# Alternative approaches to algorithm design and analysis

- **Problem:** Exhaustive search gives typically  $\mathcal{O}(n!) \approx \mathcal{O}(n^n)$ -algorithms for  $\mathcal{NP}$ -complete problems.
- So we need to get around the **worst case / best solution** paradigm:
  - worst-case  $\rightarrow$  average-case analysis
  - best solution  $\rightarrow$  approximation
  - best solution  $\rightarrow$  randomized algorithms



**Def. 1** Let *L* be an optimization problem. We say that algorithm *M* is a **polynomial-time**  $\epsilon$ -approximation algorithm for *L* if *M* runs in polynomial time and there is a constant  $\epsilon \ge 0$  such that *M* is guaranteed to produce, for all instances of *L*, a solution whose cost is within an  $\epsilon$ -neighborhood from the optimum.

**Note 1:** Formally this means that the **relative** error  $\frac{|t_M(n) - OPT|}{OPT}$  must be less than or equal to the constant  $\epsilon$ .

**Note 2:** We are still looking at the worst case, but we don't require the very best solution any more.

**Example:** TSP with triangle inequality has a polynomial-time approximation algorithm.





#### **Algorithm TSP-**△:

Phase I: Find a minimum spanning tree. Phase II: Use the tree to create a tour.



The cost of the produced solution can not be more than  $2 \cdot \text{OPT}$ , otherweise the OPT tour (minus one edge) would be a more minimal spanning tree itself. Hence  $\epsilon = 1$ .



Opt. tour



**Theorem 1** TSP has no polynomial-time  $\epsilon$ -approximation algorithm for any  $\epsilon$  unless  $\mathcal{P}=\mathcal{NP}$ .

#### **Proof:**

Idea: Given  $\epsilon$ , make a reduction from HAMILTONICITY which has only **one** solution within the  $\epsilon$ -neighborhood from OPT, namely the optimal solution itself.

$$a \rightarrow c \rightarrow d \qquad a \qquad b \qquad c \qquad d$$

$$a \qquad 2+\epsilon n \qquad 1 \qquad 2+\epsilon n \qquad 1$$

$$b \rightarrow c \qquad c \qquad b \qquad 1 \qquad 2+\epsilon n \qquad 1 \qquad 2+\epsilon n$$

$$d \qquad 1 \qquad 2+\epsilon n \qquad 1 \qquad 2+\epsilon n \qquad 1$$

$$d \qquad 1 \qquad 2+\epsilon n \qquad 1 \qquad 2+\epsilon n$$

$$K = n(=4)$$

The **error** resulting from picking a non-edge is: Approx.solutin - OPT =  $(n - 1 + 2 + \epsilon n) - n = (1 + \epsilon)n > \epsilon n$ 

Hence a polynomial-time  $\epsilon$ -approximation algorithm for TSP combined with the above reduction would solve HAMILTONICITY in polynomial time.

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#### **Example: VERTEX COVER**

- **Heuristics** are a common way of dealing with intractable (optimization) problems in practice.
- Heuristics differ from algorithms in that they have no performance guarantees, i.e. they don't always find the (best) solution.

A greedy heuristic for VERTEX COVER-opt.:

#### **Heuristic VC-H1:**

Repeat until all edges are covered:

- 1.Cover highest-degree vertex v;
- 2.Remove v (with edges) from graph;



**Theorem 2** The heuristic VC-H1 is not an  $\epsilon$ -approximation algorithm for VERTEX COVER-opt. for any fixed  $\epsilon$ .



#### **Proof:**



Show a **counterexample**, i.e. cook up an instance where the heuristic performs badly.

#### **Counterexample:**

- A graph with nodes  $a_1, \ldots, a_n$  and  $b_1, \ldots, b_n$ .
- Node  $b_i$  is only connected to node  $a_i$ .
- A bunch of *c*-nodes connected to *a*-nodes in the following way:
  - Node  $c_1$  is connected to  $a_1$  and  $a_2$ . Node  $c_2$  is connected to  $a_3$  and  $a_4$ , etc.
  - Node  $c_{n/2+1}$  is connected to  $a_1$ ,  $a_2$  and  $a_3$ . Node  $c_{n/2+2}$  is connected to  $a_4$ ,  $a_5$  and  $a_6$ , etc.
  - Node  $c_{m-1}$  is connected to  $a_1, a_2, \ldots a_{n-1}$ .
  - Node  $c_m$  is connected to all *a*-nodes.



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- The optimal solution OPT requires *n* guards (on all *a*-nodes).
- VC-H1 first covers all the *c*-nodes (starting with *c<sub>m</sub>*) before covering the *a*-nodes.
- The number of c-nodes are of order  $n \log n$ .
- Relative error for VC-H1 on this instance:

 $\frac{|\text{VC-H1}| - |\text{OPT}|}{|\text{OPT}|} = \frac{(n \log n + n) - n}{n}$  $= \frac{n \log n}{n} = \log n \neq \epsilon$ 

• The relative error **grows as a function of** *n*.

