CS711008Z Algorithm Design and Analysis

Lecture 6. Basic algorithm design technique: Dynamic programming

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Outline

- The first example: MATRIXCHAINMULTIPLICATION
- Elements of dynamic programming technique
- Various ways to describe subproblems: KNAPSACK, RNA SECONDARY STRUCTURE, HIDDEN MARKOV MODEL, SEQUENCE ALIGNMENT, and SHORTEST PATH
- Techniques to reduce space and time consumption: HIRSCHBERG algorithm and banded DP
- Connection with greedy technique: Interval Scheduling, Shortest Path

Dynamic programming and its connection with divide-and-conquer

- Dynamic programming typically applies to optimization problems if:
 - The original problem can be divided into smaller subproblems, and
 - The recursion among sub-problems has optimal-substructure property, i.e., the optimal solution to the original problem can be calculated through combining the optimal solutions to subproblems.
- Unlike the general divide-and-conquer framework, a dynamic programming algorithm usually enumerates all possible dividing strategies.
- To identify meaningful recursions, one of the key steps is to define an appropriate general form of sub-problems. For this aim, it is helpful to describe the solving process as a multistage decision process.

Revisiting the DIVIDE AND CONQUER technique

- To see whether the DIVIDE AND CONQUER technique applies on a given problem, we need to examine both input and output of the problem description.
 - Examine the input part to determine how to decompose the problem into subproblems of same structure but smaller size: It is relatively easy to decompose a problem into subproblems if the input part is related to the following data structures:
 - An array with n elements;
 - A matrix;
 - A set of n elements;
 - A tree;
 - A directed acyclic graph;
 - A general graph.
 - Examine the output part to determine how to construct the solution to the original problem using the solutions to its subproblems.

 $\label{eq:MatrixChainMultiplication problem: recursion over sequences$

MATRIXCHAINMULTIPLICATION problem

INPUT:

A sequence of n matrices $A_1, A_2, ..., A_n$; matrix A_i has dimension $p_{i-1} \times p_i$;

OUTPUT:

Fully parenthesizing the product $A_1A_2...A_n$ in a way to minimize the number of scalar multiplications.

Let's start from a simple example

$$A_1 = \begin{bmatrix} 1 & 2 \end{bmatrix} \ A_2 = \begin{bmatrix} 1 & 2 & 3 \\ 1 & 2 & 3 \end{bmatrix} \ A_3 = \begin{bmatrix} 1 & 2 & 3 & 4 \\ 1 & 2 & 3 & 4 \\ 1 & 2 & 3 & 4 \end{bmatrix}$$

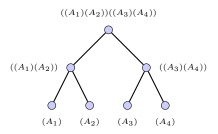
Solutions:
$$((A_1)(A_2))(A_3)$$
 $(A_1)((A_2)(A_3))$ #Multiplications: $1\times 2\times 3$ $2\times 3\times 4$ $+1\times 3\times 4$ $+1\times 2\times 4$ $=18$ $=32$

- Here we assume that the calculation of A_1A_2 needs $1\times 2\times 3$ scalar multiplications.
- The objective is to determine a calculation sequence such that the number of multiplications is minimized.



The solution space size

• Intuitively, a calculation sequence can be described as a binary tree, where each node corresponds to a subproblem.



• The total number of possible calculation sequences:

$$\binom{2n}{n} - \binom{2n}{n-1}$$
 (Catalan number)

- Thus, it takes exponential time to enumerate all possible calculation sequences.
- Question: can we design an efficient algorithm?

A dynamic programming algorithm (by S. S. Godbole, 1973)

Defining general form of sub-problems

- 1 It is not easy to solve the problem directly when n is large. Let's investigate whether it is possible to reduce into smaller sub-problems.
- Solution: a full parentheses. Let's describe the solving process as a process of multistage decisions, where each decision is to add parentheses at a position.
- **3** Suppose we have already worked out the optimal solution O, where the first **decision** adds two parentheses as $(A_1...A_k)(A_{k+1}...A_n)$.
- **①** This decision decomposes the original problem into two independent sub-problems: to calculate $A_1...A_k$ and $A_{k+1}...A_n$.
- **5** Summarizing these two cases, we define the general form of sub-problems as: to calculate $A_i...A_j$ with the minimal number of scalar multiplications.

Optimal substructure property

- The general form of sub-problems: to calculate $A_i...A_j$ with the minimal number of scalar multiplications. Let's denote the optimal solution value to the sub-problem as OPT(i,j), thus the original problem can be solved via calculating OPT(1,n).
- The optimal solution to the original problem can be obtained through combining the optimal solutions to sub-problems.
 This recursion can be stated as the following optimal substructure property:

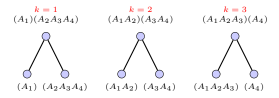
$$OPT(1, n) = OPT(1, k) + OPT(k + 1, n) + p_0 p_k p_n$$

Proof of the optimal substructure property

- "Cut-and-paste" proof:
 - Suppose for $A_1...A_k$, there is another parentheses OPT'(1,k) better than OPT(1,k). Then the combination of OPT'(1,k) and OPT(k+1,n) leads to a new solution with lower cost than OPT(1,n): a contradiction.
 - Here, the independence between $A_1...A_k$ and $A_{k+1}...A_n$ guarantees that the substitution of OPT(1,k) with OPT'(1,k) does not affect solution to $A_{k+1}...A_n$.

A recursive solution

- So far so good! The only difficulty is that we have no idea of the first splitting position k in the optimal solution.
- How to overcome this difficulty? Enumeration! We enumerate all possible options of the first decision, i.e. for all k, i < k < j.



• Thus we have the following recursion:

$$OPT(i,j) = \begin{cases} 0 & i = j \\ \min_{\mathbf{i} \leq \mathbf{k} < \mathbf{j}} \{OPT(i,k) + OPT(k+1,j) + p_{i-1}p_kp_j\} & otherwise \end{cases}$$

Implementing the recursion: trial $\boldsymbol{1}$

Trial 1: Explore the recursion in the top-down manner

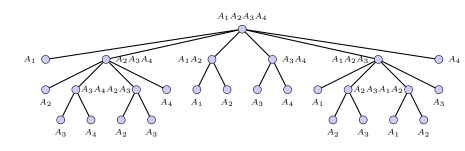
```
RECURSIVE_MATRIX_CHAIN(i, j)
 1: if i == j then
2: return 0;
3: end if
4: OPT(i, j) = +\infty;
5: for k = i to j - 1 do
    q = RECURSIVE\_MATRIX\_CHAIN(i, k)
6:
       + RECURSIVE\_MATRIX\_CHAIN(k+1, j)
7:
8: +p_{i-1}p_kp_j;
9: if q < OPT(i, j) then
10: OPT(i, j) = q;
11: end if
12: end for
13: return OPT(i, j);
```

• Note: The optimal solution to the original problem can be obtained through calling $RECURSIVE_MATRIX_CHAIN(1, n)$.

An example

$$A_1 = \begin{bmatrix} 1 & 2 \end{bmatrix} \ A_2 = \begin{bmatrix} 1 & 2 & 3 \\ 1 & 2 & 3 \end{bmatrix} \ A_3 = \begin{bmatrix} 1 & 2 & 3 & 4 \\ 1 & 2 & 3 & 4 \\ 1 & 2 & 3 & 4 \end{bmatrix} \ A_4 = \begin{bmatrix} 1 & 2 & 3 & 4 & 5 \\ 1 & 2 & 3 & 4 & 5 \\ 1 & 2 & 3 & 4 & 5 \\ 1 & 2 & 3 & 4 & 5 \\ 1 & 2 & 3 & 4 & 5 \end{bmatrix}$$

$$1 \times 2 \qquad 2 \times 3 \qquad 3 \times 4 \qquad 3 \times 4$$



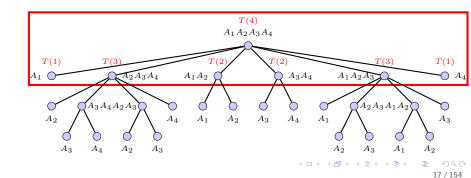
Note: each node of the recursion tree represents a subproblem.

However, this is not a good implementation

$\mathsf{Theorem}$

Algorithm RECURSIVE-MATRIX-CHAIN costs exponential time.

• Let T(n) denote the time used to calculate product of n matrices. Then $T(n) \geq 1 + \sum_{k=1}^{n-1} (T(k) + T(n-k) + 1)$ for n > 1.



Proof.

- We shall prove $T(n) \ge 2^{n-1}$ using the substitution technique.
 - Basis: $T(1) \ge 1 = 2^{1-1}$.
 - Induction:

$$T(n) \ge 1 + \sum_{k=1}^{n-1} (T(k) + T(n-k) + 1)$$
 (1)

$$= n + 2\sum_{k=1}^{n-1} T(k)$$
 (2)

$$\geq n + 2\sum_{k=1}^{n-1} 2^{k-1} \tag{3}$$

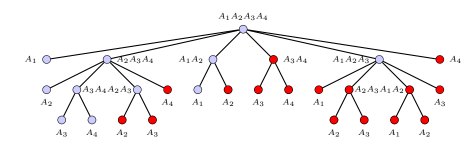
$$\geq n + 2(2^{n-1} - 1) \tag{4}$$

$$\geq n+2^n-2 \tag{5}$$

$$\geq 2^{n-1} \tag{6}$$



Why the first trial failed?



