bit-complement counterpart to Horn: a formula Φ is dual-Horn iff $\Phi[x \mapsto \neg x]$ (with every literal flipped) is Horn. Consequently, the set of satisfying assignments of a dual-Horn formula is closed under componentwise OR (join) rather than meet, and forms a join-semilattice. The LLP framework applied to the dual Boolean lattice (with $\perp = (1, 1, \dots, 1)$ and the order reversed) yields the *greatest* satisfying assignment in $O(n + m)$ time. The forbidden/advance rules dualize: a clause with one negative literal $\neg x_j$ and antecedents $x_{i_1}, \dots, x_{i_{k-1}}$ all set false forbids index j , advancing $G[j] := \text{false}$. Dual-Horn satisfiability is therefore solved by the same algorithm with the lattice flipped, and the Renamable Horn problem (Exercise) asks when this flip can be applied to selected variables to bring an arbitrary formula into one of these two classes.

17.3 Arithmetization of Horn Clauses

In this section, we give another proof that Horn clauses are lattice-linear by considering arithmetic expressions instead of Boolean expressions. We use the natural assignment of Boolean variables x_i to integer binary variables y_i . If x_i is false, then y_i is assigned a value 0, otherwise it is assigned 1. Given any clause, we translate it into an arithmetic predicate as follows. Every positive literal x_i in any clause is changed to y_i and every negative literal $\neg x_i$ is changed to $(1 - y_i)$. The clause is true iff the sum of all values so replaced in any clause is at least 1. For example, clause $(\neg x_1 \vee \neg x_2 \vee x_3)$ is equivalent to $(1 - y_1) + (1 - y_2) + y_3 \geq 1$. If all y_i take values in the set $\{0, 1\}$, then it is easy to verify that the clause is true for x 's iff the corresponding arithmetic predicate is true for y 's. Let us see how the implication Horn clauses get translated. An implication Horn clause $(x_{i_1} \wedge x_{i_2} \wedge \dots \wedge x_{i_{k-1}} \Rightarrow x_{i_k})$ gets translated into $(1 - y_{i_1}) + (1 - y_{i_2}) + \dots + (1 - y_{i_{k-1}}) + y_{i_k} \geq 1$ which is equivalent to $y_{i_k} \geq y_{i_1} + y_{i_2} + \dots + y_{i_{k-1}} - (k - 2)$. The right-hand side is a monotone function of y ; hence, the predicate is lattice-linear. A pure negative clause $(\neg x_{i_1} \vee \neg x_{i_2} \vee \dots \vee \neg x_{i_k})$ is translated into $(1 - y_{i_1}) + (1 - y_{i_2}) + \dots + (1 - y_{i_k}) \geq 1$. This predicate is equivalent to $k - 1 \geq y_{i_1} + y_{i_2} + \dots + y_{i_k}$ which, once it becomes false, stays false. Hence, predicates for negative clauses are also lattice-linear.

Example 17.9 Let us arithmetize the running Horn formula from Example 17.1,

$$B = x_1 \wedge (\neg x_1 \vee x_2) \wedge (\neg x_1 \vee \neg x_2 \vee x_3).$$

Mapping each Boolean x_i to $y_i \in \{0, 1\}$:

- The unit clause x_1 becomes $y_1 \geq 1$.
- The implication clause $(\neg x_1 \vee x_2)$ becomes $(1 - y_1) + y_2 \geq 1$, i.e., $y_2 \geq y_1$.
- The implication clause $(\neg x_1 \vee \neg x_2 \vee x_3)$ becomes $(1 - y_1) + (1 - y_2) + y_3 \geq 1$, i.e., $y_3 \geq y_1 + y_2 - 1$.

Each lower bound on the consequent is a *monotone* function of the antecedents, so each clause is lattice-linear and the conjunction B is too. The least satisfying integer point is $y = (1, 1, 1)$, matching the trace in Example 17.7.

If we further add the pure negative clause $(\neg x_2 \vee \neg x_3)$ from Example 17.8, it arithmetizes to $(1 - y_2) + (1 - y_3) \geq 1$, equivalent to $y_2 + y_3 \leq 1$. This constraint becomes false at $y = (1, 1, 1)$ and cannot be repaired by raising any y_i further — making the unsatisfiability in Example 17.8 visible as an upper-bound cut on the integer lattice.

17.4 2-SAT

A conjunctive normal form formula is a 2-SAT formula iff every clause has at most two literals. For example, formula $B_1 \equiv x_1 \wedge (x_2 \vee \neg x_3) \wedge (\neg x_1 \vee \neg x_3)$ is a 2-SAT formula.

If we consider the lattice of Boolean vectors, a 2-SAT formula may not be closed under meet. For example, consider the 2-SAT formula $(x_1 \vee x_2)$. The Boolean vectors 10 and 01 satisfy the formula, but their meet 00 does not. Hence, there is no *minimum* satisfying assignment and the simple LLP algorithm is not applicable.

This means we need a fundamentally different approach for 2-SAT. The key insight is that every 2-literal clause $(\ell_1 \vee \ell_2)$ can be read as two implications: $\neg \ell_1 \Rightarrow \ell_2$ and $\neg \ell_2 \Rightarrow \ell_1$. These implications define a directed graph—the *implication graph*—whose structure completely characterizes satisfiability.

We now outline an approach to detect if there is any element in the lattice that satisfies B , a 2-SAT formula. The key idea for the 2-SAT algorithm is to convert the predicate to an implication graph. We assume that every clause has two literals. A unit clause x_i or $\neg x_i$ forces the value of the variable. It can be replaced in B , resulting in a 2-SAT formula with fewer variables. Now, we construct an implication graph $I(B)$ for B as follows. The graph has $2n$ vertices, one for each literal. Every clause $(\ell_i \vee \ell_j)$ in B is true iff $(\neg \ell_i \Rightarrow \ell_j) \wedge (\neg \ell_j \Rightarrow \ell_i)$. Therefore, we add two directed edges to the graph — from $\neg \ell_i$ to ℓ_j and from $\neg \ell_j$ to ℓ_i .

Example 17.10 Consider the following 2-SAT formula with three variables x_1 , x_2 , and x_3 :

$$(\neg x_1 \vee x_2) \wedge (x_1 \vee x_3) \wedge (\neg x_2 \vee x_3)$$

The implication graph of B is shown in Fig. 17.2.

Each clause produces two directed edges:

- $(\neg x_1 \vee x_2)$ gives $x_1 \rightarrow x_2$ and $\neg x_2 \rightarrow \neg x_1$.
- $(x_1 \vee x_3)$ gives $\neg x_1 \rightarrow x_3$ and $\neg x_3 \rightarrow x_1$.
- $(\neg x_2 \vee x_3)$ gives $x_2 \rightarrow x_3$ and $\neg x_3 \rightarrow \neg x_2$.

This formula is satisfiable. For instance, $x_1 = x_2 = x_3 = \text{true}$ works. Note that no variable and its negation are in the same strongly connected component.

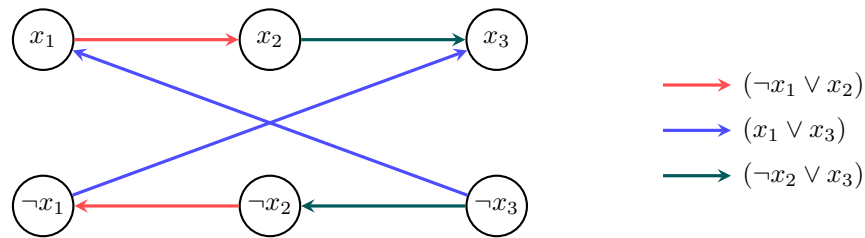


Figure 17.2: Implication graph for $(\neg x_1 \vee x_2) \wedge (x_1 \vee x_3) \wedge (\neg x_2 \vee x_3)$.

An important structural property of the implication graph is its *skew symmetry*: if there is an edge from ℓ to ℓ' , then there is also an edge from $\neg \ell'$ to $\neg \ell$. This follows directly from the construction: both edges come from the same clause. As a consequence, if there is a path from ℓ to ℓ' , there is also a path from $\neg \ell'$ to $\neg \ell$.

The following theorem captures the soundness and completeness of the SCC-based approach. It is the structural fact behind Step 3 of Algorithm 2-SAT.

Theorem 17.11 *A 2-SAT formula B is satisfiable iff there is no variable x_i such that $\neg x_i$ is reachable from x_i and x_i is reachable from $\neg x_i$ in the implication graph $I(B)$ — equivalently, iff no SCC of $I(B)$ contains both x_i and $\neg x_i$.*

Proof: (\Rightarrow) Suppose some variable x_i has both paths in $I(B)$. Pick any assignment σ . Whichever value σ gives x_i , the path from that literal to its negation forces a contradiction along an implication chain that σ must respect (because every edge of $I(B)$ encodes a clause-level implication). Hence no σ satisfies B .

(\Leftarrow) Suppose no SCC contains both x_i and $\neg x_i$. Step 4 of Algorithm 2-SAT constructs a satisfying assignment: process the SCCs in reverse topological order and set every unassigned literal in the current SCC to true. The soundness of this construction is established in the discussion immediately following the algorithm: edges go from earlier to later SCCs in topological order, so when a literal ℓ is set true, every implication $\ell \rightarrow \ell'$ has ℓ' already true. The hypothesis that no SCC contains both x_i and $\neg x_i$ guarantees that we never set both a literal and its negation to true within a single SCC.



We leave the following companion claims as exercises.

- Exercise 17.1** 1. Suppose B is satisfiable. Then, all satisfying assignments set x_j to true iff there exists a path from $\neg x_j$ to x_j in the implication graph.
2. Suppose B is satisfiable. Then, all satisfying assignments set x_j to false iff there exists a path from x_j to $\neg x_j$ in the implication graph.

Algorithm 2-SAT: 2-SAT satisfiability algorithm via SCCs.

Input: A 2-SAT formula Φ with variables x_1, \dots, x_n and m clauses.
Output: UNSATISFIABLE, or SATISFIABLE together with a satisfying assignment.

```

// Step 1: construct the implication graph
1 foreach clause  $(a \vee b)$  in  $\Phi$  do
2   | Add directed edges  $(\neg a \rightarrow b)$  and  $(\neg b \rightarrow a)$ ;
3 end
// Step 2: compute strongly connected components (SCCs)
4 Use Kosaraju's or Tarjan's algorithm to compute SCCs of the graph;
// Step 3: check for contradictions
5 foreach variable  $x_i$  do
6   | if  $x_i$  and  $\neg x_i$  belong to the same SCC then
7     |   return UNSATISFIABLE;
8   | end
9 end
// Step 4: construct an assignment
10 Topologically sort the SCCs of the implication graph (treating each SCC as a single vertex);
11 Mark every literal as unassigned;
12 foreach SCC  $S$  in reverse topological order do
13   | foreach literal  $\ell \in S$  that is unassigned do
14     |   Assign  $\ell := \text{true}$  and  $\neg \ell := \text{false}$ ;
15   | end
16 end
17 return SATISFIABLE and the constructed assignment;

```

Algorithm 2-SAT works as follows.

In the algorithm, we need to check if there exists a path from x_j to $\neg x_j$ and vice versa. One way to accomplish this is to find strongly connected components (SCCs) in the graph. An SCC is a subgraph in which every vertex is reachable from every other vertex in the same SCC. (One can use Tarjan's algorithm or Kosaraju's algorithm to find SCCs.) The expression is satisfiable if and only if for every variable x_j , x_j and $\neg x_j$ are not in the same SCC. If the expression is satisfiable, an assignment is constructed in Step 4 by processing the SCCs in *reverse* topological order: every unassigned literal in the current SCC is set to true (and its negation to false). To see why this is sound, recall that when each SCC is contracted to a single vertex, the resulting directed acyclic graph admits a topological order in which all edges go from earlier to later SCCs. So when we set a literal ℓ to true, every implication edge $\ell \rightarrow \ell'$ leads to an SCC that has already been processed and therefore ℓ' has already been set to true. The check in Step 3 guarantees that

no SCC contains both x_j and $\neg x_j$, so within a single SCC we never set both a literal and its negation to true. The sequential time complexity of the 2-SAT problem is linear in the number of clauses.

Example 17.12 Consider the 2-SAT formula:

$$(x_1 \vee x_2) \wedge (\neg x_1 \vee \neg x_2) \wedge (\neg x_1 \vee x_2) \wedge (x_1 \vee \neg x_2)$$

The implication graph has edges: $\neg x_1 \rightarrow x_2$, $\neg x_2 \rightarrow x_1$ (from clause 1); $x_1 \rightarrow \neg x_2$, $x_2 \rightarrow \neg x_1$ (from clause 2); $x_1 \rightarrow x_2$, $\neg x_2 \rightarrow \neg x_1$ (from clause 3); $\neg x_1 \rightarrow \neg x_2$, $x_2 \rightarrow x_1$ (from clause 4).

From x_1 we can reach $\neg x_1$: $x_1 \rightarrow x_2 \rightarrow \neg x_1$. From $\neg x_1$ we can reach x_1 : $\neg x_1 \rightarrow x_2 \rightarrow x_1$. So x_1 and $\neg x_1$ are in the same SCC. The formula is unsatisfiable.

Example 17.13 Consider the 2-SAT formula:

$$(\neg x_1 \vee x_2) \wedge (\neg x_2 \vee x_3) \wedge (x_1 \vee \neg x_3)$$

The implication graph has edges: $x_1 \rightarrow x_2$, $\neg x_2 \rightarrow \neg x_1$ (from clause 1); $x_2 \rightarrow x_3$, $\neg x_3 \rightarrow \neg x_2$ (from clause 2); $\neg x_1 \rightarrow \neg x_3$, $x_3 \rightarrow x_1$ (from clause 3).

The SCCs are $S_+ = \{x_1, x_2, x_3\}$ and $S_- = \{\neg x_1, \neg x_2, \neg x_3\}$, as shown in Fig. 17.3. No variable shares an SCC with its negation, so the formula is satisfiable. There are no edges between S_+ and S_- in the SCC-contracted graph, so either order is a valid topological sort. Pick the topological order (S_-, S_+) . Processing in *reverse* topological order, we first visit S_+ : all three literals x_1, x_2, x_3 are unassigned, so we set them to true (and $\neg x_1, \neg x_2, \neg x_3$ to false). When we next visit S_- , every literal is already assigned, so nothing changes. The constructed assignment is $x_1 = x_2 = x_3 = \text{true}$. Verification: clause 1 is $(\neg x_1 \vee x_2) = (F \vee T) = T$; clause 2 is $(\neg x_2 \vee x_3) = (F \vee T) = T$; clause 3 is $(x_1 \vee \neg x_3) = (T \vee F) = T$. All clauses satisfied.

17.5 Summary

In this chapter, we explored two important subclasses of Boolean formulas, Horn and 2-SAT, that form the boundary between tractable and intractable satisfiability problems.

Class	Restriction	Sequential Time	Approach
Horn SAT	≤ 1 positive literal/clause	$O(n + m)$	LLP (lattice-linear)
2-SAT	≤ 2 literals/clause	$O(n + m)$	Implication graph + SCC
3-SAT	≤ 3 literals/clause	NP-complete	—

We showed that Horn formulas are *lattice-linear predicates*, and therefore admit a solution through the LLP framework that yields the least satisfying assignment. The meet-closure property (Corollary 17.6) is a direct consequence of lattice-linearity and has important implications: the intersection of any two models of a Horn formula is again a model.

For 2-SAT formulas, we presented the implication-graph construction and demonstrated that satisfiability can be checked through the detection of strongly connected components. This approach yields a linear-time sequential algorithm.