#### **Heuristic VC-H2:**

Repeat until all edges are covered:

Pick an edge e;
Cover and remove both endpoints of e.

Since at least one endpoint of every edge must be covered, |VC-H2| ≤ 2 · |OPT|.
So VC-H2 is a polynomial-time ε-approximation algorithm for VC with ε = 1.
Surpisingly, this "stupid-looking" algorithm is the best (worst case) approximation algorithm known for VERTEX COVER-opt.

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# Average-case analysis & algorithms







- **Problem =**  $(L, P_r)$  where  $P_r$  is a probability function over the input strings:  $P_r : \sum^* \to [0, 1].$
- $\sum_{x \in \sum^{*}} P_r(x) = 1$  (the probabilities must sum up to 1).
- Average time of an algorithm:

$$T_{A}(n) = \sum_{\{x \in \sum^{*} | |x|=n\}} T_{A}(x) P_{r}(x)$$

- Key issue: How to choose  $P_r$  so that it is a realistic model of reality.
- Natural solution: Assume that all instances of length *n* are equally probable (uniform distribution).



#### **Random graphs**

#### **Uniform probability model (UPM)**

- Every graph G has equal probability
- If the number of nodes = n, then  $P_r(G) = \frac{1}{\#\text{graphs}} = \frac{1}{2^{\binom{n}{2}}}$ , where  $\binom{n}{2} = \frac{n(n-1)}{2}$
- UPM is more natural for interpretation

#### Independent edge probability model (IEPM)

- Every possible edge in a graph *G* has equal probabilility *p* of occuring
- The edges are independent in the sense that for each pair (*s*, *t*) of vertices, we make a new toss with the coin to decide whether there will be an edge between *s* and *t*.
- For  $p = \frac{1}{2}$  IEPM is identical to UPM:

$$P_r(G) = \left(\frac{1}{2}\right)^m \cdot \left(\frac{1}{2}\right)^{\binom{n}{2}-m} = \frac{1}{2^{\binom{n}{2}}}$$

• IEPM is easier to work with



#### **Example: 3-COLORABILITY**

In 3-COLORABILITY we are given a graph as input and we are asked to decide whether it is possible to color the nodes using 3 different colors in such a way that any two nodes have different colors if there is an edge between them.

**Theorem 3** 3-COLORABILITY, which is an  $\mathcal{NP}$ -complete problem, is solvable in **constant** average (expected) time on the IEPM with p = 1/2 by a branch-and-bound algorithm (with exponential worst-case complexity).

#### **Proof:**

**Strategy** (for a rough estimate): Use the indep. edge prob. model. Estimate expected time for finding a proof of non-3-colorability.



 $K_4$  (a clique of size 4) is a proof of non-3-colorability.



- The probability of 4 nodes being a  $K_4$ :  $P_r(K_4) = 2^{-\binom{4}{2}} = 2^{-6} = \frac{1}{128}$
- Expected no. of 4-vertex sets examined before a  $K_4$  is found:

$$\sum_{i=1}^{\infty} i(1-2^{-6})^{i-1} 2^{-6} = 2^{-6} \sum_{i=1}^{\infty} i(1-2^{-6})^{i-1}$$
$$\stackrel{*}{=} 2^{-6} \frac{1}{(1-(1-2^{-6}))^2}$$
$$= 2^{-6} \frac{1}{(2^{-6})^2} = \frac{2^{12}}{2^6} = 2^6 = 128$$

- $(1-2^{-6})^{i-1}2^{-6}$  is the probability that the first  $K_4$  is found after examining exactly *i* 4-vertex sets.
- (\*) is correct due to the following formula  $(q = 1 2^{-6})$  from mathematics (MA100):

$$\sum_{i=1}^{\infty} iq^{i-1} = \frac{\delta}{\delta q} \left(\sum_{i=1}^{\infty} q^i\right) = \frac{\delta}{\delta q} \left(\frac{q}{1-q}\right)$$
$$= \frac{1}{(1-q)^2}$$

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**Conlusion:** Using IEPM with  $p = \frac{1}{2}$  we need to check 128 four-vertex sets on average before we find a  $K_4$ .

**Note:** Random graphs with constant edge probability are very dense (have lots of edges). More realistic models has p as a function of n (the number of vertices), i.e.  $p = 1/\sqrt{n}$  or p = 5/n.

#### **0-1 Laws**

as a link between probabilistic and deterministic thinking.

Example: "Almost all" graphs are

- not 3-colorable
- Hamiltonian
- connected
- . . .

**Def. 2** A property of graphs or strings or other kind of problem instances is said to have a **zero-one law** if the limit of the probability that a graph/string/problem instance has that property is either 0 or 1 when n tends to infinity  $(\lim_{n\to\infty})$ .



#### **Example:** HAMILTONICITY

a linear expected-time algorithm for random graphs with p = 1/2.

 Difficulty: The probability of non-Hamiltonicity is too large to be ignored, e.g. P<sub>r</sub>(∃ at least 1 isolated vertex) = 2<sup>-n</sup>.