- Reason: There are only $O(n^2)$ subproblems. However, some subproblems (in red) were solved repeatedly.
- Solution: **memorize the solutions to subproblems** using an array OPT[1..n;1..n] for further look-up. The calculation of Fibonacci number is a good example of the power of the "memorizing" technique.

Implementing the recursion: trial 2

The "memorizing" technique

```
MEMORIZE\_MATRIX\_CHAIN(i, j)
 1: if OPT[i, j] \neq NULL then
     return OPT(i, j);
3: end if
4: if i == j then
5: OPT[i, j] = 0;
6: else
     for k = i to j - 1 do
       q = \text{MEMORIZE\_MATRIX\_CHAIN}(i, k)
8:
            +MEMORIZE_MATRIX_CHAIN(k+1, j)
9:
10:
            +p_{i-1}p_kp_i;
       if q < OPT[i, j] then
11:
          OPT[i, j] = q;
12:
13:
       end if
14: end for
15: end if
16: return OPT[i, j];
```

The "memorizing" technique cont'd

- The original problem can be solved by calling MEMORIZE_MATRIX_CHAIN(1,n) with all OPT[i,j] initialized as NULL.
- Time complexity: $O(n^3)$ (The calculation of each entry OPT[i,j] makes O(n) recursive calls in line 8.)
- Note that there are exponential ways of fully parenthesizing. The DP algorithm find the optimal solution in only $O(n^3)$ time since it avoids enumerating some redundant fully parenthesizing.

Implementing the recursion faster: trial 3

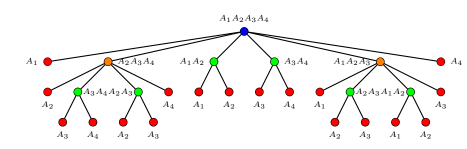
Trial 3: Faster implementation: unrolling the recursion in the bottom-up manner

MATRIX_CHAIN_MULTIPLICATION $(p_0, p_1, ..., p_n)$

```
1: for i = 1 to n do
2: OPT(i, i) = 0;
3: end for
4: for l=2 to n do
     for i = 1 to n - l + 1 do
5:
6: i = i + l - 1;
7: OPT(i, j) = +\infty;
8: for k = i to j - 1 do
         q = OPT(i, k) + OPT(k + 1, j) + p_{i-1}p_kp_j;
9:
10:
         if q < OPT(i, j) then
           OPT(i, j) = q;
11:
12:
           S(i, j) = k;
         end if
13:
       end for
14:
     end for
15:
16: end for
                                         17: return OPT(1,n):
```

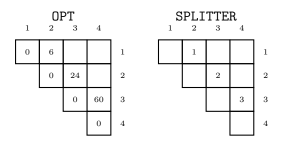
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Recursion tree: an intuitive view of the bottom-up calculation



- Solving sub-problems in a bottom-up manner, i.e.
 - Solving the sub-problems in red first;
 - Then solving the sub-problems in green;
 - Then solving the sub-problems in orange;
 - Finally we can solve the original problem in blue.

Step 1 of the bottom-up algorithm

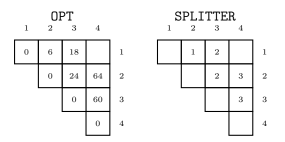


Step 1:

$$OPT[1,2] = p_0 \times p_1 \times p_2 = 1 \times 2 \times 3 = 6;$$

 $OPT[2,3] = p_1 \times p_2 \times p_3 = 2 \times 3 \times 4 = 24;$
 $OPT[3,4] = p_2 \times p_3 \times p_4 = 3 \times 4 \times 5 = 60;$

Step 2 of the bottom-up algorithm

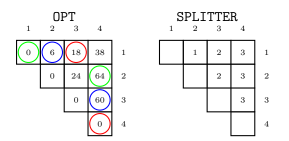


Step 2:

$$OPT[1,3] = \min \begin{cases} OPT[1,2] + OPT[3,3] + p_0 \times p_2 \times p_3 (=18) \\ OPT[1,1] + OPT[2,3] + p_0 \times p_1 \times p_3 (=32) \end{cases}$$
 Thus, $SPLITTER[1,2] = 2$.

$$OPT[2,4] = \min \begin{cases} OPT[2,2] + OPT[3,4] + p_1 \times p_2 \times p_4 (=90) \\ OPT[2,3] + OPT[4,4] + p_1 \times p_3 \times p_4 (=64) \end{cases}$$
 Thus, $SPLITTER[2,4] = 3$.

Step 3 of the bottom-up algorithm



Step 3:

$$OPT[1,4] = \min \begin{cases} OPT[1,1] + OPT[2,4] + p_0 \times p_1 \times p_4 (=74) \\ OPT[1,2] + OPT[3,4] + p_0 \times p_2 \times p_4 (=81) \\ OPT[1,3] + OPT[4,4] + p_0 \times p_3 \times p_4 (=38) \end{cases}$$
 Thus, $SPLITTER[1,4] = 3$.

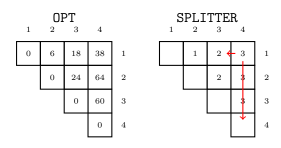
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Question: We have calculated the optimal **value**, but how to get the optimal **calculation sequence**?

Final step: constructing an optimal solution through "backtracking" the optimal options

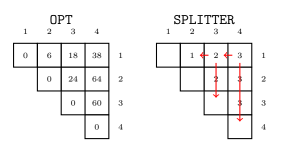
- Idea: **backtracking!** Starting from OPT[1, n], we trace back the source of OPT[1, n], i.e. which option we take at each decision stage.
- ullet Specifically, an auxiliary array S[1..n,1..n] is used.
 - Each entry S[i,j] records the optimal decision, i.e. the value of k such that the optimal parentheses of $A_i...A_j$ occurs between A_kA_{k+1} .
 - Thus, the optimal solution to the original problem $A_{1..n}$ is $A_{1..S[1,n]}A_{S[1,n]+1..n}$.
- Note: The optimal option cannot be determined before solving all subproblems.

Backtracking: step 1



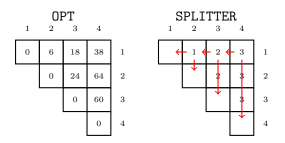
Step 1: $(A_1A_2A_3)(A_4)$

Backtracking: step 2



```
Step 1: (A_1A_2A_3)(A_4)
Step 2: ((A_1A_2)(A_3))(A_4)
```

Backtracking: step 3



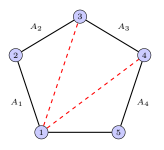
```
Step 1: (A_1A_2A_3)(A_4)
Step 2: ((A_1A_2)(A_3))(A_4)
Step 3: (((A_1)(A_2)(A_3))(A_4)
```

Summary: elements of dynamics programming

- 1 It is usually not easy to solve a large problem directly. Let's consider whether the problem can be decomposed into smaller sub-problems. How to define sub-problems?
 - Let's describe the solving process as a process of multistage decisions first.
 - Suppose that we have already worked out the optimal solution. Let's
 consider the first/final decision (in some order) in the optimal
 solution. The first/final decision might have several options.
 - We enumerate all possible options for the decision, and examine the generated sub-problems. The general form of sub-problems can be defined via summarizing all possible forms of sub-problems.
- Show that the recursion among sub-problems can be stated as the optimal substructure property, i.e. the optimal solution to the problem contains within it optimal solutions to subproblems.
- Programming: if recursive algorithm solves the same subproblem over and over, "tabular" can be used to avoid the repetition of solving same sub-problems.

Question: is $O(n^3)$ the lower bound?

An $O(n \log n)$ algorithm by Hu and Shing 1981



- One-to-one correspondence between parenthesis and partioning a convex polygon into non-intersecting triangles.
 - ullet Each node has a weight w_i , and a triangle corresponds to a product of the weight of its nodes.
 - The decomposition (red, dashed lines) has a weight sum of 38. In fact, it corresponds to the parenthesis (((A_1) (A_2) (A_3)) (A_4).
- ullet The optimal decomposition can be found in $O(n\log n)$ time.

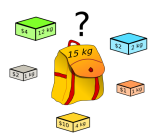
(See Hu and Shing 1981 for details)



 $0/1~\mathrm{KNAPSACK}$ problem: recursion over sets

A Knapsack instance

 Consider a set of items, where each item has a weight and a value. The objective is to select a subset of items such that the total weight is less than a given limit and the total value is as large as possible.



0/1 KNAPSACK problem

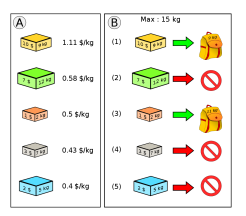
Formalized Definition:

INPUT: A set of items $S = \{1, 2, ..., n\}$. Item i has weight w_i and value v_i . A total weight limit W;

OUTPUT: A subset of items to maximize the total value with total weight below W.

