Coping with NP-hardness

You proved your problem of interest is NP-hard

Now what?

In practice, you still need to solve problems

The high level goal is to get good enough solutions by any means necessary

“Good enough” = approximation algorithms

You could simply attack the problem of designing algorithms head on – invent something ad-hoc

Or you could try out some algorithmic paradigms – general methods that seem to work well for other problems, so they may be adapted to work for your problem too
Coping with NP-hardness

Do you know who else needs to routinely solve NP-hard problems?

**NATURE!**

Borrow some inspiration for algorithmic paradigms from it

One of its most successful algorithmic paradigms is evolution:

Incremental process of

(1) introducing small mutations,
(2) trying them out, and
(3) keeping them if they work well.
Local Search Algorithmic Paradigm

Start with some solution $S$
While there is a “better” solution $S'$ in the local neighborhood of $S$
update $S$ to $S'$

possible small mutations $\Leftrightarrow$ local neighborhood
trying mutations out $\Leftrightarrow$ enumerating the entire local neighborhood
keeping mutations if they work well $\Leftrightarrow$ updating $S$ to $S'$ if $S'$ is better than $S$

Set of all possible solutions is called solution space or state space

Not to be confused with so-called “genetic algorithms” that mutate several solutions simultaneously and mutations might depend on several solutions
Local Search Applications

(1) As an approximation algorithmic paradigm with provable guarantees

(2) As an algorithmic heuristic without provable guarantees but that tends to work extremely well in practice

Examples:

Maximum flow (exact algorithms)
Finding independent sets in restricted classes of graphs (exact algorithms)
Maximum satisfiability, graph partitioning (provable guarantees)
Travelling salesman problem, VLSI design (heuristic)
Programming computer to play chess (heuristic)
As a preprocessing step for exact algorithms, etc.
Plan

(1) Exact Max-2-SAT (was briefly covered last lecture)
(2) Programming Computer to Play Chess
(3) Travelling Salesman Problem Heuristics
(4) Coping with Local Optima
Example: Exact Max-2 SAT

INPUT: CNF formula $F$
- $n$ variables $x_1, x_2, \ldots, x_n$
- $m$ clauses $C_1 \land C_2 \land \cdots \land C_m$
- each clause has exactly 2 variables

OUTPUT: assignment $\tau$ to the variables that satisfies as many clauses as possible

This problem is NP-hard to solve exactly

Solution space = \{all possible assignments to variables\}. It’s size is $2^n$
Exact Max-2-SAT

Example $n = 3$, $m = 6$

$$(\neg x_1 \lor x_2) \ (\neg x_2 \lor x_3) \ (\neg x_3 \lor \neg x_1) \ (x_1 \lor x_3) \ (\neg x_3 \lor x_2) \ (\neg x_2 \lor x_1)$$

Most natural local search algorithm:

Start with some assignment

Try flipping any bit in that assignment and see if it increases number of satisfied clauses

If so, flip the bit and repeat; otherwise terminate
**Exact Max-2-SAT**

Example $n = 3$, $m = 6$

$$(\neg x_1 \lor x_2) (\neg x_2 \lor x_3) (\neg x_3 \lor \neg x_1) (x_1 \lor x_3) (\neg x_3 \lor x_2) (\neg x_2 \lor x_1)$$

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## Exact Max-2-SAT

Example $n = 3$, $m = 6$

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Exact Max-2-SAT

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# Exact Max-2-SAT

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Landscape of Solution Space

Global max can also be flat
Approximation Guarantee

If local search algorithm terminates it ends up in a local optimum

How good is that optimum?

Worst-case approximation guarantee (locality gap) is the ratio between smallest local optimum and global optimum (for maximization problem)

Largest such ratio is the locality gap
Approximation Guarantee of Local Search for Exact MAX-2-SAT

We shall prove that our local search algorithms achieves 2/3 approximation ratio

**Proof:** Let $\tau$ be a local optimum

- $S_0$ - clauses not satisfied by $\tau$
- $S_1$ - clauses satisfied by exactly one literal by $\tau$
- $S_2$ - clauses satisfied by exactly two literals by $\tau$

Eg. $(\neg x_1 \lor x_2) \ (\neg x_2 \lor x_3) \ (\neg x_3 \lor \neg x_1) \ (x_1 \lor x_3) \ (\neg x_3 \lor x_2) \ (\neg x_2 \lor x_1)$

Local opt $\tau$: $x_1 = 1$, $x_2 = 1$, $x_3 = 1$

- $S_0 = \{ (\neg x_3 \lor \neg x_1) \}$
- $S_1 = \{ (\neg x_1 \lor x_2), (\neg x_2 \lor x_3), (\neg x_3 \lor x_2), (\neg x_2 \lor x_1) \}$
- $S_2 = \{ (x_1 \lor x_3) \}$
Approximation Guarantee of Local Search for Exact MAX-2-SAT