#### • The algorithm has 3 phases:

- Phase 1: Construct a Hamiltonian path in linear time. Fails with probability  $P_1(n)$ .
- Phase 2: Find proof of non-Hamiltonicity or construct Hamiltonian path in time  $\mathcal{O}(n^2)$ . Unsuccessful with probability  $P_2(n)$ .
- Phase 3: Exhaustive search (dynamic programming) in time  $O(2^{2n})$ .

• Expected running time is  

$$\leq \mathcal{O}(n) + \mathcal{O}(n^2) P_1(n) + \mathcal{O}(2^{2n}) P_1(n) P_2(n)$$

$$= \mathcal{O}(n) \text{ if } P_1(n) \cdot \mathcal{O}(n^2) = \mathcal{O}(n)$$
and  $P_1(n) P_2(n) \cdot \mathcal{O}(2^{2n}) = \mathcal{O}(n)$ 

- Phase 2 is necessary because  $\mathcal{O}(2^{-n}) \cdot \mathcal{O}(2^{2n}) = \mathcal{O}(2^n).$
- After failing to construct a Hamiltonian path fast in phase 1, we first reduce the probability of the instance being non-Hamiltonian (phase 2), before doing exhaustive search in phase 3.

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# **Randomized computing**

Machines that can **toss coins** (generate random bits/numbers)

- Worst case paradigm
- Always give the correct (best) solution



#### **Randomized algorithms**

Idea: Toss a coin & simulate non-determinism

#### Example 1: Proving polynomial non-identities

$$\begin{aligned} (x+y)^2 &\stackrel{?}{\neq} x^2 + 2xy + y^2 \\ &\stackrel{?}{\neq} x^2 + y^2 \end{aligned}$$

- What is the "classical" complexity of the problem?
- Fast, randomized algorithm:
  - Guess values for *x* and *y* and compute left-hand side (LHS) and right-hand side (RHS) of equation.
  - If LHS  $\neq$  RHS, then we know that the polynomials are different.
  - If LHS = RHS, then we suspect that the polynomials are identical, but we don't know for sure, so we repeat the experiment with other *x* and *y* values.
- Idea works if there are many witnesses.

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o witnesses

Let f(n) be a polynomial in n and let the probability of success after f(n) steps/coin tosses be  $\geq \frac{1}{2}$ . After f(n) steps the algorithm either

- finds a witness and says "Yes, the polynomials are different", or
- halts without success and says "No, maybe the polynomials are identical".

This sort of algorithm is called a **Monte Carlo** algorithm.



**Note:** The probability that the Monte Carlo algorithm succeeds after f(n) steps is **independent of input** (and dependent only on the coin tosses).

- Therefore the algorithm can be repeated on the same data set.
- After 100 repeated trials, the probability of failure is  $\leq 2^{-100}$  which is smaller then the probability that a meteorite hits the computer while the program is running!

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# **Metaheuristics**

### **Simulated Annealing**

- Analogy with physical annealing
- 'Temperature' T, annealing schedule
- 'Bad moves' with probability  $\exp\left(-\delta f/T\right)$

#### **Genetic algorithms**

- Analogy with Darwinian evolution
- 'individuals', 'fitness', 'cross breeding'

#### **Neural Networks**

- Analogy with human mind
- 'neurons', 'learning'

#### Taboo search

- Analogy with culture
- adaptive memory, responsive exploration



## **Parallel computing**



- some problems can be efficiently parallelized
- some problems seems inherently sequential

#### **Parallel machine models**

- Alternating TMs
- Boolean Circuits



 Boolean Circuit complexity: "time" (length of longest directed path) and hardware (# of gates)

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# • Parallell Random Access Machines (PRAMs)



— Read/Write conflict resolution strategy

— PRAM complexity: time (# of steps) and hardware (# of processors)

**Example:** Parallel summation in time  $\mathcal{O}(\log n)$ 



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# Limitations to parallel computing

#### **Good news**

parallel time  $\leftrightarrow$  sequential space

**Example:** HAMILTONICITY can easily be solved in parallel polynomial time:

- On a graph with *n* nodes there are at most *n*! possible Hamiltonian paths.
- Use *n*! processors and let each of them check 1 possible solution in polynomial time.
- Compute the the OR of the answers in parallel time  $\mathcal{O}(\log(n!)) = \mathcal{O}(n \log n)$ .

#### **Bad news**

#### Theorem 4 With polynomial many processors

parallel poly. time = sequential poly. time

#### **Proof:**

- 1 processor can simulate one step of mprocessors in sequential time  $t_1(m) = \mathcal{O}(m)$
- Let  $t_2(n)$  be the polynomial parallel time of the computation. If m is polynomial then  $t_1(m) \cdot t_2(n) =$  polynomial.



## **Parallel complexity classes**

**Def. 3** A language is said to be in class  $\mathcal{NC}$  if it is recognized in polylogarithmic,  $\mathcal{O}(\log^k(n))$ , parallel time with uniform polynomial hardware.



 $\mathcal{P}$ -hard, Ex: CIRCUIT VALUE

•  $\mathcal{P} \stackrel{?}{=} \mathcal{NC}$ 

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