- Here, "0/1" means that we should select an item (1) or abandon it (0), and we cannot select parts of an item.
- In contrast, FRACTIONAL KNAPSACK problem allow one to select a fractional, say 0.5, of an item.

0/1 Knapsack problem: an intuitive algorithm



- Intuitive method: selecting "expensive" items first.
- But this is not the optimal solution.

Defining the general form of sub-problems

- It is not easy to solve the problem with n items directly. Let's examine whether it is possible to reduce into smaller sub-problems.
- Solution: a subset of items. Let's describe the solving process as a process of multistage decisions. At the *i*-th decision stage, we decide whether item *i* should be selected.
- Suppose we have already worked out the optimal solution.
 Consider the first decision, i.e. whether the optimal solution contains item n or not (here we assume an order of the items and consider the items from end to beginning). This decision has two options:
 - Select: Then it suffices to select items as "expensive" as possible from $\{1, 2, ..., n-1\}$ with weight limit $W-w_n$.
 - **2** ABANDON: Otherwise, we should select items as "expensive" as possible from $\{1, 2, ..., n-1\}$ with weight limit W.
- In both cases, the original problem is reduced into smaller sub-problems.

Optimal sub-structure property

- Summarizing these two cases, we can set the general form of sub-problems as: to select items as "expensive" as possible from $\{1,2,...,i\}$ with weight limit w. Denote the optimal solution value as $OPT(\{1,2,...,i\},w)$.
- Then we can prove the optimal sub-structure property:

$$OPT(\{1, 2, ..., n\}, W) = \max \begin{cases} OPT(\{1, 2, ..., n-1\}, W) \\ OPT(\{1, 2, ..., n-1\}, W - w_n) + v_n \end{cases}$$

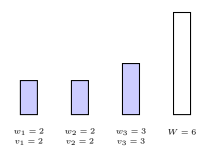
Algorithm

simplicity.

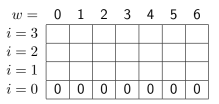
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\begin{array}{l} {\rm KNAPSACK}(n,W) \\ {\rm 1:} \ \ {\bf for} \ w=1 \ {\rm to} \ W \ \ {\bf do} \\ {\rm 2:} \ \ OPT[0,w]=0; \\ {\rm 3:} \ \ {\bf end} \ \ {\bf for} \\ {\rm 4:} \ \ {\bf for} \ i=1 \ {\rm to} \ n \ \ {\bf do} \\ {\rm 5:} \ \ \ \ {\bf for} \ w=1 \ {\rm to} \ W \ \ {\bf do} \\ {\rm 6:} \ \ \ \ OPT[i,w]=\max\{OPT[i-1,w],v_i+OPT[i-1,w-w_i]\}; \\ {\rm 7:} \ \ \ \ \ {\bf end} \ \ {\bf for} \\ {\rm 8:} \ \ \ \ {\bf end} \ \ {\bf for} \end{array}
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• Here we use OPT[i, w] to represent $OPT(\{1, 2, ..., i\}, w)$ for

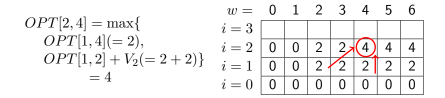
An example: Step 1



Initially all
$$OPT[0, w] = 0$$



$$OPT[1,2] = \max \{ & w = 0 & 1 & 2 & 3 & 4 & 5 & 6 \\ OPT[0,2](=0), & & i = 3 & & & & & \\ OPT[0,0] + V_1(=0+2) \} & i = 1 & 0 & 0 & 2 & 2 & 2 & 2 \\ & = 2 & & i = 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ \hline$$



$$OPT[3,3] = \max \{ & w = 0 & 1 & 2 & 3 & 4 & 5 & 6 \\ OPT[2,3](=2), & i = 2 & 0 & 0 & 2 & 2 & 4 & 4 & 4 \\ OPT[2,0] + V_3(=0+3) \} & i = 1 & 0 & 0 & 2 & 2 & 2 & 2 \\ = 3 & i = 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ \hline$$

Backtracking

$$\begin{array}{llll} OPT[3,6] = \max \{ & \\ OPT[2,6](=4), & \\ OPT[2,3] + V_3(=2+3) \} & \\ & = 5 & \\ \text{Decision: Select item 3} & w = & 0 & 1 & 2 & 3 & 4 & 5 & 6 \\ \hline Decision: Select item 3 & i = 3 & 0 & 0 & 2 & 3 & 4 & 5 & 5 \\ & i = 2 & 0 & 0 & 2 & 2 & 4 & 4 & 4 \\ \hline OPT[2,3] = \max \{ & i = 1 & 0 & 0 & 2 & 2 & 2 & 2 \\ OPT[1,3](=2), & i = 0 & 0 & 0 & 0 & 0 & 0 \\ OPT[1,1] + V_2(=0+2) \} & \end{array}$$

=2 Decision: Select item 2

Time complexity analysis

- \bullet Time complexity: O(nW). (Hint: for each entry in the matrix, only a comparison is needed; we have O(nW) entries in the matrix.)
- Notes:
 - **1** This algorithm is inefficient when W is large, say W = 1M.
 - ② Remember that a polynomial time algorithm costs time polynomial in the **input length**. However, this algorithm costs time $mW = m2^{\log W} = m2^{\text{input length}}$. Exponential!
 - **3** Pseudo-polynomial time algorithm: polynominal in the value of W rather than the **length** of W ($\log W$).
 - We will revisit this algorithm in approximation algorithm design.

Why should we consider items from end to beginning?

- Let's examine the following two types of selection of items:
 - If we consider an arbitrary item i, then the sub-problem becomes into "selecting items as expensive as possible from a subset s with weight limit w". We have the following recursion:

$$OPT(\{1, 2, ..., n\}, W) = \max \begin{cases} OPT(\{1, 2, ..., n\} - \{i\}, W) \\ OPT(\{1, 2, ..., n\} - \{i\}, W - w_i) + v_i \end{cases}$$

② In contrast, if we assume an order of the items and consider them from end to beginning, then the subproblem can be set as "selecting items as expensive as possible from $\{1,2,...,i\}$ with weight limit w" and we have the following recursion:

$$OPT(\{1, 2, ..., n\}, W) = \max \begin{cases} OPT(\{1, 2, ..., n-1\}, W) \\ OPT(\{1, 2, ..., n-1\}, W - w_n) + v_n \end{cases}$$

• In fact, the first one exploits recursion over sets, which leads to an exponential number of subproblems. In contrast, the second one is a recursion over sequences and the number of subproblems is only O(nW).

Extension: The first public-key encryption system

 Cryptosystems based on the knapsack problem were among the first public key systems to be invented, and for a while were considered to be among the most promising. However, essentially all of the knapsack cryptosystems that have been proposed so far have been broken. These notes outline the basic constructions of these cryptosystems and attacks that have been developed on them.

See The Rise and Fall of Knapsack Cryptosystems for details.

VERTEX COVER: recursion over trees

VERTEX COVER Problem

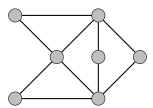
 Practical problem:
 Given n sites connected with paths, how many guards (or cameras) should be deployed on sites to surveille all the paths?

Formalized Definition:

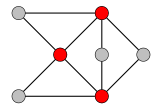
Input: Given a graph $G = \langle V, E \rangle$

Output: the minimum of nodes $S\subseteq V$, such that each edge has at least one of its endpoints in S

• For example, how many nodes are needed to cover all edges in the following graph?

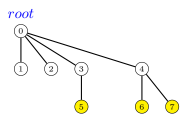


Defining general form of sub-problems



- The nodes in red form a vertex cover.
- VERTEX COVER is a hard problem for general graph.
- However, it is easy to find the minimum vertex cover for trees.

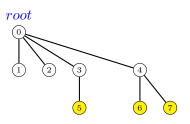
Optimal sub-structure property



- It is not easy to solve the problem with n nodes. Let's whether it is possible to reduce into smaller sub-problems.
- Solution: selection a subset of nodes. Describe the solving process as as a process of multistage decisions. At each decision stage, we decide whether a node should be selected.
- Suppose we have already worked out the optimal solution.
- Consider the first decision, i.e. whether the optimal solution contains the root node or not. The decision has two options:
 - SELECT: it suffices to consider the sub-trees;
 - ② ABANDON: we should select all the children nodes, and then consider all grand-children.

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An example



- In both cases, the original problem is reduced into smaller sub-problems.
- General form of sub-problems: find minimum vertex cover on a tree rooted at node v. Let's denote the optimal solution as OPT(v).
- Thus we have the following recursion:

$$OPT(root) = \min \begin{cases} 1 + \sum_{c} OPT(c) & c \text{ is a child} \\ \#children + \sum_{g} OPT(g) & g \text{ is a grand-child} \end{cases}$$

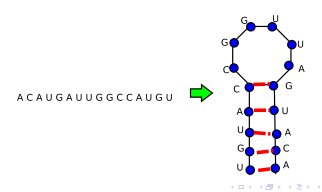
• Time complexity: O(n)



RNA SECONDARY STRUCTURE PREDICTION: recursion over trees

RNA secondary structure

- RNA is a sequence of nucleic acids. It will automatically form structures in water through the formation of bonds A-U and C-G.
- The native structure is the conformation with the lowest energy. Here, we simply use the number of base pairs as the energy function.