Let $\tau$ be a local optimum
- $S_0$ - clauses not satisfied by $\tau$
- $S_1$ - clauses satisfied by exactly one literal by $\tau$
- $S_2$ - clauses satisfied by exactly two literals by $\tau$

Our algorithm finds an assignment satisfying $|S_1| + |S_2|$ clauses.
Optimum is clearly less than the total number of clauses $|S_0| + |S_1| + |S_2|$

Thus, we need to show $\frac{|S_1| + |S_2|}{|S_0| + |S_1| + |S_2|} \geq \frac{2}{3}$. Alternatively, show $\frac{|S_0|}{|S_0| + |S_1| + |S_2|} \leq \frac{1}{3}$.
Approximation Guarantee of Local Search for Exact MAX-2-SAT

Clause involves variable $x_i$ if either $x_i$ or $\neg x_i$ occurs in it

$A_i$ - set of clauses in $S_0$ involving $x_i$

$B_i$ - set of clauses in $S_1$ involving $x_i$ and satisfied by $x_i$ in $\tau$

$C_i$ - set of clauses in $S_2$ involving $x_i$

Local optimum means $|A_i| \leq |B_i|$ for all $i$

Also we have $\sum_i |A_i| = 2|S_0|$ and $\sum_i |B_i| = |S_1|$

Hence $2|S_0| \leq |S_1|$, thus

$$\frac{|S_0|}{|S_0|+|S_1|+|S_2|} \leq \frac{|S_0|}{|S_0|+2|S_0|+|S_2|} \leq \frac{|S_0|}{3|S_0|} \leq \frac{1}{3}$$

QED
Approximation Guarantee of Local Search for Exact MAX-2-SAT

Our simple local search finds a 2/3-approximation.

This analysis and algorithm can be generalized to the case where you associate arbitrary weights with clauses and the goal is to satisfy a set of clauses of maximum weight.

In our algorithm we said a modification of an assignment is improving if it increases the number of satisfied clauses

Better algorithm can be obtained with a different criterion for “improving” step (non-oblivious local search)

See Allan’s slides for more information!
Other Interesting Applications

Claude Shannon (1916-2001)
One of the mathematical giants of the 20th century
Single-handedly introduced information theory
“A Mathematical Theory of Communication” (1948)

Less known paper
“Programming a computer for playing chess” (1950)
Programming a Computer to Play Chess

Shannon’s proposal:

Encode a valid chess configuration by a 2-dimensional array

Define a “goodness” function

- maps each chess configuration to a number
- the higher the number the better configuration is for the player, e.g., if your king is checked in this configuration its not good. If you have a lot of pieces as opposed to your opponent, the configuration is good. Can make this precise in many ways

Computer program considers various valid moves and resulting configurations, and picks a better one according to the “goodness” function
Programming a Computer to Play Chess

Shannon’s proposal is a local search algorithm!

State space: all possible valid chess configurations

Neighborhood: all chess configurations reachable from the current one in one step

Shannon also suggested extensions and improvements:

- Larger neighborhoods – chess configs resulting from several moves
- Certain moves are certainly bad and should not be explored further (pruning)
- Suggested that machine learning techniques should guide the choice of a “goodness” function and pruning strategies
Travelling Salesman Problem (TSP)

Complete graph

Positive weights on every edge
- Symmetric case: \( w(i, j) = w(j, i) \)
- Triangle inequality: \( w(i, j) \leq w(i, k) + w(k, j) \)
- Euclidean distance, i.e. nodes are located on a plane, weight of edge = dist

Find shortest tour (visit every node exactly once)

Madiera Island (west of Morocco)
Find a min distance tour visiting every city
Graph Representation

Not all edges are shown: there is an edge between every pair of vertices; weight of an edge = distance
Example of a Tour
Local Search Heuristic

Solution space = \{all possible tours, i.e., permutations, visiting \( n \) cities\}
Solution space size is \((n - 1)!\)

Pick a solution, do local improving steps until get stuck in a local opt

But what are local improving steps?

2-opt idea: take two edges out, put in two other edges in

![Diagram showing 2-opt improvement]

\[ \text{Original tour} \xrightarrow{2-opt\, improvement} \text{Improved tour} \]
Local Search for TSP

3-opt: take out 3 edges, put in 3 new edges

More generally, k-opt for any k is known as Lin-Kernighan heuristic

Bigger k bigger neighborhood!
Example

Albuquerque  Amarillo  Wichita

Tulsa  Little Rock

Dallas

El Paso

San Antonio  Houston
Example
Example
Example
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Example
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Example
### Local Search for TSP, Exact Methods, Some Benchmarks

<table>
<thead>
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<th>Year</th>
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<tbody>
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<td>1954</td>
<td>49</td>
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<td>1971</td>
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<td>1975</td>
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<td>1994</td>
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<td>1998</td>
<td>13,509</td>
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<tr>
<td>2001</td>
<td>15,112</td>
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</table>

Many of these results rely on Lin-Kernighan heuristic.