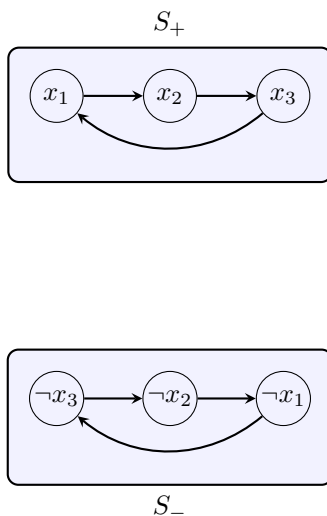


Figure 17.3: The two SCCs of the implication graph for the formula in Example 17.13. There are no edges between S_+ and S_- in the SCC-contracted graph, so either is a valid topological order. The reverse-topological-order rule of Algorithm 2-SAT sets every literal in the first-visited SCC to true.

It is instructive to compare the two approaches. Horn satisfiability exploits the *algebraic* structure of the solution space (it forms a meet-semilattice), while 2-SAT exploits the *graph-theoretic* structure of the implication relationships. The general SAT problem becomes NP-complete precisely when neither structure is present: with 3 or more literals per clause and no restriction on positive literals, neither the lattice-linear approach nor the implication-graph approach applies.

17.6 Problems

1. Is the following Horn SAT formula satisfiable? If so, give the least satisfying assignment.
 $(\neg x_1 \vee \neg x_2 \vee x_3) \wedge (\neg x_3 \vee x_1) \wedge (\neg x_3 \vee \neg x_1) \wedge (\neg x_2 \vee \neg x_3) \wedge (\neg x_1 \vee x_2)$
2. Give an efficient implementation of Algorithm LLP-HornSAT. In particular, give efficient data structures to represent Horn implications and pure negative clauses. Your algorithm should be linear in the length of the Horn formula.
3. A formula is a Renamable Horn formula if, by flipping the polarity of certain variables, the formula can be transformed to a Horn formula. For example, $(\neg x_1 \vee \neg x_2 \vee \neg x_4) \wedge (x_3 \vee x_4) \wedge (x_1 \vee x_2 \vee \neg x_4) \wedge (x_1 \vee \neg x_3)$ is not a Horn formula. However, renaming x_1 as $\neg x_1$ and x_3 as $\neg x_3$, we get a Horn formula. Give an algorithm to determine whether the given formula is a Renamable Horn formula. (Hint: Can you come up with a 2-SAT formula that is true iff the given formula is a renamable Horn formula).
4. Prove Corollary 17.6 without invoking the lattice-linearity of Horn clauses.
5. Consider a directed graph with n nodes that includes two distinguished nodes s and t . Show that there exists a 2-SAT formula B on n variables such that the formula is satisfiable iff t is reachable from s .

6. Suppose a 2-SAT formula B is satisfiable. Then, all satisfying assignments of B must have x_j as true iff there exists a path from $\neg x_j$ to x_j in the implication graph.
7. (a) Show that there exists a satisfying assignment to a 2-SAT formula B iff there is no variable x_i such that $\neg x_i$ is reachable from x_i and x_i is reachable from $\neg x_i$ in the implication graph I constructed from B . (b) Show that every literal in any strongly connected component of the implication graph must be assigned the same value, *true* or *false*.
8. (**Counting satisfying assignments.**) Show that counting the number of satisfying assignments of a 2-SAT formula is #P-complete, even though *deciding* satisfiability is polynomial. (Hint: reduce from #MONOTONE-2-SAT or use a reduction from counting independent sets in a graph.)
9. (**Horn clause and databases.**) A relational database stores facts (e.g., “Alice is Bob’s parent”) and rules (e.g., “if X is a parent of Y and Y is a parent of Z , then X is a grandparent of Z ”). Show that the problem of determining whether a given fact can be derived from a set of ground facts and rules can be modeled as a Horn satisfiability problem.
10. (**Maximum satisfiable Horn clauses.**) Given a Horn formula, the MAX-HORN-SAT problem asks for an assignment that satisfies the maximum number of clauses. Show that this problem is NP-hard. (Hint: reduce from the vertex cover problem.)
11. (**XOR-SAT.**) An XOR-SAT formula is a conjunction of XOR-clauses, where each clause is the XOR (exclusive-or) of literals. For example, $(x_1 \oplus x_2) \wedge (x_2 \oplus \neg x_3)$. Show that XOR-SAT can be solved in polynomial time. (Hint: model each clause as a linear equation over $GF(2)$ and use Gaussian elimination.)

17.7 Bibliographic Remarks

Horn formulas, a subclass of propositional formulas in which each clause has at most one positive literal, are central to efficient satisfiability algorithms. [DG84] present a linear-time algorithm to test the satisfiability of a Horn formula, using unit propagation. The 2-SAT problem, which determines the satisfiability of a CNF formula with at most two literals per clause, is solvable in linear time. [APT79] introduce a linear-time algorithm using implication graphs and strongly connected components.

Horn clauses form the logical foundation of the Prolog programming language and Datalog, a query language for deductive databases [CGT89].

The q-Horn class, which generalizes both Horn and 2-SAT, was introduced by Boros, Crama, and Hammer [BCH90] and is also solvable in polynomial time. Renamable Horn formulas were studied by Lewis [Lew78]. For a comprehensive treatment of the complexity landscape of satisfiability, the reader is referred to the Handbook of Satisfiability [BHvMW09].

Chapter 18

Algorithms in Number Theory

18.1 Introduction

Number theory is one of the oldest branches of mathematics and a rich source of algorithmic problems. Classical questions — computing the greatest common divisor, solving systems of congruences, counting integers coprime to a given number — date back more than two thousand years, yet they remain central in modern computer science. They underlie public-key cryptography (RSA, Diffie–Hellman, elliptic curve cryptography), error-correcting codes, hashing, random number generation, and secret sharing schemes.

In this chapter, we revisit several standard number-theoretic computations through the lens of Lattice-Linear Predicates (LLP). The key observation is that the set of all natural numbers forms a distributive lattice under the *divides* relation: we define $x \leq y$ for two natural numbers x and y if x divides y . In this (infinite) lattice, the bottom element is 1 since it divides all numbers. The join of any two elements is given by their *least common multiple*, and the meet of any two elements is given by their *greatest common divisor*. This lattice-theoretic viewpoint lets us re-derive classical algorithms as instances of the generic LLP framework, and it also suggests natural parallel and distributed implementations.

We proceed as follows. We first present two lattice-linear formulations of the GCD problem, one that reduces to Euclid’s classical algorithm and one that searches in the opposite direction, followed by the extended Euclidean algorithm for computing Bézout coefficients. We then apply LLP to the Chinese Remainder Theorem, obtaining both an ascending and a descending algorithm, and show that the same approach handles the dual problem (least common multiple), non-linear periodic congruences such as simultaneous quadratic residues, and the computation of multiplicative orders. Finally, we study several multiplicative arithmetical functions — the Möbius function μ , Euler’s totient ϕ , and a few related functions — and prove some classical summation identities by viewing numbers as elements of a poset of prime powers.

Section 18.2 introduces the first GCD algorithm, LLP-GCD-Descend, which is a descending LLP algorithm that starts at $G = A$ and repeatedly reduces entries using Euclidean mod operations; we analyze its sequential complexity and give a worked example. Section 18.3 presents the complementary ascending algorithm LLP-GCD-Ascend, which starts at $G = [1, \dots, 1]$ and grows each entry using ceiling operations, followed by its complexity analysis and a comparison showing inputs on which each of LLP-GCD-Descend and LLP-GCD-Ascend is preferable. Section 18.3 extends LLP-GCD-Descend into the extended Euclidean algorithm, which additionally produces Bézout coefficients (x, y) such that $ax + by = \gcd(a, b)$. Section 18.4

applies the LLP framework to the Chinese Remainder Theorem, giving two algorithms (ascending LLP-CRT and descending LLP-CRT-Max) together with their complexity analyses. Section 18.5 presents an ascending LLP algorithm for the least common multiple, dual to LLP-GCD-Descend. Section 18.6 shows that LLP-CRT generalizes beyond linear congruences to systems of non-linear periodic predicates, illustrated with simultaneous quadratic congruences. Section 18.7 develops an LLP algorithm for computing the multiplicative order of an integer modulo n , using a descending search on the divisor lattice of $\phi(n)$. The chapter closes with a *summary of algorithms* table in Section 18.8, a set of problems, and bibliographic remarks.

18.2 Greatest Common Divisor (GCD) Algorithm

Let A be an array of n natural numbers. The set of all common divisors of A forms a distributive lattice. This lattice is never empty because 1 divides all numbers. Our goal is to find the greatest common divisor of these numbers. Instead of finding just one number, we find G , an array of numbers of size n such that each number in G divides all the numbers in A . When G is the greatest such vector, then we have the required GCD.

We assume that all numbers are greater than or equal to 1. We start with $G[i]$ equal to $A[i]$ for all i . The LLP Algorithm will reduce the values in G until we reach the GCD. The top element of the lattice for the LLP algorithm would be the vector $[1, 1, \dots, 1]$. The algorithm terminates with all numbers identical and equal to the GCD.

The predicate is

$$B \equiv (\forall i : \text{gcd}(A) \text{ divides } G[i]) \wedge (\forall i, j : G[i] \geq G[j])$$

Observe that the second conjunct $(\forall i, j : G[i] \geq G[j])$ is equivalent to requiring that all entries of G are equal, which together with the first conjunct forces $G[i] = \text{gcd}(A)$ for every i .

We first need to show that B is lattice-linear.

Lemma 18.1 B is lattice-linear.

Proof: Suppose that B is false. Since the first part is invariant of the program, there exist i and j such that $G[j]$ is strictly greater than $G[i]$. Since G can only decrease, the predicate can never become true unless $G[j]$ is advanced, i.e., decreased. This decrease must be such that the invariant is maintained. ■

Suppose that $G[j]$ is greater than $G[i]$ for some i and $G[i]$ does not divide $G[j]$; then we claim that it is safe to reduce $G[j]$ to $G[j] \bmod G[i]$. This step gives us an algorithm. If all numbers are the same, we stop. Otherwise, if $G[j] > G[i]$, we replace $G[j]$ by $G[i]$ if $G[i]$ divides $G[j]$ and by $G[j] \bmod G[i]$ otherwise.

Algorithm LLP-GCD-Descend: Finding the greatest common divisor of a set of numbers

Input: A : array[1.. n] of int

Output: G : array[1.. n] of int with all entries equal to $\text{gcd}(A)$

1 G : array[1.. n] of int initially $\forall i : G[i] = A[i]$;

2 **forbidden**(j): $\exists i : G[j] > G[i]$;

3 **advance**(j): if $G[i]$ divides $G[j]$ then $G[j] := G[i]$; else $G[j] := G[j] \bmod G[i]$;

The invariant is that every common divisor of A is also a common divisor of G . The invariant holds initially because G is set to A . After every advance, we only need to consider two cases. If $G[j]$ is greater than $G[i]$ and $G[i]$ divides $G[j]$, then it is sufficient to set $G[j]$ to $G[i]$ because the divisors of $G[i]$ are common divisors for both $G[i]$ and $G[j]$. If $G[i]$ does not divide $G[j]$, then the only common divisors are those that divide $G[i]$ and $G[j] \bmod G[i]$ (prove it!).

It is easy to see that the algorithm terminates. All values are initially non-zero positive integers. They stay non-zero and integral and always decrease; therefore, the algorithm must terminate. Initially, G is the same as the given vector A . It decreases as the algorithm executes, and the least possible value is the vector with all 1's.

Example 18.2 Let $A = [12, 18, 30]$, so that $\gcd(A) = 6$. Algorithm LLP-GCD-Descend starts with $G = [12, 18, 30]$. At each step, we pick some j with $G[j] > G[i]$ for some i and advance it.

- Pick $j = 3, i = 1$: $G[3] = 30 > G[1] = 12$, and 12 does not divide 30, so $G[3] := 30 \bmod 12 = 6$. Now $G = [12, 18, 6]$.
- Pick $j = 2, i = 3$: $G[2] = 18 > G[3] = 6$, and 6 divides 18, so $G[2] := 6$. Now $G = [12, 6, 6]$.
- Pick $j = 1, i = 2$: $G[1] = 12 > G[2] = 6$, and 6 divides 12, so $G[1] := 6$. Now $G = [6, 6, 6]$.

No index is forbidden, so the algorithm terminates with $G = [6, 6, 6]$, as expected. Note that LLP-GCD-Descend interleaves Euclid-style reductions across all pairs of array entries, which allows different $G[j]$'s to be advanced in parallel whenever their advance steps are independent.

Let $M = \max_i A[i]$ and let $d = \gcd(A)$. We analyze the algorithm in terms of n , M , and d . Each advance step reduces the value of some $G[j]$ via a Euclidean mod operation. By the standard analysis of Euclid's algorithm, the sequence of reductions that affect any single entry $G[j]$ is of length $O(\log M)$ before that entry reaches d . Hence the total number of advance steps is $O(n \log M)$. Each advance has to identify an index i with $G[j] > G[i]$ and perform one mod operation, which costs $O(n)$ (a scan over the array) plus $O(1)$ arithmetic operations on integers of bit length $O(\log M)$. Therefore the sequential running time of LLP-GCD-Descend is

$$T_{\text{seq}}(\text{LLP-GCD-Descend}) = O(n^2 \log M).$$

If we charge each arithmetic operation its actual bit cost, the running time becomes $O(n^2 (\log M)^2)$.

An alternative GCD algorithm

We now give an alternative GCD algorithm based on lattice-linearity. When all elements of A are divided by the GCD, we get a quotient vector that is also a vector of natural numbers. Finding the GCD of A is equivalent to finding the minimum vector G such that there exists a number d such that for all i , $A[i] = d * G[i]$. The problem can be formulated as finding the minimum G such that $\forall i : G[i] = A[i]/d$. Equivalently, our goal is to find the minimum G such that $B_{\gcd}(G) \equiv \forall i, j : A[i]/G[i] = A[j]/G[j]$. This feasibility predicate, B_{\gcd} , is equivalent to $\forall j : G[j] \geq \max_i \{A[j]/A[i] * G[i]\}$. Since the right-hand side is a monotone function, we know that the predicate is lattice-linear. We can apply the *LLP* algorithm with *forbidden*(G, j) as $G[j] < \max_i \{A[j]/A[i] * G[i]\}$ and $\alpha(G, j) = \lceil \max_i \{A[j]/A[i] * G[i]\} \rceil$.

We now give an algorithm to compute the gcd of an array of numbers based on this idea.

Algorithm LLP-GCD-Ascend: Finding the greatest common divisor of a set of numbers

Input: A : array[1.. n] of int

Output: $\gcd(A) = A[1]/G[1]$

1 G : array[1.. n] of int initially $\forall i : G[i] = 1$;

2 **ensure**(j): $G[j] \geq \max_i \{A[j]/A[i] * G[i]\}$;

3 **return** $A[1]/G[1]$;

The above algorithm was mechanically derived from the LLP algorithm. Let us see how it relates to Euclid's algorithm to compute the GCD of two numbers: $A[1]$ and $A[2]$. Suppose that $A[1]$ and $A[2]$ are 10 and 15, respectively. We get the following steps. Initially, $G[1] = 1$ and $G[2] = 1$. Since $G[2] * A[1] < G[1] * A[2]$ (i.e., $10 < 15$), we update $G[2]$ to $\lceil (A[2]/A[1]) * G[1] \rceil = \lceil 15/10 \rceil = 2$, getting $G = [1, 2]$. Now, $G[1] * A[2] < G[2] * A[1]$ (i.e., $15 < 20$), so we update $G[1]$ to $\lceil (A[1]/A[2]) * G[2] \rceil = \lceil 20/15 \rceil = 2$, getting $G = [2, 2]$. Now $G[2] * A[1] < G[1] * A[2]$ (i.e., $20 < 30$), so we update $G[2]$ to $\lceil (A[2]/A[1]) * G[1] \rceil = \lceil 30/10 \rceil = 3$. We now have $G = [2, 3]$, and the while condition $A[1]/G[1] = 10/2 = 5$ equals $A[2]/G[2] = 15/3 = 5$, so the algorithm terminates. It returns $A[1]/G[1]$, which equals 5, the gcd of 10 and 15.

