CSE 548: (*Design and*) Analysis of Algorithms Coping with NP-Completeness

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Coping with NP-Completeness

- Sometimes you are faced with hard problems problems for which no efficient solutions exist.
- **Step 1:** Try to show that the problem is *NP*-complete
 - This way, you can avoid wasting a lot of time on a fruitless search for an efficient algorithm
- Step 2a: Sometimes, you may be able to say "let us solve a different problem"
 - you may be able leverage some special structure of your problem domain that enables a more efficient solution
- Step 2b: Other times, you are stuck with a difficult problem and you need to make the best of it.
 - We discuss different coping strategies in such cases.

Intelligent Exhaustive Search

- Exhaustive search will work for almost any problem Hamiltonian Tour: Consider an edge *e*.
 - Either e = (u, v) is part of the tour, in which case you can complete the tour by finding a path from u to v in G e.
 - Or, e is not part of the tour, in which case you can find the tour by searching G e.

Either case leads to a recurrence T(m) = 2T(m-1), i.e., $T(m) = O(2^m)$. (Here m is the number of edge in G.)

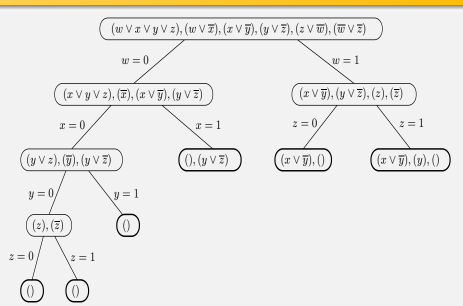
SAT: Try all 2^n possible truth assignments to the n variables in your formula.

• The key point is to be intelligent in the way this search is conducted, so that the algorithm is faster than 2^n in practice.

Backtracking

- Depth-first approach to perform exhaustive search
 - In the above example, first try to find a solution that includes e
 - Looking down further, the algorithm will make additional choices of edges to include: $e_1, e_2, ..., e_k$
 - Only when all paths that include e fail to be Hamiltonian, we consider the alternative (i.e., Hamiltonian path that doesn't include e)
- Key goal is to recognize and prune failing paths as quickly as possible.

Backtracking Approach for SAT



Backtracking Approach for SAT: Complexity

• There are two cases, based on the variable *w* chosen for branching:

Case 1: Both w and \overline{w} occur in the formula In this case, both branches are present. Moreover, both w and \overline{w} are eliminated from the formula at this point, so we have the recurrence:

$$T(n) = 2T(n-2) + O(n)$$

Case 2: Only one of them is present. In this case, only one of the branches needs exploring, so we have the recurrence

$$T(n) = T(n-1) + O(n)$$

• Clearly, case 1 will dominate, so let us ignore case 2. Case 1 yields a solution of $O(2^{n/2})$ or $O(1.414^n)$, which is much better than 2^n .

Backtracking Approach for SAT: Improvements

- We can improve the worst-case bound by choosing a variable that occurs most times
 - If it occurs *k* times, then you have the recurrence

$$T(n) = 2T(n-k)$$

whose solution is $O(2^{n/k})$.

- Of course, you won't be able to repeatedly find a variable that occurs k
 times, so this solution is meaningless in practice it just goes to show
 the exponential pruning effect of a frequently occurring variable
- Another strategy: pick a clause with fewest number of variables, and pick those variables in sequence.
- *Exercise:* Show that the backtracking algorithm solves 2SAT in polynomial time

Branch and Bound

- Generalization of backtracking to support optimization problems
- Requires a lower bound on the cost of solutions that may result from a partial solution
 - If the cost is higher than that of a previously encountered solution, then this subproblem need not be explored further.
- Sometimes, we may rely on estimates of cost rather than strict lower bounds.

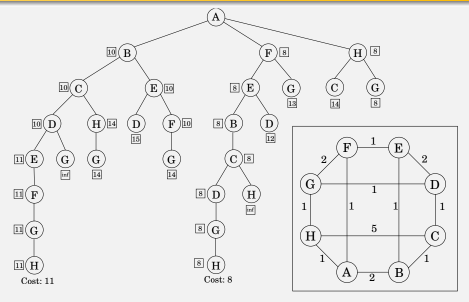
Branch and Bound for TSP

- Begin with a vertex a the goal is to compute a TSP that begins and ends at a.
- We begin the search by considering an edge from a to its neighbor x, another edge from x to a neighbor of x, and so on.
- *Partial solutions* represent a path from a to some vertex b, passing through a set $S \subset V$ of vertices.
- Completing a partial solution requires the computation of a low cost path from b to a using only vertices in V-S

Lower bound on costs of partial TSP solutions

- To complete the path from b to a, we must incur at least the following costs
 - Cost of going from b to a vertex in V-S, i.e, the minimum weight edge from b to a vertex in V-S
 - Cost of going from a V S vertex to a, i.e, the minimum weight edge from a to a vertex in V S
 - Minimal cost path in V S that visits all $v \in V S$
 - *Note:* Lower bound is the cost of MST for V S
- By adding the above three cost components, we arrive at a lower bound on solutions derivable from a partial solution.