Formulation

INPUT:

A sequence in alphabet $\Sigma = \{A, U, C, G\}$;

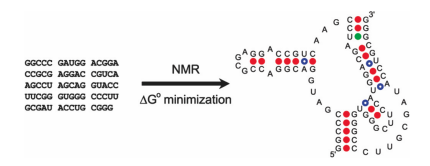
OUTPUT:

A pairing scheme with the maximum pairing number

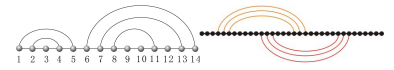
Requirements of base pairs:

- Watson-Crick pair: A pairs with U, and C pairs with G;
- There is no base occurring in more than 1 base pairs;
- No cross-over (nesting): there is no crossover under the assumption of free pseudo-knots.
- And two bases $i, j \ (|i-j| \le 4)$ cannot form a base pair.

An example

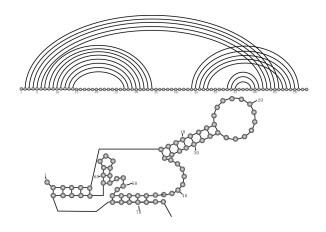


Nesting and Pseudo-knot



Left: nesting of base pairs (no cross-over); Right: pseudo-knots (cross-over).

Feymann graph

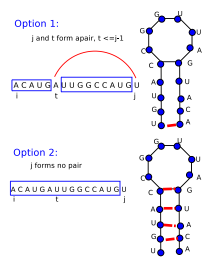


Feymann graph: an intuitive representation form of RNA secondary structure, i.e. two bases are connected by an edge if they form a Watson-Crick pair.

Defining general form of sub-problems

- Solution: a set of nested base pairs. Describe the solving process as as a process of multistage decisions. At the *i*-th decision stage, we determine whether base *i* forms pair or not.
- Suppose we have already worked out the optimal solution.
- ullet Consider the first decision made for base n. There are two options:
 - **1** Base n pairs with a base i: we should calculate optimal pairs for regions i+1..n-1 and 1..i-1. Note that these two sub-problems are independent due to the "nested" property.
 - ② Base n doesn't form a pair: we should calculate optimal pairs for regions 1..n-1.
- Thus we can design the general form of sub-problems as: to calculate the optimal pairs for region i...j. (Denote the optimal solution value as: OPT(i,j).)

Optimal sub-structure property



Optimal substructures property:

 $OPT(i,j) = \max\{OPT(i,j-1), \max_t\{1 + OPT(i,t-1) + OPT(t+1,j-1)\}\}$, where the second max takes over all

Algorithm

```
\begin{array}{lll} {\rm RNA2D}(n) \\ {\rm 1:} & {\rm Initialize \ all \ } OPT[i,j] \ {\rm with \ } 0; \\ {\rm 2:} & {\bf for \ } i=1 \ {\rm to \ } n \ {\bf do} \\ {\rm 3:} & {\bf for \ } j=i+5 \ {\rm to \ } n \ {\bf do} \\ {\rm 4:} & & OPT[i,j] = \max\{OPT[i,j-1], \max_t\{1+OPT[i,t-1]+OPT[t+1,j-1]\}\}; \\ {\rm 5:} & & /* \ t \ {\rm and \ } j \ {\rm can \ form \ Watson-Crick \ base \ pair. \ */} \\ {\rm 6:} & {\bf end \ for \ } \\ {\rm 7:} & {\bf end \ for \ } \end{array}
```

An example: Step 1

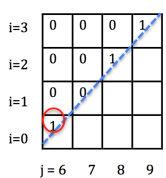
INPUT:

ACCGGUAGU

i=3	0	0	0	
i=2	0	0		
i=1	0			
i=0				
	 j = 6	7	8	9

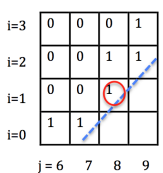
INPUT:

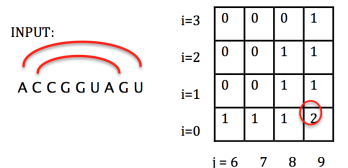






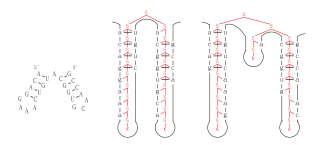






Time complexity: $O(n^3)$.

Extension: RNA is a good example of SCFG.



(see extra slides)

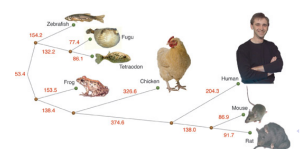
SEQUENCE ALIGNMENT problem: recursion over sequence pairs

Practical problem: genome similarity

 To identify homology genes of two species, say Human and Mouse. Human and Mouse NHPPEs (in KRAS genes) show a high sequence homology (Ref: Cogoi, S., et al. NAR, 2006).

GGGCGGTGTG
| | | | | | | | |
GGGAGG-GAG

 Having calculating the similarity of genomes of various species, a reasonable phylogeny tree can be estimated (See https://www.llnl.gov/str/June05/Ovcharenko.html)



Practical problem: spell tool to correct typos

• When you type in "OCURRANCE", spell tools might guess what you really want to type through the following alignment:

```
O-CURRANCE
| | | | | | | | |
OCCURRENCE
```

Here, '|' represent identical characters.

 This alignment suggests that "OCURRANCE" is very similar to "OCCURRENCE", i.e., "OCURRANCE" is generated from "OCCURRENCE" using several INS/DEL/MUTATION operations.

What is an alignment?

- Alignment is introduced to describe the generating process of an erroneous word from the correct word using a series of INS/DEL/MUTATION operations.
- For this aim, we make the two sequences to have the same length through adding space '-' at appropriate positions, which changes S to S', and changes T to T' with identical length |S'| = |T'|.

S': O-CURR-ANCE
| | | | | | | |
T': OCCURRE-NCE

- The spaces '-' are introduced to describe the generating process of S from T. There are three cases of aligned characters:
 - S'[i] = '-': S'[i] is simply a DELETION of T'[i].
 - 2 T'[i] = '-': S'[i] is simply an INSERTION.
 - **3** Otherwise, S'[i] is a copy of T'[i] (with possible MUTATION).

Measuring the possibility of each generating process

• For each generating process, we can measure its possibility as:

$$s(S,T) = \sum_{i=1}^{|S'|} s(S'[i], T'[i])$$

- Here we assume a simple setting of s(a,b) as follows:
 - MATCH: +1, e.g. s(C', C') = 1.
 - ② MISMATCH: -1, e.g. s(`E', `A') = -1.
 - **3** Insertion/Deletion: -3, e.g. s(C', C', C') = -3.

 $^{^1}$ Ideally, the score function is designed such that s(S,T) is proportional to $\log\Pr[S$ is generated from T]. See extra slides for the statistical model for sequence alignment, and better similarity definition, say BLOSUM62, PAM250 substitution matrix, etc.

Alignment is useful

- Observation 1: Using alignment, we can determine the most likely source of "OCURRANCE".
 - \bullet T = "OCCURRENCE":

$$s(S', T') = 1 - 3 + 1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 = 4$$

 \mathbf{O} T = "OCCUPATION":

$$s(S', T') = 1 + 1 - 3 + 1 - 3 - 1 + 1 - 3 - 3 - 3 + 1 - 3 - 3 = -28$$

 Thus, it is more likely that "OCURRANCE" comes from "OCCURRENCE" relative to "OCCUPATION".



Alignment is useful cont'd

- Observation 2: In addition, we can also determine the most likely operations changing "OCCURRENCE" into "OCURRANCE".
 - Alignment 1:

$$s(S',T') = 1 - 3 + 1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 = 4$$

② Alignment 2:

$$s(S',T') = 1 - 3 + 1 + 1 + 1 - 3 - 3 + 1 + 1 + 1 = -1$$

 Thus, the first alignment might describes the real generating process of "OCURRANCE" from "OCCURRENCE".



SEQUENCE ALIGNMENT: formulation

INPUT:

Two sequences S and T, |S| = m, and |T| = n;

OUTPUT:

To identify an alignment of ${\cal S}$ and ${\cal T}$ that maximizes a pre-defined scoring function.

Note: for the sake of simplicity, the following indexing schema is used: $S = S_1 S_2 ... S_m$.

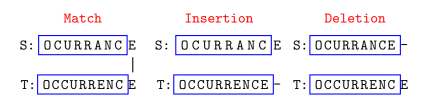
The general form of sub-problems and recursions I

- It is not easy to consider long sequences directly. Let's consider whether it is possible to reduce into smaller subproblem.
- Solution: insert '-' at appropriate positions to represent the generating process of S from T. Let's describe the solving process as a process of **multistage decisions**. At each decision stage, we decide whether S_i aligns with T_j (match), S_i aligns with a '-' (insert), or T_j aligns with a '-' (deletion).