Thus, nowadays people can solve TSP exactly with tens of thousands of vertices, but the problem is NP-hard!

This is a testament to how good these heuristics are in practice.
Why Heuristic?

Well defined algorithms, so
- What is the overall running time?
- What is its approximation ratio?

In spite of large interest in local search methods for TSP, no one knows an answer to either of those questions.
Dealing with Local Optima

Algorithm gets stuck in local optima

To improve performance want methods to push algorithm out of local optima

Simplest methods:

**Random restarts** – when algorithm gets stuck in a local optimum, restart it at a random initial solution

Add **more randomization** – when algorithms picks which solution to move to (locally), pick any improving solution with some probability

Logic for more randomization is that maybe your algorithm gets stuck in a bad local optimum due to **systematic** choices it makes during the runtime
Dealing with Local Optima

Taking idea of randomization further:

With some small probability, allow your algorithm to make a local step that is \textit{worse} than the current solution

Vary this probability with time

\begin{itemize}
  \item Initially this probability is rather large to allow your algorithm to explore solution space widely
  \item Later this probability of taking a non-improving step decreases more and more allowing the algorithm to narrow down on a promising region in the solution space
\end{itemize}
Dealing with Local Optima

Previous slide describes **simulated annealing** technique

```
let s be any starting solution
repeat
    randomly choose a solution $s'$ in the neighborhood of $s$
    if $\Delta = \text{cost}(s') - \text{cost}(s)$ is negative:
        replace $s$ by $s'$
    else:
        replace $s$ by $s'$ with probability $e^{-\Delta/T}$.
```

It is usually very successful in practice outperforming simple local search, but it is incredibly difficult to analyze
NEW TOPIC:
DUAL FITTING
What is Dual Fitting?

Technique for analyzing approximation ratio of an algorithm

In many cases, the algorithm is greedy

The algorithm is interpreted as building a primal solution

In the analysis, you build a dual solution of similar value
Set Cover Problem

We are given a universe of $n$ elements $\{1, 2, \ldots, n\}$ and $m$ subsets of the universe $S_1, S_2, \ldots, S_m$.

A set cover is a collection of these sets that covers each element at least ones.

The goal is to find a cover of minimum size. This is NP-hard.
Greedy Algorithm

Pick a subset that covers the most number of uncovered elements.
Repeat until all elements are covered.
Greedy Algorithm

Pick a subset that covers the most number of uncovered elements.

Repeat until all elements are covered.
Greedy Algorithm

Pick a subset that covers the most number of uncovered elements.
Repeat until all elements are covered.
Greedy Algorithm

Pick a subset that covers the most number of uncovered elements.
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Greedy Algorithm

Pick a subset that covers the most number of uncovered elements.
Repeat until all elements are covered.

Our goal: prove that this algorithm is within $\ln n + O(1)$ of the optimal value.
An Integer Program for Set Cover

$x_s$ is an indicator variable: 1 if $S$ is picked in the set cover and 0 otherwise

$$\min \sum_{S} x_S$$

$$\forall e: \sum_{S: e \in S} x_S \geq 1$$

$$\forall S: x_S \in \{0, 1\}$$

Every element $e$ should be included in at least one set.
LP Relaxation of Set Cover

Replace the integrality constraint $x_S \in \{0,1\}$ by a linear constraint $0 \leq x_S \leq 1$

$$\min \sum_{S} x_S$$
$$\forall e: \sum_{S: e \in S} x_S \geq 1$$
$$\forall S: x_S \in \{0,1\}$$

$$\min \sum_{S} x_S$$
$$\forall e: \sum_{S: e \in S} x_S \geq 1$$
$$\forall S: x_S \geq 0$$
Dual LP

PRIMAL

\[ \min \sum_{S} x_{S} \]
\[ \forall e: \sum_{S: e \in S} x_{S} \geq 1 \]
\[ \forall S: x_{S} \geq 0 \]

DUAL

\[ \max \sum_{e \in U} y_{e} \]
\[ \forall S: \sum_{e \in S} y_{e} \leq 1 \]
\[ \forall e: y_{e \in U} \geq 0 \]
Analysis Goal: Sandwich OPT between Algorithm’s Soln and a Feasible Dual Soln

Dual Solution (we will construct it)

Objective value

Approximation ratio

Dual feasible

primal feasible

LP Solution

OPT

Greedy Solution
Consider two entities: the algorithm and the prover

\[
\begin{align*}
\text{PRIMAL} & \quad \text{DUAL} \\
\min \sum_{S} x_{S} & \quad \max \sum_{e \in U} y_{e} \\
\forall e: \sum_{S: e \in S} x_{S} \geq 1 & \quad \forall S: \sum_{e \in S} y_{e} \leq 1 \\
\forall S: x_{S} \geq 0 & \quad \forall e: y_{e} \geq 0
\end{align*}
\]

The algorithm can be viewed as constructing a feasible solution to PRIMAL. The prover will be constructing a feasible solution to the DUAL.