Illustration of Branch-and Bound for TSP



Approximation Algorithms

- Relax optimality requirement: permit approximate solutions
 - Solutions that are within a certain distance from optimum
- *Not heuristics:* Approximate algorithms *guarantee* that solutions are within a certain distance from optimal
 - Differs from heuristics that can sometimes return very bad solutions.
- How to define "distance from optimal?"
 - Additive: Optimal solution S_O and the Solution S_A returned by approximation algorithm differ only by a constant.
 - Quality of approximation is extremely good, but unfortunately, most problems don't admit such approximations
 - Factor: S_O and S_A are related by a factor.
 - Most known approximation algorithms fall into this category.

Approximation Factors

- Constant: $S_A \leq kS_O$ for some fixed constant k.
 - Examples: Vertex cover, Facility location, ...
- Logarithmic: $S_A \leq O(\log^k n) \cdot S_O$.
 - Examples: Set cover, dominating set, ...
- Polynomial: $S_A \leq O(n^k) \cdot S_O$.
 - Examples: Max Clique, Independent set, graph coloring, ...
- PTAS: $S_A \leq (1 + \epsilon) \cdot S_O$ for any $\epsilon > 0$. ("Polynomial-time approximation scheme")
- **FPTAS:** PTAS with runtime $O(\epsilon^{-k})$ for some k. ("Fully PTAS")
 - Examples: Knapsack, Bin-packing, Euclidean TSP, ...

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Bin Packing

Problem

Pack objects of different weight into bins that have a fixed capacity in such a way that minimizes bins used.

- Obvious similarity to Knapsack
- Bin-packing is NP-hard
- Very good (and often very simple) approximation algorithms exist

First-fit Algorithm

A simple, greedy algorithm

FirstFit(x[1..n])

for i = 1 to n do

Put x[i] into the first open bin large enough to hold it

Theorem

All open bins, except possibly one, are more than half-full

Proof: Suppose that there are two bins b and b' that are less than half-full. Then, items in b' would have fitted into b, and so the FF algorithm would never have opened the bin b' — a contradiction

Theorem

First-fit is optimal within a factor of 2: specifically, $S_A < 2S_O + 1$.

Best-Fit Algorithm

- Another simple, greedy algorithm
- Instead of using the first bin that will can hold x[i], use the open bin whose remaining capacity is closest to x[i]
 - Prefers to keep bins close to full.
- Factor-2 optimality can established easily.

Other algorithms for Bin-packing

- First-fit decreasing strategy first sorts the items so that $x[i] \ge x[i+1]$ and then runs first-fit.
- Best-fit decreasing strategy first sorts the items so that $x[i] \ge x[i+1]$ and then runs best-fit.
- Both FFD and BFD achieve approximation factors of $11/9S_O + 6/9$.
- Due to the additive term, bin-packing cannot have a PTAS unless P = NP.
 - But $S_A = (1 + \epsilon)S_O + 1$ is easy to achieve for any $\epsilon > 0$

Set Cover

Problem

Given a collection $S_1, ..., S_m$ of subsets of B, find a minimum collection S_{i_1}, \ldots, S_{i_k} such that $\bigcup_{i=1}^k S_{i_i} = B$

Greedy Set Cover Algorithm

GSC(S, B) $cover = \emptyset$; $covered = \emptyset$ while covered $\neq B$ do Let *new* be the set in S-cover containing the maximum number of elements of B-coveredadd new to cover; covered = covered \cup new

return cover

Theorem

Greedy set cover is approximate with a factor of $\ln n$, where n = |B|

Proof:

- Let k be the size of optimal cover, and n_t be the number of elements left uncovered after t steps of GSC
- These n_t elements are covered by k sets in optimal cover \Rightarrow each of these k sets must cover at least n_t/k uncovered elements.
- Thus, GSC will find at least one set that covers n_t/k elements.
- This yields the recurrence for bounding uncovered elements: $U(t+1) = n_t - n_t/k = n_t(1-1/k) = U(t)(1-1/k)$
- The solution to recurrence is $n(1-1/k)^t < ne^{-t/k}$
- Thus, after $t = k \ln n$ steps, less than 1 (i.e., no) elements uncovered
- Thus, GSC computes a cover at most ln *n* times the optimal cover.

Vertex Cover

- Note that a vertex cover is a set cover for (S, E), where $S = \{\{(v, u) | v \in V \text{ and } (v, u) \in E\} | v \in V\},$
 - S contains a set for each vertex; this set lists all edges incident on v
- Thus GSC is an approximate algorithm for vertex cover.
- But ln *n* is not a factor to be thrilled about can we do better?
 - Actually, we can do much better! That too with a very simple algorithm.

Vertex Cover

Consider any edge (u, v).

- Either *u* or *v* must belong to any vertex cover.
- If we accept $S_A = 2S_O$, then we can avoid the guesswork by simply picking both vertices!