	Match	Insertion	Deletion
S:	OCURRANC E	S: OCURRANC E	S: OCURRANCE -
Т:	OCCURRENC E	T: OCCURRENCE -	T: OCCURRENC E

The general form of sub-problems and recursions II

- Suppose we have already worked out the optimal solution. Consider the **first** decision made for S_m . There are three cases:
 - S_m comes from T_n : represented as aligning S_m with T_n . Then it suffices to align S[1..m-1] and T[1..n-1].
 - ② S_m is an INSERTION: represented as aligning S_m with a space '-'. Then it suffices to align S[1..m-1] and T[1..n].
 - **3** S_m comes from T[1..n-1]: represented as aligning T_n with a space '-'. Then it suffices to align S[1..m] and T[1..n-1].



The general form of sub-problems and recursions III

- Thus, we can design the general form of sub-problems as: alignment a **prefix** of S (denoted as S[1..i]) and **prefix** of T (denoted as T[1..j]). Denote the optimal solution value as OPT(i,j).
- We can prove the following optimal substructure property:

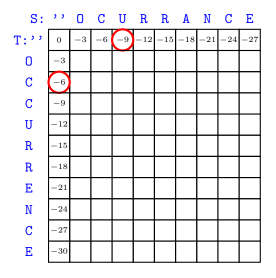
$$OPT(i, j) = \max \left\{ \begin{array}{l} s(S_i, T_j) + OPT(i - 1, j - 1) \\ s(`_', T_j) + OPT(i, j - 1) \\ s(S_i, `_') + OPT(i - 1, j) \end{array} \right.$$

Needleman-Wunsch algorithm [1970]

```
NEEDLEMAN-WUNSCH(S,T)
 1: for i = 0 to m; do
 2: OPT[i, 0] = -3 * i;
 3: end for
 4: for j = 0 to n; do
 5: OPT[0, j] = -3 * j;
 6: end for
 7: for i=1 to m do
 8: for j=1 to n do
       OPT[i, j] = \max\{OPT[i-1, j-1] + s(S_i, T_i), OPT[i-1]\}
        [1, j] - 3, OPT[i, j - 1] - 3;
     end for
10:
11: end for
12: return OPT[m, n];
Note: the first row is introduced to describe the alignment of
```

prefixes T[1..i] with an empty sequence ϵ , so does the first column.

The first row/column of the alignment score matrix



Score: s("OCU", "") = -9Alignment: S' = OCU Score: s("", "0C") = Alignment: S'= -T'= 0C

Why should we introduce the first row/column?

S:	,,	0	C	U	R	R	A	N	C	E
T:''	0	-3	-6	-9	-12	-15	-18	-21	-24	-27
0	-3	1	-2	-5	-8	-11	-14	-17	-20	-23
C	-6	-2	2	-1	-4	-7	-10	-13	-16	-19
C	-9	-5	-1	1	-2	-5	-8	-11	-12	-15
U	-12	-8	-4	0	0	-3	-6	-9	-12	13
R	-15	-11	-7	-3	1	1	-2	-5	-8	-11
R	-18	-14	-10	-6	-2	2	0	-3	-6	-9
E	-21	-17	-13	-9	-5	-1	1	-1	-4	-5
N	-24	-20	-16	-12	-8	-4	-2	2	-1	-4
C	-27	-23	-19	-15	-11	-7	-5	-1	3	0
E	-30	-26	-22	-18	-14	-10	-8	-4	0	4

Score:
$$s("0C", "0") = max \begin{cases} s("0C", "") & -3 & (=-9) \\ s("0", "") & -1 & (=-4) \\ s("0", "0") & -3 & (=-2) \end{cases}$$
Alignment: $S' = 0$

General cases

S:	, ,	0	C	U	R	R	A	N	C	E
T:''	0	-3	-6	-9	-12	-15	-18	-21	-24	-27
0	-3	1	-2	-5	-8	-11	-14	-17	-20	-23
C	-6	-2	2	-1	-4	-7	-10	-13	-16	-19
C	-9	-5	-1	1	-2	-5	-8	-11	-12	-15
U	-12	-8	-4	0	0	-3	-6	-9	-12	13
R	-15	-11	-7	-3	1	1	-2	-5	-8	-11
R	-18	-14	-10	-6	-2	2	0	-3	-6	-9
E	-21	-17	-13	-9	-5	-1	1	-1	-4	-5
N	-24	-20	-16	-12	-8	-4	-2	2	-1	-4
C	-27	-23	-19	-15	-11	-7	-5	-1	3	0
E	-30	-26	-22	-18	-14	-10	-8	-4	0	4

Score:
$$s("OCUR", "OC") = max \begin{cases} s("OCUR", "O") & -3 & (=-11) \\ s("OCU", "O") & -1 & (=-6) \\ s("OCU", "OC") & -3 & (=-4) \end{cases}$$
Alignment: $S' = OCUR$

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The final entry

S:	, ,	0	C	U	R	R	A	N	C	E
T:''	0	-3	-6	-9	-12	-15	-18	-21	-24	-27
0	-3	1	-2	-5	-8	-11	-14	-17	-20	-23
C	-6	-2	2	-1	-4	-7	-10	-13	-16	-19
C	-9	-5	-1	1	-2	-5	-8	-11	-12	-15
U	-12	-8	-4	0	0	-3	-6	-9	-12	13
R	-15	-11	-7	-3	1	1	-2	-5	-8	-11
R	-18	-14	-10	-6	-2	2	0	-3	-6	-9
E	-21	-17	-13	-9	-5	-1	1	-1	-4	-5
N	-24	-20	-16	-12	-8	-4	-2	2	-1	-4
C	-27	-23	-19	-15	-11	-7	-5	-1	3	0
E	-30	-26	-22	-18	-14	-10	-8	-4	0	4

```
Score: s("OCURRANCE", "OCCURRENCE") = max 
\begin{cases} s("OCURRANCE", "OCCURRENC") & -3 & (=-3) \\ s("OCURRANC", "OCCURRENC") & +1 & (=4) \\ s("OCURRANCE", "OCCURRENCE") & = -3 & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-3) & (=-
```

T'= OCCURRENCE

Question: how to find the alignment with the highest score?

Find the optimal alignment via backtracking

S:	, ,	0	C	U	R	R	A	N	C	E
T:''	0	-3	-6	-9	-12	-15	-18	-21	-24	-27
0	-3	1	-2	-5	-8	-11	-14	-17	-20	-23
C	-6	-2	2	-1	-4	-7	-10	-13	-16	-19
C	-9	-5	-1	1	-2	-5	-8	-11	-12	-15
U	-12	-8	-4	0	0	-3	-6	-9	-12	13
R	-15	-11	-7	-3		1	-2	-5	-8	-11
R	-18	-14	-10	-6	-2	2	0	-3	-6	-9
Е	-21	-17	-13	-9	-5	-1		-1	-4	-5
N	-24	-20	-16	-12	-8	-4	-2	2	-1	-4
C	-27	-23	-19	-15	-11	-7	-5	-1	3	0
E	-30	-26	-22	-18	-14	-10	-8	-4	0	4

Optimal Alignment: S'= O-CURRANCE T'= OCCURRENCE

Optimal alignment versus sub-optimal alignments

- In practice, there are always multiple alignments with nearly the same score as the optimal alignment. Such alignments are called sub-optimal alignments and can be divided into the following two categories.
 - Similar sub-optimal alignments: Some sub-optimal alignments
 differ from the optimal alignment in a few positions. Because
 variations might occur independently at different positions, the
 number of sub-optimal alignments grow exponentially with the
 difference to the optimal alignment. The typical sub-optimal
 alignments can be obtained through sampling over the
 dynamic programming matrix.
 - Distinct sub-optimal alignments: The other sub-optimal alignments differ completely from the optimal alignment.
 These sub-optimal alignments frequently appear when one or both sequences have repeats.

Please refer to Biological Sequence Analysis: Probabilistic Models of Proteins and Nucleic Acids for details.

Finding sub-optimal alignments

- To find similar sub-optimal alignments, we trace back through the dynamic programming matrix using sampling technique, i.e., instead of taking the highest scoring option at each step, we make probabilistic choices bases on the value of the three options.
- To find the next best alignment sharing no aligned residue pairs with the optimal alignment, we can re-calculate the dynamic programming matrix with cells corresponding to the pairs in the optimal alignment set to zero. The resulting matrix can be used to find the second best alignment. Another approach is extending cells of OPT table to store top alignments.

Space efficient algorithm: reducing the space requirement from O(mn) to O(m+n) (D. S. Hirschberg, 1975)

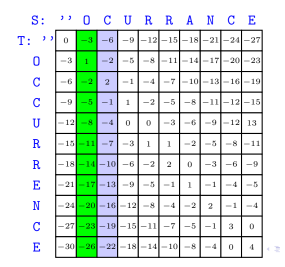
Key observation 1: it is easy to calculate the final score
 OPT(S,T) only, i.e. the alignment information are not
 recorded.