Example 18.3 Let us apply LLP-GCD-Ascend to $A = [12, 18, 30]$. Initially $G = [1, 1, 1]$. At each step, we compute $\max_i \{A[j]/A[i] * G[i]\}$ for each j and advance whichever entries are below their max. A possible execution is:

- For $j = 2$: $\max\{18/12 \cdot 1, 18/18 \cdot 1, 18/30 \cdot 1\} = 3/2$, so $G[2] := \lceil 3/2 \rceil = 2$. Now $G = [1, 2, 1]$.
- For $j = 3$: $\max\{30/12 \cdot 1, 30/18 \cdot 2, 30/30 \cdot 1\} = 10/3$, so $G[3] := \lceil 10/3 \rceil = 4$. Now $G = [1, 2, 4]$.
- Updates continue until G stabilizes at $[2, 3, 5]$, which satisfies $A[i]/G[i] = 6$ for every i .

At termination, the algorithm returns $A[1]/G[1] = 12/2 = 6 = \gcd(A)$.

Let $M = \max_i A[i]$, $d = \gcd(A)$, and let $Q = M/d$ denote the maximum quotient. The final value of each $G[j]$ in LLP-GCD-Ascend is $A[j]/d$, which is at most Q ; in particular, all intermediate values of G remain bounded by Q . Each advance of $G[j]$ at least increases $G[j]$ to $\lceil \max_i \{A[j]/A[i] \cdot G[i]\} \rceil$, and this value grows geometrically along any chain of advances of a fixed entry. Consequently, the number of advances per entry is $O(\log Q)$, and the total number of advance steps is $O(n \log Q)$. Each advance computes a maximum over n ratios, costing $O(n)$. Hence

$$T_{\text{seq}}(\text{LLP-GCD-Ascend}) = O(n^2 \log Q) = O(n^2 \log(M/d)).$$

Crucially, the arithmetic in LLP-GCD-Ascend is performed on the small integers that appear in G (bounded by Q), whereas LLP-GCD-Descend performs its mod operations on the original large numbers in A . For inputs with a large common factor, this makes LLP-GCD-Ascend's per-step cost markedly cheaper in bit complexity.

When is LLP-GCD-Descend faster, and when is LLP-GCD-Ascend faster?

The two algorithms have dual behaviors: LLP-GCD-Descend decreases a vector from A down to $d \cdot \mathbf{1}$, while LLP-GCD-Ascend increases a vector from $\mathbf{1}$ up to A/d . Each is preferable on different kinds of inputs.

Example 18.4 Consider two almost-equal, almost-coprime numbers $A = [1000001, 1000000]$. Here $d = 1$ and $Q = 10^6$. LLP-GCD-Descend finishes in two advance steps:

- $G[1] = 10^6 + 1 > G[2] = 10^6$; 10^6 does not divide $10^6 + 1$, so $G[1] := (10^6 + 1) \bmod 10^6 = 1$. Now $G = [1, 10^6]$.
- $G[2] > G[1]$ and 1 divides 10^6 , so $G[2] := 1$. Now $G = [1, 1]$ and the algorithm terminates.

In contrast, LLP-GCD-Ascend must grow G from $[1, 1]$ to $[10^6 + 1, 10^6]$, which takes on the order of $\log Q \approx 20$ advances in the best case, but with each ceiling update yielding only a very small increment because $A[1]/A[2] \approx 1$ and $A[2]/A[1] \approx 1$; in practice LLP-GCD-Ascend crawls forward by 1 at a time, requiring roughly 10^6 advance steps. LLP-GCD-Descend is better on inputs whose Euclidean remainder is small — these are exactly the inputs for which Euclid’s original algorithm was designed.

Example 18.5 Consider $A = [3 \cdot 2^{100}, 5 \cdot 2^{100}]$. Here $d = 2^{100}$ and the quotient vector is $[3, 5]$, so $Q = 5$. LLP-GCD-Descend operates on the huge numbers in A : each Euclidean mod involves integers of bit length ≈ 100 , and even though LLP-GCD-Descend only takes 3 advance steps on this pair, each step costs $\Theta(100^2)$ bit operations to compute the mod, for a total of roughly $3 \cdot 10^4$ bit operations. LLP-GCD-Ascend takes about 5 advance steps to grow G from $[1, 1]$ to $[3, 5]$, but each step is $O(1)$ bit operations because the entries of G are all single-digit integers. LLP-GCD-Ascend thus finishes in a constant number of bit operations, several orders of magnitude cheaper than LLP-GCD-Descend. More generally, LLP-GCD-Ascend is preferable whenever the numbers in A are very large but their mutual greatest common divisor is also large, so that the quotient vector is tiny.

18.3 Extended GCD Algorithm

The *extended Euclidean algorithm* strengthens LLP-GCD-Descend by computing, in addition to the greatest common divisor d of two numbers a and b , integer coefficients x and y such that

$$ax + by = d = \gcd(a, b).$$

Such a pair (x, y) is called a Bézout pair for (a, b) . The existence of Bézout pairs is guaranteed by Bézout’s identity and is essential for many applications, including: computing multiplicative inverses modulo m (needed in the Chinese Remainder Theorem and in RSA key generation), solving linear Diophantine equations of the form $ax + by = c$, and lifting relations in quotient rings.

We cast the extended algorithm as a lattice-linear one with an auxiliary array H of Bézout coefficients. We maintain the invariant

$$I \equiv \forall j : G[j] = H[j][1] \cdot a + H[j][2] \cdot b.$$

Initially, $G[1] = a$ and $G[2] = b$, and H is the 2×2 identity matrix, so the invariant holds. Whenever LLP-GCD-Descend would replace $G[j]$ by $G[j] - q \cdot G[i]$ for some integer quotient q , we perform the same subtraction on the Bézout row $H[j]$:

$$G[j] \leftarrow G[j] - q \cdot G[i], \quad H[j] \leftarrow H[j] - q \cdot H[i].$$

A direct computation shows that the invariant I is preserved: the new $G[j]$ equals

$$(H[j][1] - q H[i][1]) a + (H[j][2] - q H[i][2]) b,$$

which matches the new $H[j]$. Thus when LLP-GCD-Descend terminates with $G[1] = G[2] = d$, the first row of H gives a Bézout pair $(x, y) = (H[1][1], H[1][2])$ satisfying $ax + by = d$.

To make the reduction fit a uniform subtractive form, we set the quotient to $q = \lfloor G[j]/G[i] \rfloor$ when $G[i]$ does not divide $G[j]$, and $q = G[j]/G[i] - 1$ when $G[i]$ divides $G[j]$; in the latter case the new $G[j]$ equals $G[i]$, matching LLP-GCD-Descend's update rule. We obtain the following algorithm:

Algorithm LLP-ExtGCD: Extended Euclidean algorithm

Input: a, b : natural numbers

Output: (d, x, y) such that $d = \gcd(a, b)$ and $ax + by = d$

1 G : array[1..2] of int, initially $G := [a, b]$;

2 H : array[1..2][1..2] of int, initially $H := \begin{pmatrix} 1 & 0 \\ 0 & 1 \end{pmatrix}$;

3 **forbidden**(j): $\exists i : G[j] > G[i]$;

4 **advance**(j): (*pick such an i*) if $G[i]$ divides $G[j]$ then $q := G[j]/G[i] - 1$; else $q := \lfloor G[j]/G[i] \rfloor$;
 $G[j] := G[j] - q \cdot G[i]$; $H[j][1] := H[j][1] - q \cdot H[i][1]$; $H[j][2] := H[j][2] - q \cdot H[i][2]$;

5 **return** $(G[1], H[1][1], H[1][2])$;

Example 18.6 Let $a = 30$ and $b = 18$. Initially $G = [30, 18]$ and $H = \begin{pmatrix} 1 & 0 \\ 0 & 1 \end{pmatrix}$.

- $G[1] = 30 > G[2] = 18$, and 18 does not divide 30, so $q = \lfloor 30/18 \rfloor = 1$, and $G[1] := 30 - 18 = 12$, $H[1] := (1, 0) - 1 \cdot (0, 1) = (1, -1)$. Check: $12 = 1 \cdot 30 + (-1) \cdot 18$. Now $G = [12, 18]$, $H = \begin{pmatrix} 1 & -1 \\ 0 & 1 \end{pmatrix}$.
- $G[2] = 18 > G[1] = 12$, 12 does not divide 18, so $q = 1$, $G[2] := 18 - 12 = 6$, $H[2] := (0, 1) - 1 \cdot (1, -1) = (-1, 2)$. Check: $6 = (-1) \cdot 30 + 2 \cdot 18$. Now $G = [12, 6]$, $H = \begin{pmatrix} 1 & -1 \\ -1 & 2 \end{pmatrix}$.
- $G[1] = 12 > G[2] = 6$, and 6 divides 12, so $q = 12/6 - 1 = 1$, $G[1] := 12 - 6 = 6$, $H[1] := (1, -1) - 1 \cdot (-1, 2) = (2, -3)$. Check: $6 = 2 \cdot 30 + (-3) \cdot 18 = 60 - 54$. Now $G = [6, 6]$, and the algorithm terminates.

We return $(6, 2, -3)$: $\gcd(30, 18) = 6$, with Bézout coefficients $(x, y) = (2, -3)$.

For two input numbers a, b with $M = \max(a, b)$, the number of advance steps is the same as in classical Euclid, namely $O(\log M)$. Each advance performs a constant number of arithmetic operations on integers bounded by M in the G array and by M in the H array (the Bézout coefficients produced by extended Euclid are bounded in absolute value by the larger input). Sequential time is therefore $O(\log M)$ arithmetic operations, or $O((\log M)^2)$ bit operations. In parallel, the algorithm has no useful parallelism beyond the two indices and runs in the same $O(\log M)$ depth.

For $n > 2$ inputs, one can either iterate LLP-ExtGCD on pairs (producing an n -entry Bézout decomposition in $O(n \log M)$ arithmetic operations), or run the full n -entry version of the algorithm with an $n \times n$ matrix H of coefficients, at sequential cost $O(n^2 \log M)$ arithmetic operations.

18.4 Chinese Remainder Theorem

The Chinese Remainder Theorem (CRT) states that if m_1, m_2, \dots, m_r are pairwise coprime positive integers and b_1, \dots, b_r are any integers with $0 \leq b_i < m_i$, then the system of congruences $x \equiv b_i \pmod{m_i}$ has a unique solution modulo $M = m_1 m_2 \cdots m_r$. The theorem is more than two thousand years old — it appears in Sun Tzu's *Sun Zi Suan Jing* (3rd century CE) — and it is one of the most useful structural results in elementary number theory. It underpins algorithms for modular arithmetic with large integers, secret sharing schemes, and efficient implementations of polynomial arithmetic.

We have r coprime numbers m_1, m_2, \dots, m_r and r integers b_i such that $0 \leq b_i < m_i$. Our goal is to find the least number x such that $x \equiv b_i \pmod{m_i}$. There are r equations. We think of this as r processes such that process i is solving the equation i . Furthermore, we want each process to get the same value of x . Initially, we start with $G[i]$ equal to $b[i]$ for all i . We are searching for the least G such that

$$B \equiv (\forall i, j : G[i] \leq G[j]) \wedge (\forall i : G[i] \equiv b_i \pmod{m_i})$$

Since each process P_i can only increase the value of $G[i]$, the predicate $(G[i] \leq G[j])$ is lattice-linear. Furthermore, $(G[i] \equiv b_i \pmod{m_i})$ is a local predicate and therefore also lattice-linear. Thus, B is a lattice-linear predicate, and we can use the LLP algorithm to solve the problem. The simplest algorithm will advance on the process P_i by 1 if either $G[i] < G[j]$ for some j or if $\neg(G[i] \equiv b_i \pmod{m_i})$. Furthermore, the only way the second part would become true is if $G[i]$ is advanced to a state in which $G[i] \pmod{m_i}$ equals b_i . To simplify matters further, we initialize $G[i]$ to b_i since that is the smallest state in P_i satisfying the local predicate. We maintain the invariant that $\forall i : G[i] \equiv b[i] \pmod{m[i]}$. With this observation, we get the Figure [LLP-CRT](#).

Algorithm LLP-CRT: A Parallel Algorithm for Chinese Remainder Problem

Input: m : array[1.. n] of int; b : array[1.. n] of int

Output: least x such that $\forall j : x \equiv b[j] \pmod{m[j]}$

1 G : array[1.. n] of int initially $\forall j : G[j] := b[j]$;

2 **forbidden**(j): $\exists i : G[j] < G[i]$;

3 **advance**(j): let $\Delta := \max_i G[i] - G[j]$; $G[j] := G[j] + \lceil \Delta / m[j] \rceil \cdot m[j]$; // smallest $x \geq \max_i G[i]$ with $x \equiv b[j] \pmod{m[j]}$

4 ;

Observe that process P_i does not need access to any $b[j]$ or $m[j]$ for $j \neq i$. It needs access to $G[j]$. Hence, this algorithm can be applied in a distributed fashion if G is publicly available even though b and

m are private. Applying the Chinese Remainder Theorem to solve this problem requires public access to m .

So far we have not shown that the algorithm terminates, but we know that if the algorithm terminates it gives us the smallest number that satisfies B . For this particular example, we do know that the program terminates due to Chinese Remainder Theorem. However, if instead of linear equations we had arbitrary equations, then one could still apply this program, except that the program may not terminate.

Example 18.7 Consider the classical example $x \equiv 2 \pmod{3}$, $x \equiv 3 \pmod{5}$, $x \equiv 2 \pmod{7}$. We have $m = [3, 5, 7]$ and $b = [2, 3, 2]$, so initially $G = [2, 3, 2]$. We repeatedly advance any j for which $G[j] < \max_i G[i]$ to the next value that is congruent to $b[j]$ modulo $m[j]$:

- $G[1] = 2$ is the smallest, and $\max_i G[i] = 3$, so advance $G[1]$ to the next number $\equiv 2 \pmod{3}$ that is ≥ 3 ; that is $G[1] := 5$. Now $G = [5, 3, 2]$.
- Advance $G[2]$ to the next number $\equiv 3 \pmod{5}$ that is ≥ 5 ; that is $G[2] := 8$. Advance $G[3]$ to the next number $\equiv 2 \pmod{7}$ that is ≥ 5 ; that is $G[3] := 9$. Now $G = [5, 8, 9]$.
- Continuing this way, the entries eventually converge to $G = [23, 23, 23]$, and the algorithm returns 23. Indeed, $23 \equiv 2 \pmod{3}$, $23 \equiv 3 \pmod{5}$, and $23 \equiv 2 \pmod{7}$, matching the classical CRT answer.

Instead of searching for the solution in increasing order, one can also search for it in decreasing order provided we know where to start the search from. In the above example, let M be the product of all $m[i]$, i.e., $M = m[1]m[2]\dots m[n]$. Then, from the Chinese Remainder Theorem, we know that the solution exists between $0..M - 1$. It is easy to see that B is also dual lattice-linear. Hence, one can use the following algorithm to find the largest solution that is at most M .