Approximate Vertex Cover Algorithm

$$AVC(G = (V, E))$$

 $C = \emptyset$
while G is not empty
pick any $(u, v) \in E$
 $C = C \cup \{u, v\}$
 $G = G - \{u, v\}$

return C

k-Cluster

Problem

Given $X = \{x_1, \dots, x_n\}$ and distances between x_i , partition X into k clusters in a way that minimizes maximum cluster diameter.

Approximate *k*-Cluster Algorithm (AC)

Pick any point $\mu_1 \in X$ as the first cluster center

for
$$i = 2$$
 to k do

Choose μ_i to be the farthest point from μ_1, \ldots, μ_{i-1}

Create *k* clusters $C_i = \{x \in X | \mu_i \text{ is the closest center to } x\}$

Let x be the farthest point from μ_1, \ldots, μ_k , and let r the distance to its closest center. Then, we can say:

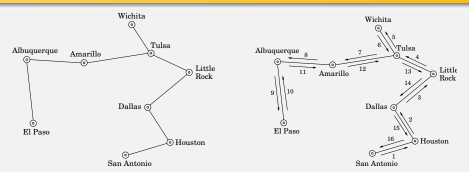
- Cluster diameter of C_1, \ldots, C_k is at most 2r
- The distance between any 2 points in $\{x, \mu_1, \dots, \mu_k\}$ is at least r. This follows from:
 - how μ_i 's was chosen to be the farthest point from μ_i for i < i,
 - this distance to μ_i must decrease with i, and
 - when i = k + 1, this distance is r
- Thus, any k-Cluster must have a diameter of at least r
 - With k circles, at least two of k+1 points must be within one of them.
 - This circle's diameter must hence be r or greater
- Thus, AC is approximate within a factor of 2.

Euclidean TSP

- Our starting point is once again the MST
- Note that no TSP solution can be smaller than MST
 - Deleting an edge from TSP solution yields a spanning tree
- Simple algorithm:
 - Start with the MST

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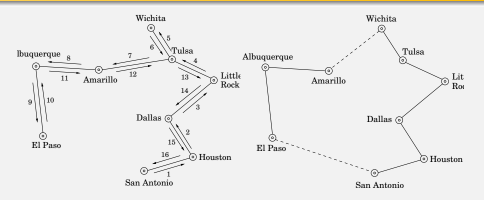
Approximating Euclidean TSP: An Illustration



- Start with the MST
- Make a tour that uses each MST edge twice (forward and backward)
 - This tour is like TSP in ending at the starting node, and differs from TSP by visiting some vertices and edges twice

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Approximating Euclidean TSP: An Illustration (2)



- Avoid revisits by short-circuiting to next unvisited vertex
- By triangle inequality, short-circuit distance can only be less than the distance following MST edges.
 - Thus, tour length less than 2xMST, i.e., approximate within a factor 2.

Knapsack

Knap01(w, v, n, W) $V = \sum_{i=0}^{n} v[j]$ $K[0, v] = 0, \forall 0 < v < V$ **for** j = 1 to n **do** for v = 1 to V do if v[j] > v then K[j, v] = K[j-1, v]

Computes minimum weight of knapsack for a given value.

else K[j, v] = min(K[j-1, v], K[j-1, v-v[j],] + w[i])

return maximum v such that K[n, v] < W

- Iterates over all possible items and all possible values: O(nV)
 - we derive a polynomial time approximate algorithm from this

$Knap01FPTAS(w, v, n, W, \epsilon)$

$$v_i' = \left\lfloor \frac{v_i}{\max_{1 \le j \le n} v_j} \cdot \frac{n}{\epsilon} \right\rfloor, \text{ for } 1 \le i \le n$$

$$Knap01(w, v', n, W)$$

- Rescaling consists of two steps:
 - Express value of each item relative to the most valuable item
 - If we worked with real values, this step won't change the optimal solution
 - Multiply relative values by a factor n/ϵ to get an integer
- Floor operation introduces an error ≤ 1 in v'_i (e.g., |3.99| = 3)
- Error in *Knap*01 output = error in $\sum v_i'$, which is at most $n \cdot 1$
- We scale each v_i' by n/ϵ , so relative error is $n/(n/\epsilon) = \epsilon$
 - Thus we have achieved the desired approximation.

FPTAS for 0-1 Knapsack: Runtime

$Knap01FPTAS(w, v, n, W, \epsilon)$

$$v'_{i} = \left\lfloor \frac{v_{i}}{\max_{1 \leq j \leq n} v_{j}} \cdot \frac{n}{\epsilon} \right\rfloor, \text{ for } 1 \leq i \leq n$$

$$Knap01(w, v', n, W)$$

- Note that we are using *Knap*01 with rescaled values, so the complexity is O(nV').
- Note: $V' = \sum_{1}^{n} v'_{i} \leq n \cdot max_{1 \leq i \leq n} v'_{i}$
- It is easy to see from definition of v'_i that $\max_{1 \le i \le n} v'_i = n/\epsilon$. Substituting this into the above equation yields a complexity of:

$$O(nV') \leq O(n(n \cdot \max_{1 \leq i \leq n} v_i')) = O(n(n \cdot (n/\epsilon))) = O(n^3/\epsilon)$$

• By varying ϵ , we can trade off accuracy against runtime.