S:	, ,	0	C	U	R	R	A	N	C	E
T:''	0	-3	-6	-9	-12	-15	-18	-21	-24	-27
0	-3	1	-2	-5	-8	-11	-14	-17	-20	-23
C	-6	-2	2	-1	-4	-7	-10	-13	-16	-19
C	-9	-5	-1	1	-2	-5	-8	-11	-12	-15
U	-12	-8	-4	0	0	-3	-6	-9	-12	13
R	-15	-11	-7	-3	1	1	-2	-5	-8	-11
R	-18	-14	-10	-6	-2	2	0	-3	-6	-9
E	-21	-17	-13	-9	-5	-1	1	-1	-4	-5
N	-24	-20	-16	-12	-8	-4	-2	2	-1	-4
C	-27	-23	-19	-15	-11	-7	-5	-1	3	0
E	-30	-26	-22	-18	-14	-10	-8	-4	0	4

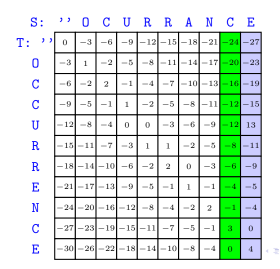
• Why? Only column j-1 is needed to calculate column i. Thus, we use two arrays score[1..n] and newscore[1..n] instead of the matrix OPT[1..n, 1..m].

S:	, ,	0	C	U	R	R	Α	N	C	E	_
T: ''	0	-3	-6	-9	-12	-15	-18	-21	-24	-27	
0	-3	1	-2	-5	-8	-11	-14	-17	-20	-23	
C	-6	-2	2	-1	-4	-7	-10	-13	-16	-19	
C	-9	-5	-1	1	-2	-5	-8	-11	-12	-15	
U	-12	-8	-4	0	0	-3	-6	-9	-12	13	
R	-15	-11	-7	-3	1	1	-2	-5	-8	-11	
R	-18	-14	-10	-6	-2	2	0	-3	-6	-9	
E	-21	-17	-13	-9	-5	-1	1	-1	-4	-5	
N	-24	-20	-16	-12	-8	-4	-2	2	-1	-4	
C	-27	-23	-19	-15	-11	-7	-5	-1	3	0	
E	-30	-26	-22	-18	-14	-10	-8	-4	0	4	4

• Why? Only column j-1 is needed to calculate column i. Thus, we use two arrays score[1..n] and newscore[1..n] instead of the matrix OPT[1..n, 1..m].



• Why? Only column j-1 is needed to calculate column i. Thus, we use two arrays score[1..m] and newscore[1..m] instead of the matrix OPT[1..m, 1..n].



Algorithm

$\label{eq:prefix_space_efficient_alignment} Prefix_Space_Efficient_Alignment(S, T, score)$

```
1: for i=0 to m do
2: score[i] = -3 * i;
3: end for
4: for i = 1 to m do
   newscore[0] = 0;
     for j=1 to n do
6:
       newscore[j] = \max\{score[j-1] + s(S_i, T_i), score[j] - 1\}
7:
        3, newscore[j-1]-3;
     end for
8:
   for j=1 to n do
9.
       score[j] = newscore[j];
10:
     end for
11:
12: end for
13: return score[n];
```

Technique 2: aligning suffixes instead of prefixes

 Key observation: Similarly, we can align suffixes of S and T instead of prefixes and obtain the same score and alignment.

4	0	-4	-10	-12	-16	-18	-22	-26	-30	0
5	3	-1	-7	-9	-13	-15	-19	-23	-27	C
3	6	2	-4	-6	-10	-12	-16	-20	-24	C
-1	2	5	-1	-3	-7	-9	-13	-17	-21	U
-5	-2	1	4	0	-4	-6	-10	-14	-18	R
-9	-6	-3	0	3	-1	-3	-7	-11	-15	R
-13	-10	-7	-4	-1	2	0	-4	-8	-12	E
-15	-12	-9	-6	-3	0	3	-1	-5	-9	N
-19	-16	-13	-10	-7	-4	-1	2	-2	-6	C
-23	-20	-17	-14	-11	-8	-5	-2	1	-3	E
-27	-24	-21	-18	-15	-12	-9	-6	-3	0	, , T
0	C	U	R	R	A	N	C	E	,,	S
									< □ →	∢ 🗇 ▶ ∢

Final difficulty: identify optimal alignment besides score

- However, the optimal alignment cannot be restored via backtracking since only the recent two columns of the matrix were kept.
- ② A clever idea: Suppose we have already obtained the optimal alignment. Let's consider the position where $S_{\left[\frac{m}{2}\right]}$ is aligned to (denoted as q). We have:

$$OPT(S,T) = OPT(S[1..\frac{m}{2}], T[1..q]) + OPT(S[\frac{m}{2} + 1..m], T[q + 1..n])$$

3 The equality holds due to the linearity of s(S,T), and $\frac{m}{2}$ is chosen for the sake of time-complexity analysis.

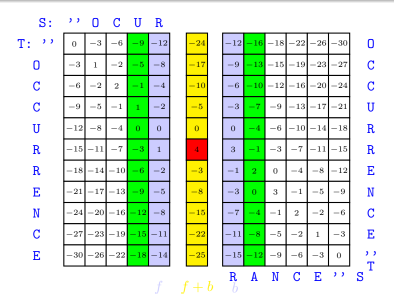
$$\frac{m}{2}$$
S: OCUR RANCE
T: OCCUR RENCE
 $1 < q < n$

Hirschberg's algorithm for alignment

LINEAR_SPACE_ALIGNMENT(S, T)

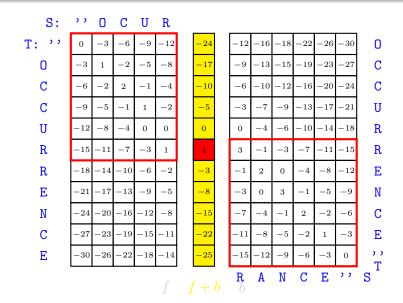
- 1: Allocate two arrays f and b; each array has a size of m .
- 2: Prefix_Space_Efficient_Alignment($S[1...\frac{m}{2}], T, f$);
- 3: Suffix_Space_Efficient_Alignment($S[\frac{m}{2}+1..m], T, b$);
- 4: Let $q = argmax_i(f[i] + b[i]);$
- 5: Free arrays f and b;
- 6: Record aligned-position $<\frac{m}{2},q>$ in an array A;
- 7: LINEAR_SPACE_ALIGNMENT $(S[1...\frac{m}{2}], T[1..q]);$
- 8: LINEAR_SPACE_ALIGNMENT($S[\frac{m}{2}+1..m], T[q+1..n]$);
- 9: **return** A;
 - Key observation: at each iteration step, only 2n space is needed.
 - How to determine q? Identifying the largest entry in f[i] + b[i].

Step 1: Determine the optimal aligned position of $S_{\left[\frac{m}{2}\right]}$



The value of the largest item is 4, which is actually the optimal score

Step 2: Recursively solve sub-problems



We generate two sub-problems according to the position of the

Space complexity analysis

- The total space requirement: O(m+n).
 - Prefix_Space_Efficient_Alignment($S[1..\frac{m}{2}), T, f$) needs only O(n) space;
 - SUFFIX_SPACE_EFFICIENT_ALIGNMENT $(S[\frac{m}{2}+1..m),T,b)$ needs only O(n) space;
 - Line 4 (Record $<\frac{n}{2},q>$ in array A) needs only O(m) space;

Time complexity analysis

Theorem

Algorithm Linear_Space_Alignment (S, T) still takes O(mn) time.

Proof.

- The algorithm implies the following recursion: $T(m,n) = cmn + T(\frac{m}{2},q) + T(\frac{m}{2},n-q);$
- Difficulty: we have no idea of q before algorithm ends; thus, the master theorem cannot apply directly. **Guess and substitution!!!**
 - Guess: $T(m', n') \le km'n'$ follows for any m' < m and n' < n.
 - Substitution:(猜啊)

SIGN HERE

$$T(m,n) = cmn + T(\frac{m}{2},q) + T(\frac{m}{2},n-q)$$
 (7)

$$\leq cmn + kq\frac{m}{2} + k(n-q)\frac{m}{2} \tag{8}$$

$$= cmn + kq\frac{m}{2} + kn\frac{m}{2} - kq\frac{m}{2}$$
 (9)

$$\leq (c + \frac{k}{2})mn \tag{10}$$

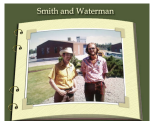
$$(set k = 2c) (11$$



Extended Reading 1: From global alignment to local alignment

From global alignment to local alignment: Smith-Waterman algorithm

- Global alignment: to identify similarity between two entire sequences.
- Local alignment: It is often that we wish to find similar segments (sub-sequences).
- Needleman-Wunsch global alignment algorithm was developed by biologists in 1970s, about twenty years later than Bellman-Ford algorithm was developed. Then Smith-Waterman local alignment algorithm was proposed (Please refer to Smith and Waterman1981 for details.).