Algorithm LLP-CRT-Max: A Parallel Algorithm for Chinese Remainder Problem

Input: m : array[1.. n] of int; b : array[1.. n] of int

Output: largest $x < M$ such that $\forall j : x \equiv b[j] \pmod{m[j]}$

- 1 G : array[1.. n] of int initially $\forall j : G[j] :=$ greatest number at most $M - 1$ that is congruent to $b[j] \pmod{m[j]}$;
 - 2 **forbidden**(j): $\exists i : G[j] > G[i]$;
 - 3 **advance**(j): let $\Delta := G[j] - \min_i G[i]$; $G[j] := G[j] - \lceil \Delta/m[j] \rceil \cdot m[j]$; // largest $x \leq \min_i G[i]$ with $x \equiv b[j] \pmod{m[j]}$
 - 4 ;
-

Computational complexity of LLP-CRT and LLP-CRT-Max

Let $M = m_1 m_2 \dots m_r$ and let x^* denote the unique solution of the CRT system in $\{0, 1, \dots, M - 1\}$. By construction, the final value of every $G[j]$ in LLP-CRT equals x^* . The number of advance steps performed by process P_j is at most $\lfloor (x^* - b_j)/m_j \rfloor$, which is at most M/m_j . Each advance in LLP-CRT consists of a

scan over the other entries to compute $\max_i G[i]$ and one arithmetic step, costing $O(r)$. Hence:

$$T_{\text{seq}}(\text{LLP-CRT}) = O\left(r \sum_{j=1}^r \frac{M}{m_j}\right) = O\left(r M \sum_{j=1}^r \frac{1}{m_j}\right).$$

LLP-CRT-Max is symmetric: its sequential complexities have the same form, except that each process starts from the largest admissible value below M and descends toward x^* . The number of advance steps is bounded by $\lfloor (M - 1 - x^*)/m_j \rfloor + 1 \leq M/m_j$ per process.

These bounds are polynomial in the *numeric values* M and m_j but only pseudopolynomial in the input size, which is the total number of bits needed to write the moduli. For applications with small moduli (e.g., residue number systems in signal processing or small secret-sharing thresholds), this is perfectly adequate; for large moduli one should instead use the constructive CRT solution based on the extended Euclidean algorithm of Section 18.3, which runs in polynomial time in the bit size of the input.

A polynomial-time CRT via LLP-ExtGCD

The pseudopolynomial bound of LLP-CRT and LLP-CRT-Max is unavoidable for those algorithms: each process advances $G[j]$ by jumps of m_j , so the number of advances is proportional to the magnitude of the answer. When the moduli are large — as in cryptographic applications — we need a genuinely polynomial-time procedure. The classical constructive CRT delivers this in $O(r^2)$ arithmetic operations on integers bounded by M , equivalently $O(r^2 \log^2 M)$ bit operations using schoolbook multiplication.

The construction uses the LLP-ExtGCD of Section 18.3 as a subroutine. For each j , let $M_j = M/m_j$. Since the m_j are pairwise coprime, $\gcd(m_j, M_j) = 1$, and LLP-ExtGCD on (m_j, M_j) returns coefficients (s_j, y_j) with $s_j m_j + y_j M_j = 1$. Reducing modulo m_j gives $y_j M_j \equiv 1 \pmod{m_j}$, so y_j is the multiplicative inverse of M_j modulo m_j . The combination

$$x = \left(\sum_{j=1}^r b_j M_j y_j \right) \pmod{M}$$

satisfies $x \equiv b_j \pmod{m_j}$ for every j : in the sum, the j -th term reduces to b_j modulo m_j (since $M_j y_j \equiv 1$), and every other term contains the factor $M_k = M/m_k$ for some $k \neq j$, which is divisible by m_j and so vanishes mod m_j .

Algorithm CRT-via-ExtGCD: Polynomial-time CRT via LLP-ExtGCD

Input: m : array[1.. r] of pairwise-coprime int; b : array[1.. r] of int with $0 \leq b[j] < m[j]$

Output: the unique $x \in \{0, 1, \dots, M - 1\}$ with $\forall j : x \equiv b[j] \pmod{m[j]}$, where $M = \prod_j m[j]$

```

1  $M := \prod_{j=1}^r m[j]$ ;
2 forall  $j \in [1..r]$  in parallel do
3    $M_j := M/m[j]$ ;
4    $(s_j, y_j) := \text{LLP-ExtGCD}(m[j], M_j)$ ;           // satisfies  $s_j m[j] + y_j M_j = 1$ 
5 end
6  $x := \left( \sum_{j=1}^r b[j] \cdot M_j \cdot y_j \right) \pmod{M}$ ;
7 return  $x$ ;
```

Each LLP-ExtGCD call costs $O(\log M)$ arithmetic operations (or $O(\log^2 M)$ bit operations with schoolbook multiplication); the final summation does r multiplications and additions on integers of bit length $O(\log M)$. Total sequential cost is $O(r \log^2 M)$ bit operations. The r LLP-ExtGCD calls are independent and may all run in parallel, yielding parallel time $O(\log M)$ arithmetic operations on r processors.

Example 18.8 For the system $x \equiv 2 \pmod{3}$, $x \equiv 3 \pmod{5}$, $x \equiv 2 \pmod{7}$, we have $M = 105$, and $M_1 = 35$, $M_2 = 21$, $M_3 = 15$. LLP-ExtGCD gives $35 \cdot 2 \equiv 1 \pmod{3}$, so $y_1 = 2$; $21 \cdot 1 \equiv 1 \pmod{5}$, so $y_2 = 1$; $15 \cdot 1 \equiv 1 \pmod{7}$, so $y_3 = 1$. Hence

$$x = (2 \cdot 35 \cdot 2 + 3 \cdot 21 \cdot 1 + 2 \cdot 15 \cdot 1) \bmod 105 = (140 + 63 + 30) \bmod 105 = 233 \bmod 105 = 23,$$

matching the value 23 found by LLP-CRT in the previous example.

18.5 Least Common Multiple (LCM) Algorithm

Since gcd and lcm are the meet and join of the divisibility lattice, it is natural that the LCM problem admits an LLP algorithm that is the exact dual of LLP-GCD-Descend. Given an array A of n natural numbers, we wish to find their least common multiple. We maintain a vector G of n numbers with the invariant that each $A[i]$ divides $G[i]$ throughout, and we search for the least such vector in which all entries are equal.

The feasibility predicate is

$$B_{\text{lcm}} \equiv (\forall i : A[i] \mid G[i]) \wedge (\forall i, j : G[i] \leq G[j]).$$

The second conjunct forces all entries of G to be equal, and the first forces this common value to be a common multiple of every $A[i]$; by minimality of the LLP solution, the common value is the least common multiple.

We initialize $G[i] = A[i]$ for all i . The invariant $A[i] \mid G[i]$ is satisfied initially and is preserved by advances that only increase $G[i]$ to the next multiple of $A[i]$. The predicate is lattice-linear (ascending): if $G[j] < G[i]$ for some i , then $G[j]$ must increase.

Algorithm LLP-LCM: Finding the least common multiple of a set of numbers

Input: A : array[1.. n] of int

Output: lcm(A)

1 G : array[1.. n] of int initially $\forall i : G[i] = A[i]$;

2 **forbidden**(j): $\exists i : G[j] < G[i]$;

3 **advance**(j): $G[j] := G[j] + A[j] \cdot \lceil (G[i] - G[j])/A[j] \rceil$; // next multiple of $A[j]$ that is $\geq G[i]$

Example 18.9 Let $A = [4, 6, 10]$. We have $\text{lcm}(A) = 60$. The algorithm starts with $G = [4, 6, 10]$.

- $G[1] = 4 < G[3] = 10$: advance $G[1]$ to the next multiple of 4 that is ≥ 10 , i.e., $G[1] := 12$. Now $G = [12, 6, 10]$.
- $G[2] = 6 < G[1] = 12$: advance $G[2]$ to the next multiple of 6 that is ≥ 12 , i.e., $G[2] := 12$. Also $G[3] = 10 < 12$: advance $G[3]$ to the next multiple of 10 that is ≥ 12 , i.e., $G[3] := 20$. Now $G = [12, 12, 20]$.
- $G[1] = 12 < G[3] = 20$: $G[1] := 20$. $G[2] = 12 < 20$: $G[2] := 24$. Now $G = [20, 24, 20]$.
- Continuing, the entries keep leapfrogging until they stabilize at $G = [60, 60, 60]$, at which point $4 \mid 60$, $6 \mid 60$, $10 \mid 60$, and the algorithm terminates.

Computational complexity of LLP-LCM

Let $L = \text{lcm}(A)$. Each entry $G[j]$ takes values in $\{A[j], 2A[j], 3A[j], \dots, L\}$ and increases at each advance by at least $A[j]$, so process j makes at most $L/A[j]$ advances. The total number of advances is $\sum_j L/A[j]$. Each advance scans n entries, so the sequential time is $O\left(n \sum_j L/A[j]\right)$.

These bounds are pseudopolynomial in L . For inputs whose LCM is very large relative to the input sizes, an alternative is to compute $\text{lcm}(A)$ via the identity $\text{lcm}(a, b) = ab / \text{gcd}(a, b)$, using LLP-GCD-Descend or LLP-ExtGCD on pairs.

18.6 Non-Linear Congruences via LLP

The LLP-CRT algorithm (Section 18.4) uses the local predicate $G[j] \equiv b_j \pmod{m_j}$ for each process P_j , but neither the LLP framework nor the algorithm itself depends on this predicate being a *linear* congruence. Any *eventually periodic* predicate $P_j(x)$ can serve as the local constraint, and the algorithm will search for the least x satisfying all P_j simultaneously, provided such an x exists.

Definition 18.10 A predicate $P(x)$ on the natural numbers is eventually periodic with period m if there exists x_0 such that $P(x) = P(x+m)$ for all $x \geq x_0$. The standard CRT predicate $P_j(x) \equiv [x \equiv b_j \pmod{m_j}]$ has period m_j and $x_0 = 0$.

A natural non-linear application is the system of *quadratic* congruences

$$x^2 \equiv a_j \pmod{m_j}, \quad j = 1, \dots, r,$$

where a_j is a quadratic residue modulo m_j . The local predicate $P_j(x) = [x^2 \equiv a_j \pmod{m_j}]$ is periodic with period m_j .

The advance step requires finding the next quadratic-residue solution, which for a prime modulus $m_j = p$ amounts to advancing to the next x in $\{s_j, m_j - s_j, m_j + s_j, 2m_j - s_j, \dots\}$ where s_j is a square root of a_j modulo p . For composite moduli, the set of solutions in one period is found by the standard CRT lift of solutions modulo prime powers.

Algorithm LLP-QuadCRT: LLP algorithm for simultaneous quadratic congruences

Input: m : array[1.. r] of int; a : array[1.. r] of int

Output: least x such that $\forall j : x^2 \equiv a_j \pmod{m_j}$

- 1 G : array[1.. r] of int initially $\forall j : G[j] :=$ smallest non-negative x with $x^2 \equiv a_j \pmod{m_j}$;
 - 2 **forbidden**(j): $(\exists i : G[j] < G[i]) \vee (G[j]^2 \pmod{m_j} \neq a_j)$;
 - 3 **advance**(j): $G[j] :=$ next $x > G[j]$ with $x^2 \equiv a_j \pmod{m_j}$ and $x \geq \max_i G[i]$;
-

Example 18.11 Consider $x^2 \equiv 1 \pmod{5}$ and $x^2 \equiv 1 \pmod{7}$. The solutions modulo 5 are $\{1, 4\}$ (period 5), and modulo 7 are $\{1, 6\}$ (period 7). So G starts at $[1, 1]$ — already equal, and the algorithm terminates immediately with $x = 1$. Indeed, $1^2 \equiv 1$ modulo both 5 and 7.

For the next solution, we could restart with $G[1] := 4$ (the other root mod 5) and $G[2] := 1$:

- $G[2] = 1 < G[1] = 4$: advance to next $x \geq 4$ with $x^2 \equiv 1 \pmod{7}$; that is $G[2] := 6$. Now $G = [4, 6]$.
- $G[1] = 4 < 6$: next $x \geq 6$ with $x^2 \equiv 1 \pmod{5}$; $x = 6$ since $6^2 = 36 \equiv 1 \pmod{5}$. Now $G = [6, 6]$.

The algorithm returns $x = 6$. Check: $6^2 = 36 \equiv 1 \pmod{5}$ and $36 \equiv 1 \pmod{7}$. ✓

This approach extends to any family of periodic local predicates — for instance, $x^3 \equiv a_j \pmod{m_j}$ (cubic residues), $f(x) \equiv 0 \pmod{m_j}$ for a fixed polynomial f , or even predicates defined by a lookup table. The LLP framework handles them all uniformly: the only requirement is that each P_j be computable and eventually periodic, so that the advance step is well-defined.

18.7 Multiplicative Order via LLP

Given a natural number a and a modulus n with $\gcd(a, n) = 1$, the *multiplicative order* of a modulo n , written $\text{ord}_n(a)$, is the smallest positive integer k such that $a^k \equiv 1 \pmod{n}$. By Lagrange's theorem, $\text{ord}_n(a)$ divides $\phi(n)$. Computing the order is important for primality testing, finding primitive roots, and determining the period of linear recurrences modulo n .

The order can be computed by a *descending* LLP on the *divisor lattice* of $\phi(n)$. Recall that the set of divisors of $\phi(n)$, ordered by divisibility, forms a finite distributive lattice. The top element is $\phi(n)$ itself, the bottom is 1, and meets and joins are gcd and lcm. We want the *least* element k of this lattice such that $a^k \equiv 1 \pmod{n}$.

The feasibility predicate is

$$B_{\text{ord}}(k) \equiv a^k \equiv 1 \pmod{n}.$$

This predicate is *upward closed* (lattice-linear for a descending search) in the divisor lattice: if $a^k \equiv 1 \pmod{n}$ and $k \mid m$, then $a^m = (a^k)^{m/k} \equiv 1 \pmod{n}$. Hence, if the predicate holds at k , it holds at every multiple of k . Equivalently, if it fails at k , then it fails at every divisor of k , so we should try to reduce k .

We represent the current candidate k by its prime factorization: if $\phi(n) = p_1^{e_1} p_2^{e_2} \cdots p_s^{e_s}$, then $k = p_1^{f_1} p_2^{f_2} \cdots p_s^{f_s}$ with $0 \leq f_i \leq e_i$. This gives us a vector $G = (f_1, \dots, f_s) \in [0, e_1] \times \cdots \times [0, e_s]$ on which we run a descending LLP:

Algorithm LLP-Order: Computing the multiplicative order of a modulo n

Input: a, n : natural numbers with $\gcd(a, n) = 1$; $\phi(n) = p_1^{e_1} \cdots p_s^{e_s}$: factorization of $\phi(n)$

Output: $\text{ord}_n(a)$

1 G : array[1.. s] of int initially $\forall i : G[i] = e_i$;

2 $k := \phi(n)$;

3 **forbidden**(i): $a^{k/p_i} \equiv 1 \pmod{n}$;

// can reduce the exponent of p_i

4 **advance**(i): $G[i] := G[i] - 1$; $k := k/p_i$;

5 **return** k ;

Lemma 18.12 B_{ord} is lattice-linear on the divisor lattice of $\phi(n)$, and Algorithm LLP-Order terminates with $k = \text{ord}_n(a)$.

Proof: As argued above, B_{ord} is upward closed in the divisor lattice: it holds for k if and only if $\text{ord}_n(a) \mid k$. If the current k is not the order, then $\text{ord}_n(a)$ strictly divides k , so there exists some prime p_i with $\text{ord}_n(a) \mid k/p_i$, i.e., $a^{k/p_i} \equiv 1 \pmod{n}$: this is exactly the forbidden condition. Hence B_{ord} is lattice-linear.

The algorithm starts at $k = \phi(n)$, which satisfies the predicate, and each advance strictly decreases k in the divisor lattice (by reducing one exponent). By the correctness of LLP, the algorithm terminates at the minimum k satisfying the predicate, which is $\text{ord}_n(a)$. ■

Example 18.13 Find $\text{ord}_{13}(2)$. We have $\phi(13) = 12 = 2^2 \cdot 3$, so $s = 2$, $p_1 = 2$, $e_1 = 2$, $p_2 = 3$, $e_2 = 1$, and $G = [2, 1]$ with $k = 12$.