If the prover succeeds at constructing a feasible solution within \( \alpha \) fraction of the primal solution, it means that the approximation ratio is \( \alpha \).

Prover is completely imaginary – we play the role of the prover.
The Algorithm vs the Prover

<table>
<thead>
<tr>
<th>PRIMAL</th>
<th>DUAL</th>
<th>Consider a particular step in the algorithm</th>
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</thead>
<tbody>
<tr>
<td>(\min \sum_{S} x_S)</td>
<td>(\max \sum_{e \in U} y_e)</td>
<td>- Algorithm selects a new set (S)</td>
</tr>
<tr>
<td>(\forall e: \sum_{S : e \in S} x_S \geq 1)</td>
<td>(\forall S: \sum_{e \in S} y_e \leq 1)</td>
<td>- This set covers (k) new elements</td>
</tr>
<tr>
<td>(\forall S: x_S \geq 0)</td>
<td>(\forall e: y_e \geq 0)</td>
<td>The algorithm can be viewed as constructing a feasible solution to PRIMAL: (x_S = 1)</td>
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<td>Note: this adds value 1 to the primal objective</td>
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As a prover, we need to create some dual soln, i.e., assign \(y_e\) values to all newly covered elements \(e\).
First try: assign \(y_e = \frac{1}{k}\)
The good: we added the total of \(k(1/k) = 1\) value to the dual objective
The bad: our dual vars violate dual constraints (set \(S\) can have prev. covered elts)
Making Prover’s Dual Solution Feasible

We set \( y_e = \frac{1}{k} \) where \( k \) is the number of new elements covered at the time element \( e \) was covered.

Consider an arbitrary set \( S \). We would like to enforce \( \sum_{e \in S} y_e \leq 1 \).

Assume without loss of generality that if greedy covers \( e_j \) before \( e_k \) then \( j < k \).

When greedy picked a set to cover \( e_j \), it could also have picked \( S \) and covered \(|S| - j + 1\) new elements.

So \( y_e \leq \frac{1}{(|S| - j + 1)} \)

\[ \sum_{e \in S} y_e \leq \left(1 + \frac{1}{2} + \frac{1}{3} + \cdots + \frac{1}{|S|}\right) \leq \ln|S| + O(1) \]
Making Prover’s Dual Solution Feasible

We set $y_e = 1/k$ where $k$ is the number of new elements covered at the time element $e$ was covered.

When variables are set as above, we have

$$\sum_{e \in S} y_e \leq (1 + \frac{1}{2} + \frac{1}{3} + \cdots + \frac{1}{|S|}) \leq \ln|S| + O(1) \leq \ln n + O(1)$$

But we want $\sum_{e \in S} y_e \leq 1$

So…? Rescale the variables! Set $\hat{y}_e = \frac{y_e}{\ln n + O(1)}$

Then the variables $\hat{y}_e$ form a feasible dual solution.
Making Prover’s Dual Solution Feasible

We set $y_e = 1/k$ where $k$ is the number of new elements covered at the time element $e$ was covered.

Setting $\hat{y}_e = \frac{y_e}{\ln n + O(1)}$ made the prover’s solution feasible.

What happened to the objective value?

Recall that adding $y_e$ for all $e$ gave exactly the same value as algo’s primal soln

Hence, adding $\hat{y}_e$ for all $e$ gives a solution of value within $\ln n + O(1)$ of primal’s.

This finishes the proof, since opt is sandwiched between feasible dual and greedy solution.
NEW TOPIC:
RANDOMIZED ALGORITHMS
Deterministic Algorithms

Goal:
Correctness – algo always solves the problem correctly
Efficiency – algo always solves the problem quickly, e.g., polytime
Randomized Algorithms

Behavior of the algorithm can vary even for a given fixed input!

Goals:
Correctness – algo often solves the problem correctly (for every input!)
Efficiency – algo often solves the problem quickly, e.g., expected polytime
Probabilistic Analysis of Algorithms

Goals:

Correctness – algo solves the problem correctly on most inputs

Efficiency – algo solves the problem quickly, e.g., polytime, on most inputs
Reasons for Studying Randomized Algorithms

Simplicity
Performance

For many problems, a randomized algorithm is either the simplest known, the fastest known, or both.

Turning randomized algorithms into deterministic is called “derandomization”.

A randomized algorithm is often discovered first, then it is derandomized, but in the process either runtime or approximation ratio might suffer.

In certain cases, people don’t know if an algorithm can be derandomized.