Local alignment vs. global alignment: two difference

 The objective of local alignment is to identify similar segments of two sequence. The other regions can be treated as independent and thus form "random matches". To distinguish random matches and true matches, the scoring schema was designed to assign random matches with negative expected score, i.e.,

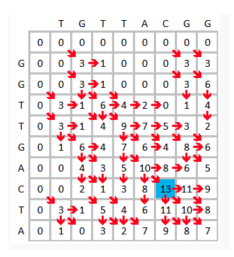
$$\sum_{a,b} q_a q_b s(a,b) < 0$$

The recursion was changed by adding an extra possibility:

$$OPT(i, j) = \max \begin{cases} \frac{\mathbf{0}}{s(S_i, T_j) + OPT(i - 1, j - 1)} \\ s(`_', T_j) + OPT(i, j - 1) \\ s(S_i, `_') + OPT(i - 1, j) \end{cases}$$

Taking the option 0 corresponds to starting a new alignment: if we obtain a negative score for OPT(i,j), this means that the subsequences S[1..i] and T[1..j] are independent. Thus, it is better to start a new alignment rather than extend the old one.

An example



• Note that the consequence of the 0 option is that the top row and left column are set as 0.

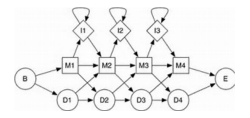
Local alignment vs. global alignment: two difference

		Т	G	Т	Т	Α	С	G	G
	0	0	0	0	0	0	0	0	0
G	0	0	3	1	0	0	0	3	3
G	0	0	3	1	0	0	0	3	6
Т	0	3	1	6	4	2	0	1	4
Т	0	3	1	4	9	7	5	3	2
G	0	1	6	4	7	6	4	8	6
Α	0	0	4	3	5	10	8	6	5
С	0	0	2	1	3	8	13	11	9
Т	0	3	1	5	4	6	11	10	8
Α	0	1	0	3	2	7	9	8	7

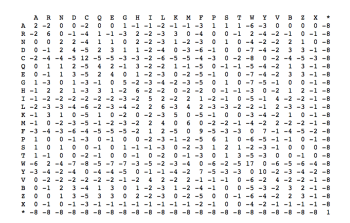
- As the local alignment aims to represent similar segments, it can end at any cell in the matrix. Thus, instead of taking the bottom-right corner in global alignment, we look for the highest value in the matrix and start backtracking from there.
- The backtrack ends when we meets a cell with value 0, which_{108/154}

Extended Reading 2: How to derive a reasonable scoring schema?

Profile-HMM: a generative model of multiple-sequence alignment



PAM250: one of the most popular substitution matrices in Bioinformatics



Please refer to "PAM matrix for Blast algorithm" (by C. Alexander, 2002) for the details to calculate PAM matrix.

Extended Reading 3: How to measure the significance of an alignment?

Measure the significance of a segment pair

• When two random sequences of length m and n are compared, the probability of finding a pair of segments with a score greater than or equal to S is $1-e^{-y}$ where $y=Kmne^{-\lambda S}$.

Please refer to Altschul1990 for details.

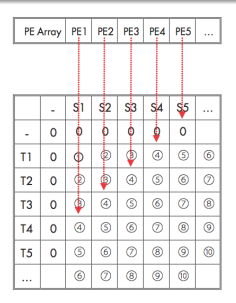
Extended Reading 4: An FPGA implementation of Smith-Waterman algorithm

The potential parallelity of SmithWaterman algorithm

	-	S 1	S2	S3	S4	S5	
-	0	Q	0	0	0	0	
T1	0-	1	2	3	4	(5)	6
T2	0	2	3	4	(5)	6	7
Т3	0	3	4	D	6	7	8
T4	0	4	(5)	6	Ø	8	9
T5	0	(5)	6	7	8	Ø	10
		6	7	8	9	10	×

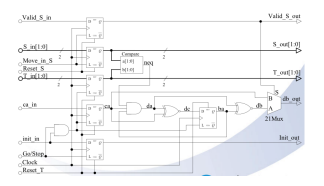
For example, in the first cycle, only one element marked as (1) could be calculated. In the second cycle, two elements marked as (2) could be calculated. In the third cycle, three elements marked as (3) could be calculated, etc., and this feature implies that the algorithm has a very good potential parallelity.

Mapping Smithg-Waterman algorithm on PE



Pacantiques bla Cunarcomputing Platform for datails

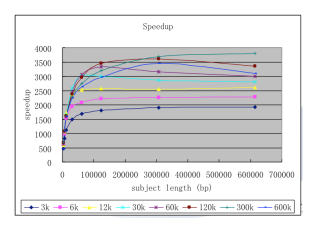
PE design of a card for Dawning 4000L



Smith-Waterman card for Dawning 4000L



Performance of Dawning 4000L



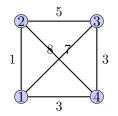
2

Bellman-Held-Karp for TSP problem: recursion over graphs

TRAVELLING SALESMAN PROBLEM

INPUT: a list of n cities (denoted as V), and the distances between each pair of cities d_{ij} $(1 \le i, j \le n)$;

OUTPUT: the shortest tour that visits each city exactly once and returns to the origin city



#Tours: 6

- Tour 1: $1 \to 2 \to 3 \to 4 \to 1$ (12)
- Tour 2: $1 \rightarrow 2 \rightarrow 4 \rightarrow 3 \rightarrow 1$ (21)
- Tour 3: $1 \to 3 \to 2 \to 4 \to 1$ (23)
-

Consider a tightly related problem

Definition

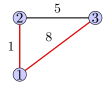
s(S,e)= the minimum distance, starting from city 1, visiting all cities in S, and finishing at city $e\in S$.



- It suffices to calculate s(S,e) for any $S \in \{1,2,...,n\}$ and city e since:
 - There are 3 cases of the city from which we return to 1.
 - Thus, the shortest tour can be calculated as: $\min\{D(\{1,2,3,4\},2)+d_{2,1},\\D(\{1,2,3,4\},3)+d_{3,1},\\D(\{1,2,3,4\},4)+d_{4,1}\}$

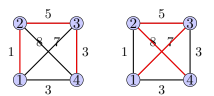
Let's start from the smallest problem

 \bullet It is trivial to calculate D(S,e) when S consists of only 1 cities.



- $D(\{2\},2)=d_{12};$
- $D({3},3)=d_{13};$
- But how to solve a large problem, say $D(\{2,3,4\},4)$?

Divide a larger problem into smaller problems



- $D(\{2,3,4\},4) = \min\{D(\{2,3\},3) + d_{34}, D(\{2,3\},2) + d_{24}\};$
- Optimal substructure property:

$$s(S,e) = \begin{cases} d_{1e} & \text{if } S = \{e\} \\ \min_{m \in S - \{e\}} (D(S - \{e\}, m) + d_{me}) & \text{otherwise} \end{cases}$$



Bellman-Held-Karp algorithm [1962]

```
function s(S, e)
 1: if S = \{e\} then
 2: return d_{1e};
 3: end if
 4: d=\infty:
 5: for all city m \in S, and m \neq e do
    if D(S - \{e\}, m) + d_{me} < d then
        d = D(S - \{e\}, m) + d_{me};
    end if
 8:
 9: end for
10: return d:
```

- Space complexity: $\sum_{k=2}^{n-1} k \binom{n-1}{k} + n 1 = (n-1)2^{n-2}$
- Time complexity: $\sum_{k=2}^{n-1} k(k-1) {n-1 \choose k} + n 1 = O(2^n n^2)$.

SINGLESOURCESHORTESTPATH problem: recursion over graphs

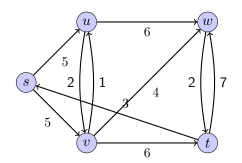
SINGLESOURCESHORTESTPATH problem

INPUT:

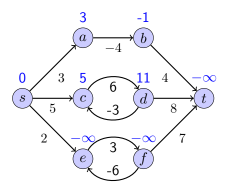
A directed graph G=< V, E>. Each edge e=(u,v) has a weight or distance d(u,v). Two special nodes: source s, and destination t;

OUTPUT:

A shortest path from s to t; that is, the sum weight of the edges is minimized.



SHORTESTPATH problem: cycles



- Here d(i, j) might be negative; however, there should be no negative cycle, i.e. the sum weight of edges in any cycle should be greater than 0.
- In fact, a negative cycle means an $-\infty$ shortest-path weight. Since e and f form a negative-weight cycle reachable from s, they have shortest-path weight of $-\infty$ from s.

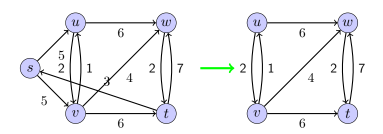
Trial 1: describing the sub-problem as finding the shortest path in a graph

Trial 1: recursion over graphs

- Solution: a path. Describe the solving process as series of decisions. At each decision stage, we need to determine an edge to the subsequent node.
- Suppose we have already worked out the optimal solution O.
- Consider the first decision in O. The options are:
 - All edges starting from s: Suppose we use an edge (s,v). Then it suffices to calculate the shortest path in graph G'=< V', E'>, where node s and related edges are removed.
- Thus, the original problem can be reduced into smaller sub-problems.
- General form of sub-problem: to find the shortest path from node v to t in graph G.