- Test $i = 1$: Is $2^{12/2} = 2^6 = 64 \equiv 64 - 4 \cdot 13 = 12 \equiv -1 \not\equiv 1 \pmod{13}$? No, $G[1]$ is not forbidden.
- Test $i = 2$: Is $2^{12/3} = 2^4 = 16 \equiv 3 \not\equiv 1 \pmod{13}$? No, $G[2]$ is not forbidden.

No index is forbidden, so $k = 12$ is the order. Indeed, $\text{ord}_{13}(2) = 12$, confirming that 2 is a primitive root modulo 13.

Example 18.14 Find $\text{ord}_{13}(3)$. Again $\phi(13) = 12 = 2^2 \cdot 3$, $G = [2, 1]$, $k = 12$.

- Test $i = 1$: $3^{12/2} = 3^6 = 729 \equiv 729 - 56 \cdot 13 = 729 - 728 = 1 \pmod{13}$. Forbidden! Advance: $G[1] := 1$, $k := 6$.
- Test $i = 1$ again: $3^{6/2} = 3^3 = 27 \equiv 27 - 2 \cdot 13 = 1 \pmod{13}$. Forbidden! Advance: $G[1] := 0$, $k := 3$.
- Test $i = 1$: $G[1] = 0$, cannot reduce further. Test $i = 2$: $3^{3/3} = 3^1 = 3 \not\equiv 1 \pmod{13}$. Not forbidden.

The algorithm terminates with $k = 3$. Indeed, $3^1 = 3$, $3^2 = 9$, $3^3 = 27 \equiv 1 \pmod{13}$, so $\text{ord}_{13}(3) = 3$.
✓

Computational complexity of Ord

Let $\phi(n) = p_1^{e_1} \cdots p_s^{e_s}$. The total number of advances is at most $\sum_{i=1}^s e_i$, since each advance reduces some $G[i]$ by 1 and $G[i]$ starts at e_i . Each advance requires one modular exponentiation $a^{k/p_i} \pmod{n}$, which costs $O(\log(k/p_i) \cdot M(n))$ where $M(n)$ is the cost of a multiplication modulo n . Using fast exponentiation, this is $O(\log \phi(n) \cdot M(n))$ per advance.

Sequential complexity. $O(\sum_i e_i \cdot \log \phi(n) \cdot M(n)) = O(\log^2 \phi(n) \cdot M(n))$, since $\sum_i e_i \leq \log_2 \phi(n)$.

18.8 Summary

This chapter applied the Lattice-Linear Predicate framework to a range of number-theoretic computations: the greatest common divisor of an array of numbers (LLP-GCD-Descend and LLP-GCD-Ascend), the extended Euclidean algorithm (LLP-ExtGCD), the Chinese Remainder Theorem (LLP-CRT and LLP-CRT-Max), the least common multiple (LLP-LCM), non-linear congruences (LLP-QuadCRT), and the multiplicative order (Ord). In each case the LLP framework delivered not just a sequential algorithm but also an immediately parallelizable one: since the feasibility predicate is lattice-linear, any set of forbidden indices can be advanced simultaneously, and correctness is independent of the order in which they are processed.

The algorithms illustrate two recurring themes. First, *ascending / descending duality*: every feasibility predicate has both an ascending and a descending lattice-linear formulation, giving rise to complementary algorithms with dual performance profiles. LLP-GCD-Descend descends from A using Euclidean mod, while LLP-GCD-Ascend ascends from the vector of ones using ceiling updates. The same duality reappears in LLP-CRT vs. LLP-CRT-Max, and in LLP-LCM (ascending) vs. a potential descending LCM variant. Second, *generality of the local predicate*: LLP-CRT's correctness depends only on the lattice-linearity of the global conjunction, not on the form of the individual congruences. Replacing linear congruences $x \equiv b_j \pmod{m_j}$ by quadratic ones $x^2 \equiv a_j \pmod{m_j}$, or by any eventually periodic predicate, requires no change to the framework.

The second half of the chapter developed the poset-of-prime-powers representation of natural numbers and used it to prove the classical summation identities for the Möbius function, Euler's totient function, and several other multiplicative arithmetical functions. The proofs exploited the fact that each multiplicative

Algorithm	Problem	Direction	Sequential Time	Parallel Time
LLP-GCD-Descend	GCD of n numbers	descending	$O(n^2 \log M)$	depth $O(\log M)$
LLP-GCD-Ascend	GCD of n numbers	ascending	$O(n^2 \log(M/d))$	depth $O(\log(M/d))$
LLP-ExtGCD	GCD + Bézout pair	descending	$O(\log M)$ (for $n = 2$)	depth $O(\log M)$
LLP-CRT	CRT solution, smallest	ascending	$O\left(rM \sum_j 1/m_j\right)$	depth $O(M/\min_j m_j)$
LLP-CRT-Max	CRT solution, largest $< M$	descending	$O\left(rM \sum_j 1/m_j\right)$	depth $O(M/\min_j m_j)$
LLP-LCM	LCM of n numbers	ascending	$O\left(n \sum_j L/A[j]\right)$	depth $O(L/\min_j A[j])$
LLP-QuadCRT	Simult. quad. congruences	ascending	same as LLP-CRT	same as LLP-CRT
LLP-Order	Multiplicative order	descending	$O(\log^2 \phi(n) \cdot M(n))$	depth $O(\log^2 \phi(n) \cdot M(n))$

Table 18.1: Summary of the LLP-based number-theory algorithms of this chapter. Here n is the number of inputs to the GCD/LCM problems, $M = \max_i A[i]$, $d = \gcd(A)$, $L = \text{lcm}(A)$, r is the number of congruences in the CRT problem, m_1, \dots, m_r are the moduli with product $M = \prod_j m_j$, and $M(n)$ denotes the cost of one multiplication modulo n . Parallel times assume one processor per index of the G array.

function is determined by its values on prime powers, so that a divisor sum over n factors as a product of divisor sums over each prime-power chain.

Throughout the chapter, the LLP framework served as a unifying bridge: number-theoretic structure on one side (divisibility, residues, prime factorization), and a uniform parallel-algorithmic recipe on the other (identify the feasibility predicate, verify lattice-linearity, read off the algorithm). This mirrors the treatment of graph and optimization problems in earlier chapters and sets the stage for the remaining chapters of the book.

18.9 Problems

- Trace Algorithm LLP-GCD-Descend on the array $A = [24, 36, 60, 84]$. Write down the value of G after each advance step, and verify that the algorithm terminates with $G[i] = \gcd(A) = 12$ for every i .
- Prove carefully that if $G[j] > G[i]$ and $G[i]$ does not divide $G[j]$, then the set of common divisors of $G[i]$ and $G[j]$ equals the set of common divisors of $G[i]$ and $G[j] \bmod G[i]$. Use this to conclude that the invariant “every common divisor of A is a common divisor of G ” is preserved by advance in LLP-GCD-Descend.
- Prove that Algorithm LLP-GCD-Descend performs at most $O(n \log M)$ advance steps, where $M = \max_i A[i]$. (Hint: show that the potential function $\Phi(G) = \sum_i \log G[i]$ strictly decreases by a constant after any sequence of advances by a fixed pair (i, j) .)
- Show that if $\gcd(a, b) = 1$, then $\gcd(a + b, ab) = 1$, and that $\gcd(a + b, a - b)$ divides 2.
- Use the Chinese Remainder Theorem to solve the system $x \equiv 1 \pmod{5}$, $x \equiv 2 \pmod{7}$, $x \equiv 3 \pmod{11}$. Trace Algorithm LLP-CRT on this input, and compare with the classical constructive proof of CRT.

6. Adapt LLP-CRT to the case where the moduli are not pairwise coprime. Show that the algorithm still runs, but that it may fail to terminate. Characterize the inputs on which it does terminate, and give the resulting solution.
7. Prove that Euler's totient function satisfies $\phi(mn) = \phi(m)\phi(n)$ whenever $\gcd(m, n) = 1$, by giving an explicit bijection between the residues coprime to mn and pairs of residues coprime to m and to n , respectively. (This is essentially the Chinese Remainder Theorem.)
8. Möbius inversion formula. Let f and g be arithmetical functions. Prove that if $g(n) = \sum_{d|n} f(d)$ for all $n \geq 1$, then $f(n) = \sum_{d|n} \mu(d)g(n/d)$ for all $n \geq 1$. Use this to derive the formula $\phi(n) = \sum_{d|n} \mu(d)(n/d)$.
9. Compute $\mu(n)$ and $\phi(n)$ for all n from 1 to 30 using the poset representation of numbers as prime powers. Verify numerically that $\sum_{d|n} \mu(d) = [n = 1]$ and $\sum_{d|n} \phi(d) = n$.
10. Let $\sigma(n) = \sum_{d|n} d$ denote the sum of the divisors of n . Show that σ is multiplicative, and derive a closed-form formula for $\sigma(p^k)$. What does the Lemma in this chapter say about $\sigma(n)$ in terms of N ?
11. A positive integer n is called *square-free* if no prime appears more than once in its factorization. Using μ and λ , write down an LLP-style algorithm that, given n , decides whether n is square-free.
12. Trace Algorithm LLP-LCM on $A = [6, 8, 12]$. Write down G after each advance and verify that the algorithm terminates with $G[i] = \text{lcm}(A) = 24$ for every i .
13. Find $\text{ord}_{17}(2)$ using Algorithm Ord. Show each step.
14. Use Algorithm Par-QuadCRT to find the smallest positive x satisfying $x^2 \equiv 2 \pmod{7}$ and $x^2 \equiv 3 \pmod{11}$.
15. Let $A = [10, 15]$. Trace Algorithm GCD3 from our description to confirm that it returns 5, and compare its behavior with the classical Euclidean algorithm applied to the same pair.

18.10 Bibliographic Remarks

Euclid's algorithm for the greatest common divisor appears in Book VII of Euclid's *Elements*; it is one of the oldest algorithms known. Knuth's *The Art of Computer Programming*, Volume 2 [Knu97b], provides a thorough modern analysis of Euclid's algorithm, its extended form, and variants such as the binary GCD algorithm, and is the standard reference for algorithmic number theory. Complexity-theoretic aspects and randomized variants are discussed in detail by Shoup [Sho09] and by Bach and Shallit [BS96].

The Chinese Remainder Theorem, in its arithmetic form, was first stated by Sun Tzu in the third century CE. Its modern formulation and the extension to general principal ideal domains were developed in the nineteenth century; [Gau01] treats it systematically in his *Disquisitiones Arithmeticae*. Modern number-theory textbooks such as [Ros11] and [HW08] give accessible treatments with many applications, including secret sharing (Mignotte's and Asmuth–Bloom's schemes) and fast modular multiplication for large integers. Applications to cryptography, in particular the RSA cryptosystem, are discussed in Rivest, Shamir, and Adleman's original paper [RSA78].

Multiplicative arithmetical functions, including μ , ϕ , N , u , and λ , are classical. A systematic treatment can be found in Apostol [Apo76] and in Hardy and Wright [HW08]. The poset-of-prime-powers view that

we emphasize in this chapter is implicit in many treatments of multiplicative functions and is closely related to Rota's theory of Möbius functions on posets [Rot64], which generalizes the number-theoretic Möbius function to arbitrary locally finite partially ordered sets. *Concrete Mathematics* by Graham, Knuth, and Patashnik [GKP94] offers a friendly introduction to these ideas at the undergraduate level.

The lattice-linear predicate framework used here to derive parallel versions of Euclid's algorithm and the Chinese Remainder Theorem is due to Garg [Gar20b]; it unifies a variety of parallel and distributed algorithms, including those for shortest paths, stable matching, and scheduling, within a single algorithmic paradigm.

Chapter 19

Linear Programming

19.1 Introduction

In this chapter, we consider linear programming, which is a general technique for solving many problems. We note here that linear programming has also served as a unifying tool for many combinatorial optimization problems. In particular, the man-optimal stable marriage problem, the shortest path problem, and the least market clearing price can all be modeled as integer linear programs with total unimodularity. However, it is not known whether linear programming is strongly polynomial. In contrast, the LLP method results in algorithms that are close to the best known strongly polynomial algorithms for all these problems. There are other differences in these two approaches. Algorithms for linear programming typically stay inside the feasible space and improve the objective function with every iteration. Our approach keeps the objective function as extremal as possible while making progress towards a feasible solution. The linear programming method uses real variables with linear objective function and linear constraints. Our method assumes that the search is for the infimum element of a feasible space in a distributive lattice such that the feasible space satisfies a lattice-linear predicate.

Linear constraints for linear programming are defined using the conjunction of half-spaces. Lattice-linear predicates are incomparable to half-spaces. There are half-spaces that are not lattice-linear, and there are lattice-linear predicates that are not half-spaces.

Linear programming is a general technique to solve combinatorial optimization problems. Many optimization problems can be modeled using linear or integer linear programs.

Suppose that there is an objective function which we are trying to minimize,

$$7x_1 + x_2 + 5x_3$$

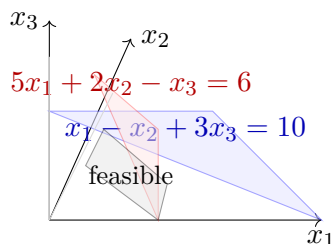
subject to constraints,

$$x_1 - x_2 + 3x_3 \geq 10$$

$$5x_1 + 2x_2 - x_3 \geq 6$$

where x_1, x_2, x_3 are the decision variables and the assumption is that

$$x_1, x_2, x_3 \geq 0$$



Suppose that we have a solution, $x = (2, 1, 3)$. On substituting in the objective function, we get the objective value to be $14 + 1 + 15 = 30$. The reader can verify that this solution is also feasible, as it satisfies the constraints,

$$2 - 1 + 3(3) = 10$$

$$5(2) + 2(1) - 3 = 9$$

Trying to find a bound on primal:

$7x_1 + x_2 + 5x_3 \geq x_1 - x_2 + 3x_3 \geq 10$ by comparing coefficients point-wise and $x_i \geq 0$ solution has to be ≥ 10 but we are trying to get a better or greater lower bound

Add up the constraint equations,

$$6x_1 + x_2 + 2x_3 \geq 16$$

The objective function is still greater than the above equation. Hence new lower bound is 16.

The goal of primal was to minimize the objective function; the goal of its dual will be to maximize its objective function. We have to figure out what combination of each constraint should we add such that it gives the tightest bound on the objective function.

We have n decision variables and m constraints in our primal. So, the dual will have m variables (one variable for each constraint in the primal) and n constraints (since there are n decision variables in the primal).

The objective function is $10y_1 + 6y_2$ subject to constraints,

$$y_1 + 5y_2 \leq 7$$

$$-y_1 + 2y_2 \leq 1$$

$$3y_1 - y_2 \leq 5$$

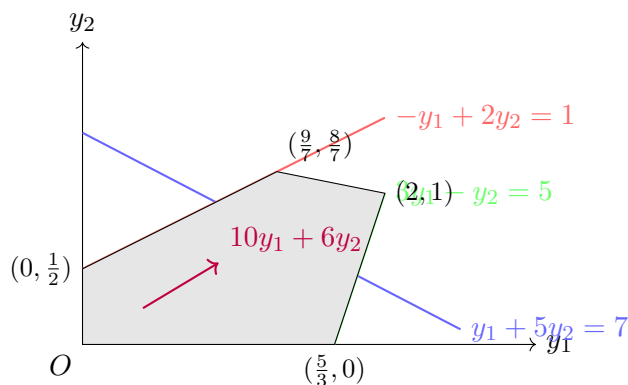


Figure 19.1: Feasible region of the dual LP.