Lastly, in some problem settings (e.g., sublinear time algorithms) randomization is provably necessary!
Applications of Randomized Algorithms

Number theory (primality testing)
Data structures (hashing)
Algebraic identities (polynomial and matrix identity verification)
Mathematical programming (rounding linear program relaxations)
Counting and enumeration (approximating counting of perfect matchings)
Networking (deadlock avoidance)
Probabilistic method for proving theorems
etc...
This Lecture: Randomized Algos

Plan:
(1) Probability overview
(2) Types of randomized algorithms
(3) Polynomial identity testing
Probability Theory Overview

Mathematical theory of chance

We will only consider finite discrete probability spaces

**Probability space** is a pair \((\Omega, p)\), where

- \(\Omega\) – sample space, \(|\Omega| < \infty\)
- \(p: \Omega \rightarrow [0,1]\) – probability distribution, i.e., a function such that

\[
\sum_{\omega \in \Omega} p(\omega) = 1
\]
Probability Theory Overview

**Event** is a subset of the sample space, $A \subseteq \Omega$.

How many distinct events are there? Answer: $2^{|\Omega|}$ (prove it!)

Probability distribution extends to events:

Let $A \subseteq \Omega$ then $p(A) = \sum_{\omega \in \Omega} p(\omega)$

$\omega \in \Omega$ is called an **elementary event**
Probability Theory Overview

Example: throw a fair six-sided dice once

Sample space $\Omega = \{1,2,3,4,5,6\}$

Probability distribution $p(1) = p(2) = p(3) = p(4) = p(5) = p(6) = \frac{1}{6}$

The above distribution is called **uniform**

More generally, uniform distribution is such that $p(\omega) = 1/|\Omega|$

Event $A$: “even number shows up on the top of the dice”

Formally, $A = \{2,4,6\}$, so $p(A) = p(2) + p(4) + p(6) = \frac{3}{6} = \frac{1}{2}$
Probability Theory Overview

Back to general definitions

Fix probability space \((\Omega, p)\)

**Real-valued random variable** is a function \(X: \Omega \rightarrow \mathbb{R}\)

It is a misnomer: random variable is

- Neither random
- Nor a variable
Real-valued random variable “collapses” elementary events into groups and associates real numbers with those groups.

These groups are subsets of sample space, i.e., events, which we denote by “$X = value$”
Probability Theory Overview

For example, consider a throw of a fair six-sided dice
Let $X$ be the random variable defined as follows:
   If the number that comes up is odd, $X$ is equal to that number
   If the number that comes up is even, $X$ is equal to half of that number
In other words,

\[
\begin{align*}
X(1) &= 1 \\
X(2) &= 2/2 = 1 \\
X(3) &= 3 \\
X(4) &= 4/2 = 2 \\
X(5) &= 5 \\
X(6) &= 6/2 = 3
\end{align*}
\]
Events corresponding to outcomes of random variable:

\[ P(X = 1) = P(\{1,2\}) = p(1) + p(2) = \frac{2}{6} = \frac{1}{3} \]

\[ P(X = 2) = P(\{4\}) = p(4) = \frac{1}{6} \]

\[ P(X = 3) = P(\{3,6\}) = p(3) + p(6) = \frac{2}{6} = \frac{1}{3} \]

\[ P(X = 5) = P(\{5\}) = p(5) = \frac{1}{6} \]

**NOTE:**

\[ P(X = 1) + P(X = 2) + P(X = 3) + P(X = 5) = 1 \]
Thus, we get an auxiliary probability space!

Sample space
outcomes of random variable $X$, e.g., \{1,2,3,5\}

Probability distribution
probability of outcomes, e.g.,

$$p(1) = p(3) = \frac{1}{3}, p(2) = p(5) = \frac{1}{6}$$
Probability Theory Overview

Start with a probability distribution here.

Pass it “through” a random variable.

Get a probability distribution here.

distribution of $X$
Probability Theory Overview

**Expected value** of a random variable $X$ is defined as

$$E(X) = \sum_{\omega \in \Omega} X(\omega)p(\omega)$$

Equivalently,

$$E(X) = \sum_i i P(X = i)$$

Exercise: prove the equivalence

Observe that second definition does not require you to know the original distribution, only distribution of $X$
Probability Theory Pitfalls

Probability theory models something very natural – chance!

It is easy to think that you sort-of-understand it, and dive into problems right away, start calculating answers, and getting results (sometimes even correct).

Without understanding basic definitions you won’t get very far – even beginner problems would seem incomprehensible.

You should be very clear what are probability spaces, events, random variables, expectations, etc.

To keep yourself in check, get into habit of asking yourself “What is the sample space? Is it a set of numbers/names/dates/apples? What is the size of the sample space? What is the distribution? Is it uniform?”
Examples of Pitfalls

Events have probabilities!
Random variables have expectations!
“Expectation of event A” does not make sense! Never say it!
“Probability of random variable X” does not make sense! Never say it!
Probabilities can never be negative!
Probabilities can never be greater than 1!
Things You Should Know to Understand and Analyze Randomized Algorithms

Conditional probabilities
Independent random variables
Conditional expectations
Law of complete probability
Moments of random variables
Standard discrete distr: uniform, Bernoulli, Binomial, Geometric, Poisson, ...
Standard continuous distr: uniform, Gaussian, exponential, ...
...

Three Pillars of Probability Theory

- Linearity of Expectation
- Union Bound
- Chernoff Bound

Deceptively simple!
Incredibly powerful!
If you understand and practice these techniques, you would be able to solve MANY MANY MANY MANY MANY MANY probability theory problems!
Can think of randomness as an infinite supply of random bits, e.g.,
1101000010100011 1111101010000001011111001100111010110101...

Algorithm accesses them sequentially, so if it runs in finite time, it only uses finitely many of the bits.

Polytime algorithm uses polynomially many random bits
Randomness in Randomized Algorithms

Suppose that on input \( x \) your algorithm uses exactly \( N \) random bits (WLOG)

What is the sample space?
\[ \Omega = \{0,1\}^N \]

What is the probability distribution?
\( p \) is uniform, i.e., for any random string \( r \in \Omega \) we have \( p(r) = 1/2^N \)
Randomness in Randomized Algorithms

But why do we only allow independent coin flips?

What if our algorithm requires a different distribution?

Turns out you can create samples from most other distributions (either exactly or approximately) we care about

For example, if you wanted to get a uniform random number between [0,1] to 5 bits of accuracy, you can take 5 coin flips, e.g., 01101, and put a zero followed by dot after it:

0.01101 – interpret it as a binary fractional number
Randomness in Randomized Algorithms

Another example:

Start with a probability space of \( n \) flips of a fair coin

Define a random variable \( X \) to be the number of times heads come up

Then \( X \) has binomial distribution

\( X \) can take on values 0, 1, 2, ..., \( n \)

For \( k \in \{0,1,\ldots,n\} \) we have

\[
P(X = k) = \binom{n}{k}2^{-n}
\]

Suppose in designing an algorithm you need a sample from binomial distribution with parameter \( n = 5 \)

Take 5 random bits from the random string and add them together
Randomness in Randomized Algorithms

More generally, any random variable whose cumulative distribution function is efficiently invertible can be efficiently sampled.

Bottom line:

with just uniform random bits we can approximately simulate almost all other distributions we care about.
Random Variables of Interest

\( x \) denotes input, \( r \) denotes a random string

Main random variables of interest:

\( T(x, r) = \) runtime of your algorithm on input \( x \) with randomness \( r \)

\( C(x, r) = 1, \) if your algorithm gives the right output on input \( x \) when it uses randomness \( r; \) 0, otherwise.
Desired Properties

\[ T(x, r) = \text{runtime of your algorithm on input } x \text{ with randomness } r \]
\[ C(x, r) = 1, \text{ if your algorithm gives the right output on input } x \text{ when it uses randomness } r; 0, \text{ otherwise.} \]

(1) Runtime

\[ T(x, r) \leq \text{poly}(|x|) \text{ for every input } x \text{ and random string } r \]

Weaker requirement: \( E_r(T(x, r)) \leq \text{poly}(|x|) \) for every input \( x \)

(2) Probability of success

\[ P_r(C(x, r) = 1) \geq 9/10 \text{ for every input } x \]
Monte Carlo vs Las Vegas Algorithms

**Monte Carlo algorithms** always run for a small number of steps (e.g., \( T(x, r) \leq \text{poly}(|x|) \)) and produce a correct answer with high enough probability (e.g., \( P_r(C(x, r) = 1) \geq 9/10 \)).

**Las Vegas** algorithms run for expected small number of steps (e.g., \( E_r(T(x, r)) \leq \text{poly}(|x|) \)), but always produce a correct answer (i.e., \( P_r(C(x, r) = 1) = 1 \)).

Thus, Las Vegas algorithms for some values of randomness might run for much longer time.
Monte Carlo vs Las Vegas

Monte Carlo algs
- Also called an algo with 2-sided error
- Probability of success in the definition is not important, as long as it is $\frac{1}{2} + \varepsilon$ for some constant $\varepsilon > 0$

Las Vegas algs
- Also called an algo with 0-sided error
Example of a Las Vegas Algorithm

Quicksort

(1) to sort an array of elements, pick a random number in the array - pivot

(2) put all numbers less than the pivot in one array, all numbers bigger than the pivot in another array

(3) recursively sort the two arrays

Note that

Quicksort always produces the right answer

Its expected runtime is $O(n \log n)$

Sometimes it might run for $\Omega(n^2)$ steps if it was unlucky in pivot selections
First Serious Example: Polynomial Identity Testing

Polynomial of degree $d$ in one variable $x$ is a function of the form

$$p(x) = a_d x^d + a_{d-1} x^{d-1} + \cdots + a_1 x + a_0$$

where $a_i \in \mathbb{Q}$ (for example) and $a_d \neq 0$
Polynomial Identity Testing

Two polynomials \( p(x) \) and \( q(x) \) are equal if all the coefficients are the same.