In a generalized representation of the primal, objective function can be written as

$$\sum_{i=1}^n c_i x_i$$

constraints can be written as

$$\sum_j A_{ij} x_j \geq b_i$$

where A is a matrix of dimension $m \times n$, c and b are vectors of size n and m respectively.

For dual, b is translated to c , c is translated to b and matrix A becomes A^T .

This chapter is organized as follows. Section 19.2 discusses duality in linear programming, including strong duality, weak duality, and complementary slackness conditions. Section 19.3 shows how to formulate combinatorial optimization problems such as maximum matching, stable marriage, and shortest path as linear programs. Section 19.4 compares linear programming with the lattice-linear predicate approach.

19.2 Duality in Linear Programming

Strong Duality

Theorem 19.1 *The primal program has finite optimum if and only if its dual has finite optimum. Moreover, if x^* and y^* are optimal solutions for (P) and (D) then,*

$$\sum_{j=1}^n c_j x_j^* = \sum_{i=1}^m b_i y_i^*$$

This theorem translates into many min-max theorems such as max flow = min cut, max matching = min vertex cover and min number of chains to cover a poset = max sized anti-chain.

Weak Duality

Theorem 19.2 *If x and y are feasible solutions for (P) and (D) then,*

$$\sum_{j=1}^n c_j x_j \geq \sum_{i=1}^m b_i y_i$$

Proof:

$$\sum_{j=1}^n c_j x_j \geq \sum_{j=1}^n \left(\sum_{i=1}^m a_{ij} y_i \right) x_j$$

This claim is true because y is feasible and $x \geq 0$

$$\sum_{i=1}^m \left(\sum_{j=1}^n a_{ij} x_j \right) y_i \geq \sum_{i=1}^m b_i y_i$$

This claim is true because x is feasible and $y \geq 0$

Hence proved,

$$\sum_{j=1}^n c_j x_j \geq \sum_{i=1}^m b_i y_i$$

■

Complementary Slackness Conditions

Theorem 19.3 *Given that x, y are (P) and (D) feasible solutions. Then x and y are both optimal if and only if the following complementary slackness conditions are satisfied:*

$$\forall j \quad \text{either}(x_j = 0) \quad \text{or} \quad \sum_{i=1}^m a_{ij} y_i = c_j$$

$$\forall i \quad \text{either}(y_i = 0) \quad \text{or} \quad \sum_{j=1}^n a_{ij} x_j = b_i$$

19.3 Formulating Combinatorial Optimization Problems as Linear Programs

Maximum Matching Problem

Primal:

$$\max \sum_{i,j} W_{ij} x_{ij}$$

subject to:

$$\sum_j x_{ij} \leq 1$$

and

$$\sum_i x_{ij} \leq 1$$

where x_{ij} is an edge between A and B, i is an index from A and j is an index from B. For each edge, there is a weight W_{ij} . We need to pick the edges such that the sum of their weights is maximum.

Replacing integral constraint by linear constraint,

$$x_{ij} \in \{0, 1\} \Rightarrow x_{ij} \geq 0$$

The above transformation is called LP-Relaxation. This can be used only when we can prove that the optimal solution is always going to be integral.

When $|A| = |B| = n$, the number of decision variables equals n^2 , and the number of constraints is $2n$.

Dual:

The number of decision variables equals $2n$ and the number of constraints equals n^2 .

Objective function (minimizing vertex labels):

$$\min \sum u_i + \sum v_j$$

subject to: $u_i + v_j \geq w_{ij}$
 $u_i \geq 0$ and $v_j \geq 0$
 where u_i is constraints for side A and v_j is constraints for side B

The Transportation Problem

We have n supply nodes and n demand nodes. The supply at node i is given by a_i and the demand at node j is given by b_j . We assume that the total supply is equal to the total demand. If the supply and demand do not match, we can create fake nodes to ensure the equality.

Let w_{ij} be the cost function when an item is shipped from i to node j . Let the variable x_{ij} be the number of items i shipped from the source node i to the demand node j . Our goal is to choose x_{ij} such that the cost of transportation is minimized. $\sum_{i,j} w_{ij}x_{ij}$ is minimized subject to:

$$\begin{aligned} \sum_j x_{ij} &= a_i \text{ for all } i \\ \sum_i x_{ij} &= b_j \text{ for all } j \\ \text{and } x_{ij} &\geq 0. \end{aligned}$$

The dual of the relaxed linear program is:

$$\begin{aligned} &\text{maximize } \sum_i a_i u_i + \sum_j b_j v_j \text{ subject to} \\ &u_i + v_j \leq w_{ij} \text{ for all } i, j, \\ &u_i \geq 0 \text{ for all } i, \\ &v_j \geq 0 \text{ for all } j. \end{aligned}$$

The Stable Marriage Problem

Let $x_{i,j}$ denote that man i is matched to woman j . We require that $x_{i,j} \in \{0, 1\}$. The constraint that every man and every woman is matched to a single partner is given as follows. For each man i ,

$$\sum_j x_{ij} = 1.$$

Similarly, for each woman j ,

$$\sum_i x_{ij} = 1.$$

Furthermore, we would like this assignment to not have any blocking pair. For any man i and woman j , they do not form a blocking pair

This condition can be expressed as follows. For any man i and woman j , if j is matched to a man who is less preferred than i , then man i must be matched to a woman who he prefers to j . Otherwise, (i, j) will form a blocking pair for the matching. Let $Q(j, i)$ be the set of men who are less preferable than i to woman j , i.e.,

$$Q(j, i) = \{i' \mid \text{rank}[j][i'] > \text{rank}[j][i]\}$$

Then, the expression $\sum_{i' \in Q(j, i)} x_{i', j}$ is one iff j is matched to someone who is less preferable than i . Similarly, let $R(i, j)$ be the set of women who are more preferable than woman j to man i . Then, the expression $\sum_{j' \in R(i, j)} x_{i, j'}$ is one iff i is matched to someone who is more preferable than woman j . Given these two expressions, we can write the condition that (i, j) is not a blocking pair as

$$\sum_{i' \in Q(j, i)} x_{i', j} - \sum_{j' \in R(i, j)} x_{i, j'} \leq 0.$$

Thus, we have $O(n^2)$ constraints.

The Shortest Path Problem

Let x_i denote the distance from the source vertex x_0 to x_i . Let t be the destination vertex. We assume that there are no incoming edges to x_0 or outgoing edges from x_t .

We would like to maximize $x_t - x_0$ such that for all edges $(i, j) \in E$: $x_j \leq x_i + w[i, j]$. Hence, the LP formulation is as follows.

$$\begin{aligned} & \text{maximize} && x_t - x_0 \\ & \text{subject to} && x_j \leq x_i + w[i, j] \quad \forall (i, j) \in E, \\ & && x_i \geq 0 \quad \forall i \end{aligned} \tag{19.1}$$

This formulation is identical to that we used in designing an LLP based algorithm (or equivalently, BellmanFord algorithm). We assume that there is no incoming edge to x_0 . In this linear program, there are n variables, one for each node, and there are m constraints, one for each edge. The variables x_i are nonnegative.

The dual of this linear program is as follows.

We have m variables, $y_{i,j}$ for each edge $(v_i, v_j) \in E$. The variable $y_{i,j}$ can be viewed as the flow on the edge (v_i, v_j) .

We would like to minimize $\sum_{(i,j) \in E} w[i, j] * y[i, j]$ as follows.

$$\begin{aligned} & \text{minimize} && \sum_{(i,j) \in E} w[i, j] * y[i, j] \\ & \text{subject to} && \\ & && \text{for each node } v_i \text{ other than } v_0 \text{ and } v_t: \sum_j y_{j,i} - \sum_k y_{i,k} = 0. \\ & && \text{for } v_t, \sum_j y_{j,t} = 1. \\ & && \text{for } v_0, -\sum_j y_{0,j} = -1. \\ & && \text{for all } i, j: y[i, j] \geq 0. \end{aligned}$$

19.4 Comparison of Linear Programming with Lattice-Linear Predicates

Linear programming can also be viewed as a search for an optimal feasible solution. However, there are many important differences.

- **Vector Space vs Distributive Lattice:** First, the underlying space in linear programming is the set of real valued vectors whereas the underlying space in the lattice-linear predicate detection method is a distributive lattice.

In the domain of distributive lattices, we do not have addition or the scalar multiplication as in vector spaces. All of lattice-linear predicate algorithms use the following two operations: meet of the underlying lattice and the “advance” operation. The advance operation maps an element of the lattice to a bigger element in the lattice.

- **Polyhedron vs Meet-Closed Predicates:** In linear programming, the feasible space is characterized by a polyhedron (or the set of vectors x such that $Ax \leq 0$ for some matrix A). There is no lattice

structure required on the feasible space. It is not guaranteed that if two vectors are feasible, then their component-wise minimum vector is also feasible. Lattice-linear predicate detection requires the feasible space to be closed under meets.

Linear programming has its optimization objective as minimization or maximization of a linear cost function. Lattice linear predicate detection simply uses the underlying order operation to define optimization. Since feasible space is closed under meets, the infimum of the feasible space is well-defined.

- **Efficiency of the General Algorithm:** Even though many problems studied in this book can also be solved via linear programming, the algorithms derived in that manner are not as efficient as the LLP algorithm. The following list gives the algorithms that are used for Linear Programming.
 1. **Simplex Method:** Developed by George Dantzig, it is widely used for solving linear programming problems by moving from one vertex of the feasible region to an adjacent one with a higher objective value until the optimum is reached. The *dual Simplex method* is a variant of the simplex method, used when the initial solution is not feasible. It modifies the constraints to move towards feasibility and optimality simultaneously. The worst-case time complexity is exponential, but in practice, it is often efficient.
 2. **Interior Point Methods:** These methods, including Karmarkar's algorithm, approach the optimal solution from within the feasible region. They can be faster than the simplex method for large-scale problems.
 3. **Ellipsoid Method:** Introduced by Khachiyan, this was the first polynomial-time algorithm for linear programming. It's more of theoretical interest as it's outperformed by other methods in practice.
 4. **Cutting Plane Method:** Adds additional linear constraints (cuts) to the problem to exclude regions that do not contain the optimal solution, gradually cutting closer to the optimal point.
 5. **Branch and Bound Method:** Often used for mixed-integer linear programming, this method explores branches of possible solutions and bounds them to identify the optimal solution.
- **Parallelism of the General Algorithm:** The LLP algorithm determines which components in G are forbidden and advances on all of those components in parallel. The worst case complexity is given by the height of the lattice.

19.5 Summary

This chapter introduced linear programming as a general technique for solving combinatorial optimization problems and compared it with the lattice-linear predicate approach.

Topic	Method	Time Complexity
Linear Programming	Simplex Method	Exponential (worst case)
Linear Programming	Ellipsoid Method	Polynomial
Linear Programming	Interior Point Methods	Polynomial
LLP Detection	Lattice-Linear Predicate	Height of the lattice

19.6 Problems

1. A company produces two products, X and Y , with profits of \$3 and \$4 per unit, respectively. Production requires labor and materials, with constraints:
 - Labor: $2X + Y \leq 8$ hours.
 - Materials: $X + 2Y \leq 8$ units.
 - Non-negativity: $X, Y \geq 0$.

Formulate and solve the linear program to maximize profit, using the graphical method.

2. A factory produces three items, A , B , and C , with profits \$5, \$3, and \$4 per unit. Constraints are:
 - Machine hours: $3A + B + 2C \leq 12$.
 - Labor hours: $A + 2B + C \leq 10$.
 - Non-negativity: $A, B, C \geq 0$.

Formulate and solve using the simplex method to maximize profit.

3. Formulate the max-flow problem for the network below as a linear program and solve it.

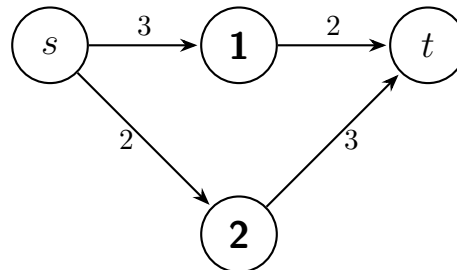


Figure 19.2: Flow network for Problem 3.

19.7 Bibliographic Remarks

Linear programming optimizes a linear objective subject to linear constraints. [Dan63] introduced the simplex method, a cornerstone algorithm, and developed foundational theory. [Kar84] proposed the interior-point method, establishing polynomial-time solvability. For a comprehensive modern treatment, including duality and applications, see [Van20].

Chapter 20

The Predicate Detection Problem

20.1 Introduction

Debugging and testing multithreaded software is widely acknowledged to be a hard task. Sometimes it takes a programmer days to locate a single bug, especially when the bug appears in one thread schedule but not in others. The current debugging and testing method for multithreaded programs is as follows. The programmer tries the program with multiple inputs in the hope of finding a faulty execution. However, the behavior of a multithreaded program depends not only on the external user input, but also the thread schedule and the order in which locks are obtained by the program. It is easy for the testing process to miss a bug that arises with an alternate schedule. One of the fundamental problems in debugging these systems is to check if the user-specified condition exists in *any* global state of the system that can be reached by an alternative thread schedule. This problem, called *predicate detection*, takes a concurrent computation (in an online or offline fashion) and a condition that denotes a bug (for example, violation of a safety constraint), and outputs a schedule of threads that exhibits the bug if possible. Predicate detection is predictive because it generates inferred reachable global states from the computation; an inferred reachable global state might not be observed during the execution of the program, but is possible if the program is executed in a different thread interleaving.

This chapter is organized as follows. We first discuss the complexity of the predicate detection problem, establishing NP-completeness results for general boolean predicates. Section 20.3 presents a work-efficient algorithm for detecting conjunctive predicates using the State Rejection Graph technique. We then give an NC algorithm for conjunctive predicate detection that runs in $O(\log mn)$ time on the CRCW PRAM. We show that detecting a conjunctive predicate at a given level is NP-complete. Finally, we address the problem of recognizing lattice-linear and regular predicates, showing that these recognition problems are co-NP-complete.

20.2 Complexity of Predicate Detection Problem

In this section, we explore the complexity-theoretic results for predicate detection. It is not surprising that given a parallel program computation and a boolean predicate, it is computationally hard to determine if the execution went through a global state in which the predicate became true. We also show that

given a boolean predicate b and an execution, determining whether b is lattice-linear for that execution is co-NP-complete.

Since there are N processes, the total number of global states possible is m^N , where m is the number of state intervals at any process. Consider a boolean predicate B . Even when B is a boolean expression, and processes do not communicate, the problem of detecting *possibly*: B is NP-complete.

We show in this section that the problem of global predicate detection is NP-complete. In fact, we show that it is NP-complete even in the absence of messages between processes.

The global predicate detection problem is a decision problem. It can be written as:

Input instance: a poset S of N sequences, a set of variables X partitioned into N subsets X_1, \dots, X_N , and a predicate B defined on X .

Problem: Determine whether there exists a consistent cut $G \in S$ such that $B(G)$ has the value true.

We now show:

Theorem 20.1 *The global predicate detection problem is NP-complete.*

Proof: First note that the problem is in NP. The verification that the cut is consistent can easily be done in polynomial time (for example, using vector clocks and examining all pairs of states from the cut). Therefore, if the predicate itself can be evaluated in polynomial time, then the detection of that predicate belongs to the set NP.

We show NP-completeness of the simplified predicate detection problem where all program variables are restricted to taking the values “true” or “false”, and at most one variable from each X_i can appear in B . We reduce the satisfiability problem of a boolean expression (SAT) to the global predicate detection problem by constructing an appropriate poset.

The poset is constructed as shown in Figure 20.1. For each variable $u_i \in U$, we define a process P_i that hosts variable x_i (i.e., $X_i = \{x_i\}$). Let the sequence S_i consist of exactly two states. In the first state, x_i has the value false. In the second state, x_i has the value true.

It is easily verified that the predicate B is true for some cut in S if and only if the expression is satisfiable. ■

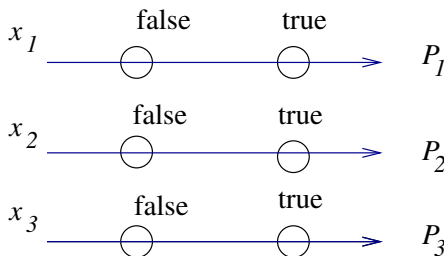


Figure 20.1: Transformation from SAT to global predicate detection

The above result shows that detection of a general global predicate is intractable even for simple distributed computations.

20.3 An Algorithm for Detecting Conjunctive Predicates

In this section we describe a work-efficient algorithm to detect a conjunctive predicate $B = l_1 \wedge l_2 \wedge \dots \wedge l_n$. To detect B , we need to determine if there exists a consistent global state G such that B is true in G . Note that given a computation on n processes each with m states, there can be as many as m^n possible consistent global states. Therefore, enumerating and checking the condition B for all consistent global states is not feasible. Since B is conjunctive, it is easy to show [GW92] that B is true iff there exists a set of states s_1, s_2, \dots, s_n such that (1) for all i , s_i is a state on P_i , (2) for all i , l_i is true on s_i and (3) for all i, j : $s_i \parallel s_j$. Our detection algorithm will either output such local states or guarantee that it is not possible to find them in the computation. When the global predicate B is true, there may be multiple G such that B holds in G . We are interested in algorithms that return the minimum G that satisfies B . The minimum G corresponds to the smallest counter-example to a programmer's understanding because B typically represents the violation of a safety constraint.

Our algorithm is based on the setting where the execution traces for all processes have been collected at one process. For example, in the centralized algorithm for conjunctive predicate detection, one process serves as a checker and collects the traces. All other processes involved in detecting the conjunctive predicate, referred to as application processes, check for local predicates during the computation. Each process P_i also maintains the vector clock algorithm. Whenever the local predicate of a process becomes true for the *first* time since the most recently sent message (or the beginning of the trace), it generates a debug message containing its local timestamp vector and sends it to the checker process [GW92].

The checker process uses queues of incoming messages to hold incoming local snapshots from application processes. We require that messages from an individual process be received in FIFO order. If the underlying system is non-FIFO, then sequence numbers can be attached with messages to ensure FIFO delivery. At the end of the computation, the checker process has a sequence of local states from each process where its local predicate is true. We now describe a sequential and a algorithm to detect B on these traces. The sequential algorithm is an adaptation of the algorithm from [GW92]. We include it here because it is instrumental in understanding the algorithm. Moreover, the correctness of the algorithm is shown by assuming the correctness of the sequential algorithm.

The algorithm in Fig. 20.3 takes as input n traces each of size m as shown in Fig. 20.2. Since conjunctive predicates are lattice-linear, we simply use LLP algorithm to detect them. We use G for the consistent global state. A local state $G[j]$ is forbidden if it is less than $G[i]$ for some i . Since we are interested only in global states where the local predicate is true for all $G[i]$, we assume that for any i , we only consider $G[i]$ such that the local predicate is true in $G[i]$.

20.4 Detecting a Conjunctive Predicate at the Given Level

We now show that, in general, asking for a conjunctive predicate on a particular level is **NP**-complete.

Theorem 20.2 *Given a distributed computation, deciding whether there exists a global state with k events satisfying a given conjunctive predicate is **NP**-complete.*

Proof: We first show that the problem is in **NP**. The global state itself provides a succinct certificate. We can check that all local predicates are true in that global state and that the global state is at level k .

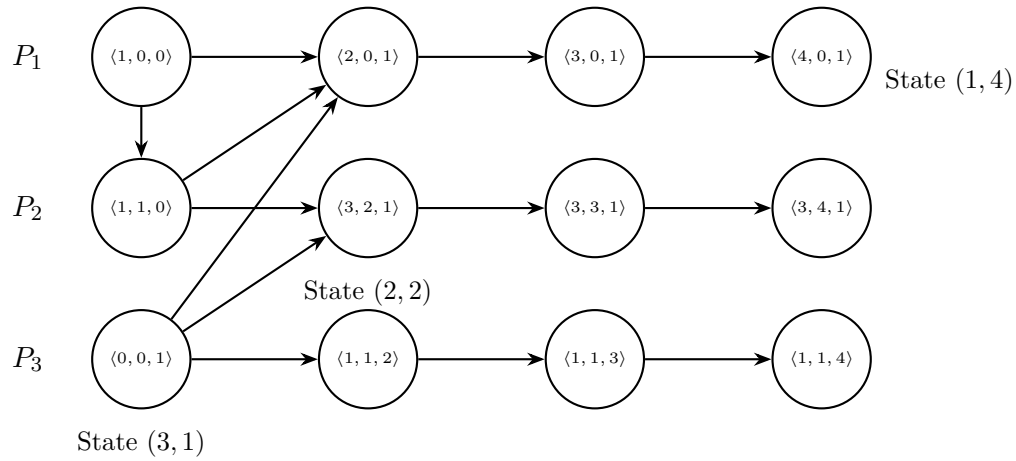


Figure 20.2: State-Based Model of a Distributed Computation

```

function ConjunctiveAlgorithm()
var
   $G$ : array[1.. $n$ ] of int initially 1;
   $T = (m_1, m_2, \dots, m_n)$ ; //maximum number of proposals at  $P_i$ 
always
   $forbidden(G, j, B) \equiv \exists i : G[j] \rightarrow G[i]$ 
while  $\exists j : forbidden(G, j, B)$  do
  for all  $j$  such that  $forbidden(G, j, B)$  in parallel:
    if ( $G[j] = T[j]$ ) then return null;
    else  $G[j] := G[j] + 1$ ;
endwhile;
return  $G$ ; // satisfying global state

```

Figure 20.3: Conjunctive Predicate Detection Algorithm.

```

function ParallelCut()
Input: states : array[1...n][1...m] of vectorClock // Sequence of local states at each process
Output: Consistent Global State as array cut[1...n]

Step 1: Create F: set of states rejected in the first round
  var F : array[1...n] of 0...1 initially 0;
  for all (i ∈ 1...n, j ∈ 1...n) in parallel do
    if ((i, 1) → (j, 1)) then
      F[i] := 1;

Step 2: Create R: State Rejection Graph // Represented as an Adjacency Matrix
  var R : [(1...n, 1...m), (1...n, 1...m)] of 0...1;
  for all (i ∈ 1...n, j ∈ 1...m) in parallel do
    R[(i, j), (i, j)] := 1;
  for all (i ∈ 1...n, j ∈ 1...m - 1, i' ∈ 1...n, j' ∈ 1...m)
    such that i ≠ i' in parallel do
      if ((i', j') → (i, j + 1)) then
        R[(i, j), (i', j')] := 1;
      else
        R[(i, j), (i', j')] := 0;

Step 3: Create RT: transitive closure of R
  var RT : array[(1...n, 1...m), (1...n, 1...m)] of 0...1;
  RT := TransitiveClosure(R);

Step 4: Create valid: replace invalid states by 0
  var valid : array[1...n][1...m] of 0...1;
  for all (i ∈ 1...n, j ∈ 1...m) in parallel do
    valid[i][j] := 1;
  for all (i ∈ 1...n, i' ∈ 1...n, j' ∈ 1...m) in parallel do
    if (F[i] = 1) ∧ (RT[(i, 1), (i', j')] = 1) then
      valid[i'][j'] := 0;

Step 5: Create cut: First Consistent Global State
  var cut : array[1...n] of 0...m initially 0;
  for all (i ∈ 1...n, j ∈ 1...m) in parallel do
    if (valid[i][j] ≠ 0) then
      if (j = 1) ∨ ((j > 1) ∧ (valid[i][j - 1] = 0)) then
        cut[i] := j;
  for all (i ∈ 1...n) in parallel do
    if (cut[i] = 0) then
      output("No satisfying Consistent Cut");

return ConsistentCut := cut;

```

Figure 20.4: The ParallelCut algorithm to find the first consistent cut.

For hardness, we use the subset sum problem. Given a subset problem on n positive integers, x_1, x_2, \dots, x_n with the requirement to choose a subset that adds up to k , we create a computation on n processes as follows. Each process P_i has x_i events. The local predicate on P_i is true initially and also after it has executed x_i events. Thus, the local predicate is true at each process exactly twice. The problem asks us if there is a global state with k events in which all local predicates are true. Such a global state, if it exists, would choose for every process either the initial local state or the final local state. All the final states that are chosen correspond to the numbers that have been chosen.

To avoid the expansion of the numbers in binary to unary construction, we encode the representation of events on a process as follows: Since the conjunctive predicates can only be true when the local predicates are true, we keep only local states which satisfy their corresponding local predicate and store the number of local events executed so far with them. This leaves two local states at each process: The initial state with zero events executed until that point, and the final state of the process with the number of events equal to x_i for the i^{th} process. Now, the construction of the computation from the subset sum problem is polynomial in the size of the input.

■

20.5 Recognizing Lattice-Linear and Regular Predicates

As discussed earlier, efficient detection algorithms exist for various classes of predicates. Thus, given a boolean expression B , one would like to determine if it belongs to a tractable subclass, in which case detection of the predicate may be performed efficiently. We first consider the classes of Lattice-Linear and regular predicates. In this section, we show that determining whether a given boolean expression is Lattice-Linear with respect to a given distributed computation is a co-NP-complete problem. We also show that this problem is co-NP-complete for regular predicates, as well as dual-Lattice-Linear predicates.

We define the decision problems of predicate recognition for Lattice-Linear and regular predicates as follows.

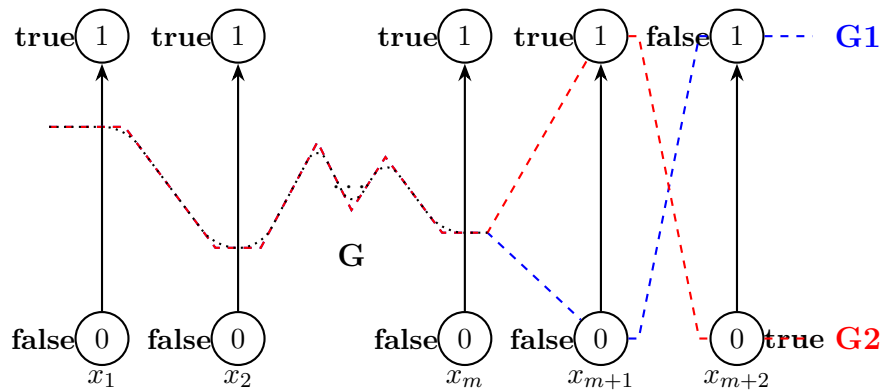


Figure 20.5: Transformation for Theorems 20.3 and 20.4.

Lattice-Linearity: Given a boolean expression b and a program computation, is b a Lattice-Linear predicate?

Regularity: Given a boolean expression b and a program computation, is b a regular predicate?

Theorem 20.3 *Lattice-Linearity is co-NP-complete.*

Proof: *Lattice-Linearity is in co-NP:* Given a pair of candidate global states G and H in which the predicate is true, it can be easily verified in polynomial time that the predicate is false in the global state $G \cap H$. Thus, Lattice-Linearity is in co-NP.

Lattice-Linearity is co-NP-hard: To show co-NP-hardness, we transform an arbitrary instance of TAUTOLOGY to an instance of Lattice-Linearity.

Let b be a boolean expression involving variables $x_1, x_2, \dots, x_m. \forall i = 1..m$, we place each x_i on a separate process, P_i . Each of these m processes has two local states, a *true* state and a *false* state, which corresponds to the value taken by the variable x_i in that local state. We also define two new variables, x_{m+1} and x_{m+2} , and place them on processes P_{m+1} and P_{m+2} respectively. Process P_{m+1} has two local states: an initial *false* state and a final *true* state, and process P_{m+2} has two local states: an initial *true* state and a final *false* state. Figure 20.5 shows this transformation. It is evident that this transformation can be achieved in polynomial time.

We define

$$B = b \vee x_{m+1}x_{m+2} \vee \bar{x}_{m+1}\bar{x}_{m+2}$$

We claim that B is Lattice-Linear iff b is a tautology. If b is a tautology, then B is trivially Lattice-Linear, since B will be true for all global states. Conversely, if b is not a tautology, then there exists a subcut involving processes $P_1 \dots P_m$ in which b evaluates to false. Let us call this subcut G . We can now extend the subcut G to form two cuts, G_1 and G_2 , in which the predicate B is true, as shown in Figure 20.5.

$$G_1 = (G, 0, 1) \text{ and } G_2 = (G, 1, 0)$$

However, $G_1 \cap G_2 = (G, 0, 0)$ in which the predicate B is false. Thus, B is not Lattice-Linear. ■

Theorem 20.4 *Regularity is co-NP-complete.*

Proof: *Regularity is in co-NP:* Given two candidate global states in which the predicate is true, and their union and intersection such that the predicate is false in either the union or intersection, it can be easily verified in polynomial time that the predicate is not regular. Thus, Regularity is in co-NP.

Regularity is co-NP-hard: The transformation in Theorem 20.3 holds for Regularity as well. That is, b is a tautology iff $B = b \vee x_{m+1}x_{m+2} \vee \bar{x}_{m+1}\bar{x}_{m+2}$ is regular. If b is a tautology, then B is trivially regular since B is true for all global states. Conversely, if b is not a tautology, then B is not regular since both the intersection and union of G_1 and G_2 result in a global state in which B is false. Thus, Regularity is co-NP-complete. ■

Note that the above transformation can also be used to show that the problem of deciding whether a given boolean predicate is dual-Lattice-Linear is co-NP-complete.

We stated earlier that efficient detection of Lattice-Linear predicates relies on the assumption that the given predicate satisfies the efficient advancement property, that is, the forbidden state can be identified

in polynomial time. The question that arises is, do all Lattice-Linear predicates satisfy the efficient advancement property? If not, then is it possible to efficiently detect Lattice-Linear predicates that do not satisfy this property? We show here that, unless $RP=NP$, polynomial-time detection cannot be performed for all Lattice-Linear predicates.

We use a result by Valiant and Vazirani [VV85], which states that satisfiability is NP-hard under randomized reductions even for instances that have at most one satisfying assignment (USAT). Valiant and Vazirani's proof uses a randomized polynomial-time algorithm that reduces a given instance of SAT to an instance of USAT.

Theorem 20.5 (Valiant-Vazirani) *If there exists a randomized polynomial-time algorithm for solving instances of SAT having at most one satisfying assignment, then $NP=RP$.*

We know that a predicate having at most one satisfying assignment is Lattice-Linear, so every instance of USAT is a Lattice-Linear predicate. Given any instance B of USAT, involving variables x_1, x_2, \dots, x_m , one can create a distributed computation as shown in Figure 20.1, such that detecting *possibly* : B is equivalent to solving USAT for B . Since we know that Lattice-Linear predicates that satisfy efficient advancement can be detected in polynomial-time, this indicates that Lattice-Linear predicates that do not exhibit efficient advancement may not be detected in polynomial-time, even by a randomized algorithm. Furthermore, since every instance of USAT is also a regular predicate, detection of regular predicates is also NP-hard under randomized reductions.

20.6 Summary

This chapter studied the complexity of predicate detection and predicate recognition in distributed computations. We showed that general predicate detection is NP-complete and presented efficient algorithms for the important subclass of conjunctive predicates. We also established co-NP-completeness results for recognizing tractable predicate classes.