Sometimes, a polynomial is not represented in the above form right away.

A polynomial could be represented by an implicit formula, e.g.:

\[
q(x) = (x - 3)(10x + 5)(3x^5 - 3x + 8) \ldots (x + 48) + (x^{27} - 3)(2x - 8)
\]

We know it is a polynomial, but without opening up the brackets, we don’t know its coefficients! There could be nontrivial cancellations.

Note that when \( q(x) \) is given implicitly as above, its degree is at most the size of such an implicit representation. In what follows, we assume \( d \) is both a bound on degree and size of representation.
Polynomial Identity Testing

Given two polynomials $p(x)$ and $q(x)$ in such an implicit form, we would like to test if they are the same polynomial.

Suppose that the degree of each polynomial is at most $d$.

We could open up the brackets and simplify the polynomials, but it is too much work – fast and complicated algorithms for doing this require time $\Omega(d \log d)$.

Can we do better?

Can we check if two polynomials given implicitly are the same in time $O(d)$ and without opening up the brackets?
Polynomial Identity Testing to Polynomial Zero Testing

Instead of checking $p(x) = q(x)$ we can check if $p(x) - q(x) = 0$.

If $p(x)$ and $q(x)$ are degree at most $d$ polynomials then $p(x) - q(x)$ is also degree at most $d$ polynomial.

So, now the problem becomes

Given a polynomial $p(x)$ of degree at most $d$ represented implicitly, check if it is the identically 0 polynomial without opening up the brackets and simplifying all the coefficients.
Problem: given a polynomial \( p(x) \) of degree at most \( d \) represented implicitly, check if it is the identically 0 polynomial without opening up the brackets and simplifying all the coefficients

Key observations:

(1) polynomial \( p(x) \) of degree \( d \) has at most \( d \) roots – those numbers where the polynomial evaluates to zero

(2) evaluating a polynomial of degree at most \( d \) can be done with \( O(d) \) basic arithmetic operations (addition, subtraction, multiplication) – Exercise! (recall that \( d \) is both the bound on the implicit representation and on the degree)
Randomized Polynomial Zero Testing

Problem: given a polynomial $p(x)$ of degree at most $d$ represented implicitly, check if it is the identically 0 polynomial without opening up the brackets and simplifying all the coefficients.

Randomized algorithm:

- Pick a number $r$ uniformly at random from $\{1, 2, \ldots, 10d\}$
- Evaluate $p(r)$. If it returns non-zero, output “NON-ZERO POLY”; otherwise, output “ZERO POLY”
Analysis of Randomized Zero Testing Algo

Runtime: $O(d)$ – always

Correctness:

if $p(x)$ is the zero polynomial, it always evaluates to zero, so our algorithm is always correct!

If $p(x)$ is non-zero, it has at most $d$ roots, so at most $d$ out of $10d$ first numbers evaluate to zero. With probability $9/10$ we select a number on which $p(x)$ does not evaluate to zero, so we output the right answer with probability $9/10$

This is a Monte Carlo algorithm with one-sided error.

Error probability can be boosted by either repetition or selecting a bigger range
Boosting the Probability of Success

Repeat the algorithm $k$ times with fresh randomness

Output “ZERO” only if all runs output “ZERO”

Probability of success?

If $p(x)$ is the zero polynomial, we always output “ZERO”

If $p(x)$ is not the zero polynomial, we make an error if and only if ALL runs make an error. Since runs are independent, probabilities multiply:

Probability that a single run makes an error is $1/10$, then the probability that all runs make errors is $1/10^k$. 
Cosmic Rays

From Wiki:

“Cosmic rays are high-energy radiation, mainly originating outside the Solar System and even from distant galaxies. Upon impact with the Earth's atmosphere, cosmic rays can produce showers of secondary particles that sometimes reach the surface.”

When they hit a RAM module on your laptop they can corrupt main memory. Depending on where cosmic rays hit, your program can crash, or simply continue execution with erroneous data.
Boosting the Probability of Success

If your program runs for 1 minute and occupies 20 MB of RAM, IBM estimates that there is roughly $1/10^7$ chance that cosmic rays will hit the RAM module and introduce errors to your program.

If you run the repeated zero polynomial testing algorithm for $k = 10$ iterations, the probability of error is $1/10^{10}$, but it still runs in $10 \mathcal{O}(d) = \mathcal{O}(d)$ time.