Problem/Topic	Algorithm/Result	Time Complexity
General predicate detection	NP-complete	—
Conjunctive predicate detection	ParallelCut (CRCW PRAM)	$O(\log mn)$
Conjunctive predicate at level k	NP-complete	—
Recognizing Lattice-Linear	co-NP-complete	—
Recognizing Regular predicates	co-NP-complete	—

20.7 Problems

1. In this chapter, we discuss finding the least consistent cut that satisfies a conjunctive predicate. (a) Show that conjunctive predicates are also closed under joins. (b) Give the predicate $rForbidden(G)$ to detect the largest consistent cut that satisfies a conjunctive predicate.
2. Give an algorithm that finds a consistent cut satisfying a conjunctive predicate by combining the forward search with the backward search.

20.8 Bibliographic Remarks

The detection of conjunctive predicates was discussed by Garg and Waldecker in [GW92]. Distributed on-line algorithms for detecting conjunctive predicates were presented in Garg and Chase [GC95]. Observer-independent predicates were introduced by Charron-Bost, Delporte-Gallet, and Fauconnier [CBDGF95]. Hurfin, Mizuno, Raynal and Singhal [HMRS95] gave a distributed algorithm that does not use any additional messages for predicate detection. Distributed algorithms for offline evaluation of global predicates are also discussed in Venkatesan and Dathan [VD92]. Stoller and Schneider [SS95] have shown how Cooper and Marzullo's algorithm can be integrated with that of Garg and Waldecker's to detect a conjunction of global predicates.

The second algorithm for detecting conjunctive predicates is from [GG19]. The **NP**-completeness of detecting a conjunctive predicate at the level k is shown in [GS24].

The complexity of detecting a boolean predicate is taken from [CG98]. The complexity of checking whether a predicate is lattice-linear (or regular) is from [KG05].

Chapter 21

Appendix: Lattice Theory

This appendix covers the definitions of partial orders and lattices used throughout this book. For a thorough treatment, see [DP90, Grä03, Gar15].

21.1 Relations

A partial order is simply a relation with certain properties. A **relation** R over a set X is a subset of $X \times X$. For example, on $X = \{a, b, c\}$, one possible relation is

$$R = \{(a, c), (a, a), (b, c), (c, a)\}.$$

It is sometimes useful to visualize a relation as a directed graph on the vertex set X with an edge from x to y whenever $(x, y) \in R$.

A relation is **reflexive** if $(x, x) \in R$ for every $x \in X$, and **irreflexive** if $(x, x) \notin R$ for every $x \in X$. On the natural numbers \mathbb{N} , the relation *divides* is reflexive, while *less than* is irreflexive. Note that a relation need not be either reflexive or irreflexive.

A relation R is **symmetric** if $(x, y) \in R$ implies $(y, x) \in R$, and **antisymmetric** if $(x, y) \in R$ and $(y, x) \in R$ implies $x = y$. On \mathbb{N} , *less than or equal to* is antisymmetric. R is **transitive** if $(x, y) \in R$ and $(y, z) \in R$ imply $(x, z) \in R$; the relations *less than* and *equal to* on \mathbb{N} are both transitive.

A relation that is reflexive, symmetric, and transitive is an **equivalence** relation. When R is an equivalence, we write $x \equiv_R y$ for $(x, y) \in R$, and the **equivalence class** $[x]_R = \{y \mid y \equiv_R x\}$. The equivalence classes form a **partition** of X . For example,

$$R = \{(x, y) \mid x \bmod 5 = y \bmod 5\} \tag{21.1}$$

is an equivalence on \mathbb{N} that partitions it into five classes.

21.2 Partial Orders

A relation R is a **reflexive partial order** (or **non-strict partial order**) if it is reflexive, antisymmetric, and transitive. A relation R is an **irreflexive partial order** (or **strict partial order**) if it is irreflexive and transitive. The *divides* relation on \mathbb{N} is a reflexive partial order, and *less than* on \mathbb{N} is an irreflexive

partial order. When R is a reflexive partial order, we write $x \leq_R y$ to denote $(x, y) \in R$. A reflexive partially ordered set, **poset** for short, is denoted by (X, \leq) . When R is irreflexive, we write $x <_R y$. Given an irreflexive partial order, we can define $x \leq y$ as $x < y$ or $x = y$, and vice versa. A relation is a **total order** if it is a partial order and for all distinct $x, y \in X$, either $(x, y) \in R$ or $(y, x) \in R$. The natural order on the integers is a total order, but *divides* on \mathbb{N} is only a partial order.

Finite posets are often depicted using **Hasse diagrams**. For any $x, y \in X$, y **covers** x if $x < y$ and there is no z with $x < z < y$. We write $x <_c y$ to denote that y covers x (y is an **upper cover** of x ; x is a **lower cover** of y). A Hasse diagram draws an edge from x to y whenever $x <_c y$, with x below y . For example, consider the poset

$$X \stackrel{\text{def}}{=} \{p, q, r, s\}; \quad \leq \stackrel{\text{def}}{=} \{(p, q), (q, r), (p, r), (p, s)\}, \quad (21.2)$$

whose Hasse diagram is shown in Figure 21.1.

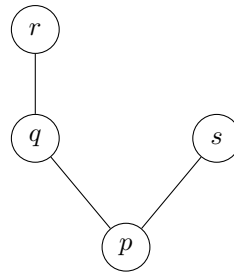


Figure 21.1: Hasse diagram of a poset

Elements x, y are **comparable** if $x < y$ or $y < x$; otherwise they are incomparable, written $x \parallel y$. A poset is a **chain** if every pair is comparable, and an **antichain** if every pair is incomparable. In Figure 21.1, $\{p, q, r\}$ is a chain and $\{q, s\}$ is an antichain. The **height** of a poset is the size of a longest chain, and the **width** is the size of a largest antichain. We use \underline{n} for a chain of n elements and A_n for an antichain of n elements. A total order Q that extends a poset P is called a **linear extension** of P . For example, ordering the poset of Figure 21.1 as $p < q < r < s$ gives one linear extension.

An element x is a **bottom element** (denoted \perp) if $x \leq y$ for all y , and a **top element** (denoted \top) if $y \leq x$ for all y . An element x is **minimal** if there is no y with $y < x$, and **maximal** if there is no y with $y > x$.

21.3 Lattices

Let (X, \leq) be a poset. We define two operators on subsets of X : **meet** (also called **infimum**, or **inf**) and **join** (also called **supremum**, or **sup**). For $Y \subseteq X$ and $m \in X$, we say that $m = \inf Y$ iff

1. $\forall y \in Y : m \leq y$, and
2. $\forall m' \in X : (\forall y \in Y : m' \leq y) \Rightarrow m' \leq m$.

Condition (1) says that m is a lower bound of Y , and condition (2) says that any other lower bound is at most m . For this reason, m is also called the **greatest lower bound** (glb) of Y . The infimum is unique whenever it exists, and m need not itself belong to Y . Dually, for $s \in X$, we say that $s = \sup Y$ iff

1. $\forall y \in Y : y \leq s$, and
2. $\forall s' \in X : (\forall y \in Y : y \leq s') \Rightarrow s \leq s'$.

s is also called the **least upper bound** (lub) of Y . We denote the glb of $\{a, b\}$ by $a \sqcap b$ and the lub of $\{a, b\}$ by $a \sqcup b$. In the natural numbers ordered by *divides*, the meet of two numbers is their gcd and the join is their lcm. The meet or join need not always exist: in Figure 21.1, $\{q, s\}$ has no upper bound; in poset (iii) of Figure 21.2 below, $\{b, c\}$ has upper bounds d and e but no least upper bound.

Definition 21.1 (Lattice) A poset (X, \leq) is a **lattice** iff $\forall x, y \in X : x \sqcup y$ and $x \sqcap y$ exist.

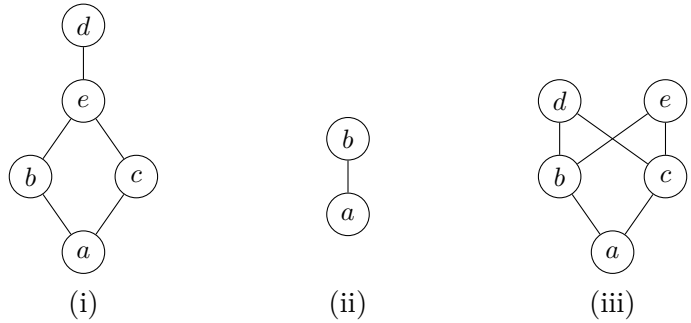


Figure 21.2: Only the first two posets are lattices. In (iii), $\{b, c\}$ has no least upper bound.

The first two posets in Figure 21.2 are lattices; the third is not, since $\{b, c\}$ has upper bounds d and e but no least upper bound. If only \sqcup (resp. \sqcap) exists for all pairs, we have a *sup semilattice* (resp. *inf semilattice*). A lattice in which every subset has both a sup and inf is a *complete lattice*.

A nonempty $S \subseteq L$ is a **sublattice** of a lattice L iff $\forall a, b \in S : \sup(a, b) \in S$ and $\inf(a, b) \in S$, where sup and inf are computed in L .

A subset $Y \subseteq X$ of a poset is a **down-set** (or **order ideal**) if $z \in Y$ and $y \leq z$ implies $y \in Y$. The set $D[x] = \{y \mid y \leq x\}$ is a **principal order ideal**. Dually, Y is an **up-set** (or **order filter**) if $y \in Y$ and $y \leq z$ implies $z \in Y$; $U[x] = \{y \mid x \leq y\}$ is a **principal order filter**. In Figure 21.1, $D[r] = \{p, q, r\}$ and $U[p] = \{p, q, r, s\}$. We also write $D(x) = \{y \mid y < x\}$ and $U(x) = \{y \mid x < y\}$.

21.4 Distributive Lattices and Birkhoff's Theorem

An element x in a poset P is **join-irreducible** if $\forall Y \subseteq P : x = \sup Y$ implies $x \in Y$. The bottom element (if it exists) is not join-irreducible. The set of join-irreducible elements is denoted $J(P)$; in a finite lattice, $x \in J(P)$ iff x has exactly one lower cover. In poset (i) of Figure 21.2, $J = \{b, c, d\}$ (and $e \notin J$ since $e = b \sqcup c$). By duality, **meet-irreducible** elements are denoted $M(P)$. The join-irreducibles form a basis: for any $x \in P$, $x = \sup(D[x] \cap J(P))$ (see [DP90]).

Definition 21.2 (Distributive Lattice) A lattice L is **distributive** if $\forall a, b, c \in L : a \sqcap (b \sqcup c) = (a \sqcap b) \sqcup (a \sqcap c)$.

Any power-set lattice is distributive, as is the lattice of natural numbers under *divides*. A finite lattice is nondistributive iff it contains a sublattice isomorphic to the Pentagon N_5 or the Diamond M_3 [Bir67, DP90]:

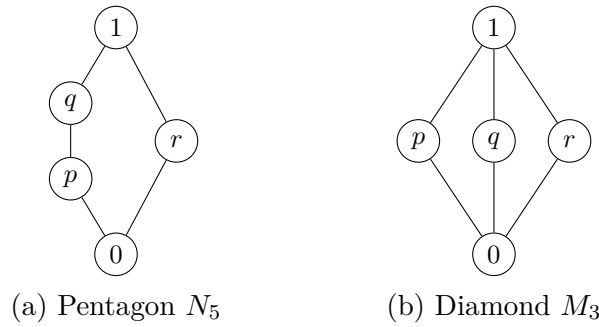


Figure 21.3: The two smallest nondistributive lattices.

Let $L_I(P)$ denote the set of order ideals of a poset P , ordered by inclusion. Birkhoff’s Theorem establishes that every finite distributive lattice is isomorphic to the lattice of order ideals of its join-irreducible elements.

Theorem 21.3 [Birkhoff’s Theorem [Bir67]] *Let L be a finite distributive lattice. Then the mapping*

$$f: L \rightarrow L_I(J(L)), \quad a \mapsto f(a) = \{x \in J(L) : x \leq a\}$$

is an isomorphism from L onto $L_I(J(L))$.

The poset $J(L)$ can be exponentially more succinct than L itself. An example is shown in Figure 21.4: the five-element lattice L has three join-irreducibles a, b, c (with $a < b$ and $a < c$ in $J(L)$), and $L_I(J(L)) \cong L$.

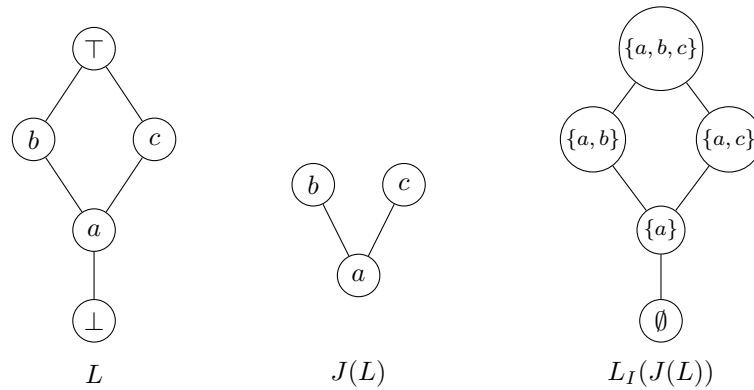


Figure 21.4: A lattice L , its join-irreducibles $J(L)$, and the lattice of ideals $L_I(J(L))$.

Birkhoff’s Theorem also translates properties between L and $P = J(L)$: the height of L equals $|P| + 1$,

L is Boolean iff P is an antichain, and L is a chain iff P is a chain. See [DP90, Grä03] for proofs.

21.5 Problems

21.1. Give an example of a nonempty binary relation which is symmetric and transitive but not reflexive.

21.2. Show that if P and Q are posets defined on set X , then so is $P \cap Q$.

21.3. Draw the Hasse diagram of all natural numbers less than 10 ordered by the relation *divides*.

21.4. Show that if C_1 and C_2 are down-sets for any poset $(E, <)$, then so is $C_1 \cap C_2$.

21.5. Show that the join and meet operators are associative, commutative, and satisfy the absorption laws:
 $a \sqcup (a \sqcap b) = a$ and $a \sqcap (a \sqcup b) = a$.

21.6. Show that in a finite lattice, an element is join-irreducible iff it has only one lower cover.

21.7. Show that for a finite distributive lattice, L is Boolean iff $J(L)$ is an antichain.

21.6 Bibliographic Remarks

The first book on lattice theory was written by Garrett Birkhoff [Bir40, Bir48, Bir67]. The reader will find a discussion of origins of lattice theory and an extensive bibliography in the book by Grätzer [Grä71, Grä03]. The book by Davey and Priestley [DP90] provides an easily accessible account of the field. A treatment with an emphasis on computer science applications is given by Garg [Gar15]. Other books include those by Caspard, Leclerc, and Monjardet [CLM12], and Roman [Rom08].

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ALGORITHMS THROUGH THE LENS OF LATTICE-LINEAR PREDICATES

This book presents the classical algorithms of computer science from two complementary perspectives. The first is the traditional one—greedy choices, divide-and-conquer recursions, dynamic-programming tables, augmenting paths. The second recasts each problem as the search for an extremal element of a distributive lattice that satisfies a lattice-linear predicate. From this single, unifying viewpoint, the book derives algorithms for sorting, graph traversal, shortest paths, minimum spanning trees, dynamic programming, network flow, bipartite matching, stable matching, housing allocation, assignment, satisfiability, number theory, and more.

The lattice-search formulation makes correctness proofs shorter, exposes natural parallelism, and turns algorithmic design into a systematic activity: identify the lattice, write down the predicate, check lattice-linearity, and read off the advance rule. Every algorithm in the book ships as runnable code on a companion website—in Java, Python, C++, and JavaScript—together with interactive demos that step through each LLP loop on small concrete instances. A solution manual is available to instructors on request.

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