For all practical purposes, the result of this randomized algorithm is as dependable as any other computation that you run on your computer

BUT the algorithm is super simple and super efficient!
Extending the Algorithm to Multiple Vars

Example: polynomial in 3 variables $x_1, x_2, x_3$

$$p(x_1, x_2, x_3) = 10x_1^3x_2^4x_3 + 8x_1x_2x_3^4 - x_1x_3 + x_2 - 6$$

Total degree is 8 (maximum sum of degrees in a single term)

More generally, we will consider polynomials in many variables

These polynomials can also be represented implicitly

We want to see when two polynomials in several variables are the same

Alternatively, when a single polynomial in several variables is the zero polynomial
Nontrivial Example

Consider a square $n \times n$ matrix whose entries are variables $x_{ij}$:

$$ M = \begin{pmatrix} x_{11} & \cdots & x_{1n} \\ \vdots & \ddots & \vdots \\ x_{n1} & \cdots & x_{nn} \end{pmatrix} $$

The determinant of $M$ is defined as follows:

$$ \det(M) = \sum_{\sigma} (-1)^{sgn(\sigma)} x_{1\sigma(1)} x_{2\sigma(2)} \cdots x_{n\sigma(n)} $$

Where $\sigma$ ranges over all permutations of \{1,2, ..., $n$\}

Thus, the determinant is a polynomial in $n^2$ variables of total degree $n$ and with $n!$ terms
Nontrivial Example

$$\text{det}(M) = \sum_{\sigma} (-1)^{sgn(\sigma)} x_{1\sigma(1)} x_{2\sigma(2)} \ldots x_{n\sigma(n)}$$

$n^2$ variables, total degree $n$, $n!$ Terms

For $n = 100$, there is not enough material in the observable universe to create the paper necessary to write down this polynomial explicitly

Nonetheless, for a particular setting of values to variables $x_{ij}$ we can evaluate this polynomial efficiently using Gaussian Elimination!

This is an implicit representation of a polynomial: evaluation is quick, but we cannot expand the polynomial coefficients
Multivariate Polynomial Zero Testing

Polynomial Zero Testing with one variable required a bound on the number of roots of a polynomial of degree \( n \)

Thus, we need a generalization of such a bound to several variables and total degree

**Schwartz-Zippel Lemma.** Let \( q(x_1, x_2, \ldots, x_n) \) be a non-zero multivariate polynomial of total degree \( d \) over \( \mathbb{Q} \). Let \( S \) be a finite subset of \( \mathbb{Q} \), and let \( r_1, r_2, \ldots, r_n \) be chosen independently and uniformly at random from \( S \). Then

\[
P(q(r_1, r_2, \ldots, r_n) = 0) \leq \frac{d}{|S|}
\]
Multivariate Polynomial Zero Testing

**INPUT:** polynomial $q$ on $n$ variables of total degree $d$ given implicitly

**OUTPUT:** yes if $q$ is identically zero polynomial, no otherwise

**Algorithm:**

Pick $r_1, r_2, ..., r_n$ uniformly at random from \{1, 2, ..., 10d\}

Output yes if and only if $q(r_1, r_2, ..., r_n) = 0$

**Analysis:**

If $q$ is zero then we always output yes

If $q$ is nonzero, by Schwarz-Zippel, we output no with probability at least $9/10$
Multivariate Polynomial Zero Testing

Algorithm:

Pick $r_1, r_2, \ldots, r_n$ uniformly at random from $\{1, 2, \ldots, 10d\}$

Output yes if and only if $q(r_1, r_2, \ldots, r_n) = 0$

As before, we can boost the probability of success to be almost 1 by repetition

Example of a problem for which the best known algorithm is randomized

Can this algorithm be derandomized? Huge open question in theoretical CS
Interesting Application

The Vandermonde matrix $M(x_1, x_2, \ldots, x_n)$

$$
\begin{pmatrix}
1 & x_1 & x_1^2 & \ldots & x_1^n \\
1 & x_2 & x_2^2 & \ldots & x_2^n \\
\vdots & \vdots & \vdots & \ddots & \vdots \\
1 & x_n & x_n^2 & \ldots & x_n^n \\
\end{pmatrix}
$$

Vandermonde’s identity: $\det(M) = \prod_{j<i} (x_i - x_j)$ (EXERCISE – prove it!)
Vandermonde’s Identity

\[
\begin{vmatrix}
1 & x_1 & x_1^2 & \ldots & x_1^n \\
1 & x_2 & x_2^2 & \ldots & x_2^n \\
\vdots & \vdots & \vdots & \ddots & \vdots \\
1 & x_n & x_n^2 & \ldots & x_n^n \\
\end{vmatrix} = \prod_{j<i}(x_i - x_j)
\]

This is polynomial identity testing!

Cannot write down and simplify the poly on the left – it has \(n!\) Terms

**BUT** can evaluate it efficiently using Gaussian elimination

Thus, can use our algo to verify equality of polynomials with high probability

The difference with a mathematical proof is that in pure math we are not satisfied with knowing that a theorem holds with 99% certainty :)