Trial 1: recursion over graphs cont'd

- General form of sub-problem: to find the shortest path from node v to t in graph G. Denote the optimal solution value as OPT(G,v).
- Optimal substructure: $OPT(G,s) = \min_{v:(s,v) \in E} \{OPT(G',v) + s(S,v)\}$



• Infeasible! The number of sub-problems is exponential.

Trial 2: another problem form with a new variable

Trial 2: simplifying sub-problem form via limiting path length

- Solution: the shortest path from node s to t is a path with at most n nodes (Why? no negative cycle ⇒ removing cycles in a path can shorten the path). Let's describe the solving process as a process of multistage decisions; at each decision stage, we decide the subsequent node from current node.
- Suppose we have already worked out the optimal solution O.
- Consider the first decision in O. The feasible options are:
 - All adjacent nodes of s: Suppose we choose an edge (s,v) to node v. Then the left-over is to find the shortest path from v to t via at most n-2 edges.
- Thus the general form of subproblem can be designed as: to find the shortest path from node v to t with **at most** k edges $(k \le n-1)$. Denote the optimal solution value as OPT(v,k).
- Optimal substructure:

$$OPT[v,k] = \min \begin{cases} OPT[v,k-1], \\ \min_{(v,w) \in E} \{OPT[w,k-1] + d(v,\bar{w})\} \end{cases}$$

Bellman-Ford algorithm [1956, 1958]

```
Bellman_Ford(G, s, t)
 1: for any node v \in V do
 2: OPT[v, 0] = \infty;
 3: end for
 4: for k = 0 to n - 1 do
 5: OPT[t, k] = 0;
 6: end for
 7: for k = 1 to n - 1 do
      for all node v (in an arbitrary order) do
        OPT[v,k] = \min \begin{cases} OPT[v,k-1] \\ \min_{(v,w) \in E} \{OPT[w,k-1] + d(v,w) \} \end{cases}
      end for
10:
11: end for
12: return OPT[s, n-1];
Note that the algorithm actually finds the shortest path from every
```

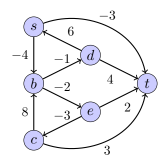
possible source to t (or from s to every possible destination).

Richard Bellman



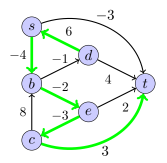
See "Richard Bellman on the birth of dynamic programming" (S. Dreyfus, 2002) and "On the routing problem" (R. Bellman, 1958) for details.

An example



Source node	k = 0	k = 1	k = 2	k = 3	k = 4	k = 5
t	0	0	0	0	0	0
8	-	-3	-3	-4	-6	-6
b	-	-	0	-2	-2	-2
c	-	3	3	3	3	3
d	-	4	3	3	2	0
e	1	2	0	0	0	0

Shortest path tree



Source node	k = 0	k=1	k=2	k=3	k = 4	k=5
t	0	0	0	0	0	0
8	-	-3	-3	-4	-6	-6
b	-	-	0	-2	-2	-2
c	-	3	3	3	3	3
d	-	4	3	3	2	0
e	-	2	0	0	0	0

Note: the shortest paths from all nodes to t form a shortest path tree.

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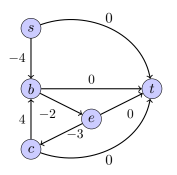
Time complexity

- Cursory analysis: $O(n^3)$. (There are n^2 subproblems, and for each subproblem, we need at most O(n) operations in line 7.
- 2 Better analysis: O(mn). (Efficient for sparse graph, i.e. $m << n^2$.)
 - For each node v, line 7 need $O(d_v)$ operations, where d_v denotes the degree of node v;
 - Thus the inner for loop (lines 6-8) needs $\sum_v d_v = O(m)$ operations;
 - Thus the outer for loop (lines 5-9) needs O(nm) operations.

Extension: detecting negative cycle

Theorem

If t is reachable from node v, and v is contained in a negative cycle, then we have: $\lim_{k\to\infty} OPT(v,k) = -\infty$.



Intuition: a traveling of the negative cycle leads to a shorter length. Say,

$$length(b \rightarrow t) = 0$$

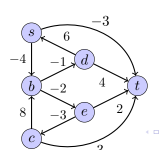
$$length(b \rightarrow e \rightarrow c \rightarrow b \rightarrow t) = -1$$

.

Corollary

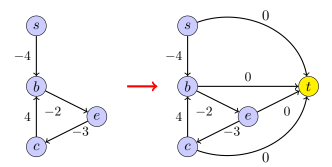
If there is no negative cycle in G, then for all node v, and $k \ge n$, OPT(v,k) = OPT(v,n).

Source node	k=0	k=1	k=2	k=3	k=4	k=5	k=6	k=7	k=8	k=9
t	0	0	0	0	0	0	0	0	0	О
s	-	-3	-3	-4	-6	-6	-6	-6	-6	-6
b	-	-	0	-2	-2	-2	-2	-2	-2	-2
c	-	3	3	3	3	3	3	3	3	3
d	-	4	3	3	2	0	0	0	0	C
e	-	2	0	0	0	0	0	0	0	C

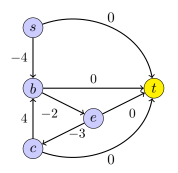


Detecting negative cycle via adding edges and a node t

- Expanding G to G' to guarantee that t is reachable from the negative cycle:
 - \bigcirc Adding a new node t;
 - ② For each node v, adding a new edge (v,t) with d(v,t)=0;
- Property: G has a negative cycle C (say, $b \rightarrow e \rightarrow c \rightarrow b$) \Rightarrow t is reachable from a node in C. Thus, the above theorem applies.



An example of negative cycle



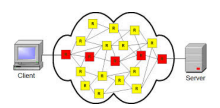
Source node	k=0	k=1	k=2	k=3	k=4	k=5	k=6	k=7	k=8	
t	0	0	0	0	0	0	0	0	0	
s	-	0	-4	-6	-9	-9	-11	-11	-12	
b	-	0	-2	-5	-5	-7	-7	-8	-8	
c	-	0	0	0	-2	-3	-3	-3	-4	
e	-	0	-3	-3	-5	-5	-6	-6	-6	

Application of Bellman-Ford algorithm: Internet router protocol

Internet router protocol

Problem statement:

- Each node denotes a route, and the weight denotes the **transmission delay** of the link from router *i* to *j*.
- ullet The objective to design a protocol to determine the quickest route when router s wants to send a package to t.



Internet router protocol: Dijkstra's algo vs. Bellman-Ford algo

- Choice: Dijkstra algorithm.
- However, the algorithm needs global knowledge, i.e. the knowledge of the whole graph, which is (almost) impossible to obtain.
- In contrast, the Bellman-Ford algorithm needs only local information, i.e. the information of its neighboorhood rather than the whole network.

Application: Internet router protocol

```
AsynchronousShortestPath(G, t)
```

1: Initially, set OPT[t, t] = 0, and $OPT[v, t] = \infty$; 2: Label node t as "active"; //Here a node v is called "active" if OPT[v,t] has been changed; 3: while exists an active node do Select an active node w arbitrarily; Remove w's "active" label: 5. 6: **for all** edges (v, w) (in an arbitrary order) **do** $OPT[v, t] = \min \begin{cases} OPT[v, t] \\ OPT[w, t] + d(v, w) \end{cases}$ 7: if OPT[v,t] was updated then 8: Label v as "active": 9: end if 10: end for 11: 12: end while

A related problem: LongestPath problem

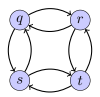
LONGESTPATH problem

INPUT:

A directed graph G=< V, E>. Each edge (u,v) has a distance d(u,v). Two nodes s and t.

OUTPUT:

The longest simple path from s to t.



Hardness: LongestPath problem is NP-hard. (Hint: it is obvious that LongestPath problem contains HAMILTONIANPATH as its special case.)

Subtlety: LongestPath problem | 1

- Divide: The subproblems are not independent.
- Consider dividing problem to find a path from q to t into two subproblems: to find a path from q to r, and to find a path from r to t.



- Suppose we have already solved the sub-problems. Let's try to combine the solutions to the two sub-problems:
 - $P(q,r) = q \to s \to t \to r$
 - $P(r,t) = r \to q \to s \to t$

We will obtain a path $q \to s \to t \to r \to q \to s \to t$, which is not simple.

Subtlety: LongestPath problem II

• In other words, the use of s in the first subproblem prevents us from using s in the second subproblem. However, we cannot obtain the optimal solution to the second subproblem without using s.

LongestPath versus ShortestPath

- In contrast, the ShortestPath problem does not have this difficulty.
- Why? The solutions to the subproblems **share no node**. Suppose the shortest paths P(q,r) and P(r,t) share a node $w(w \neq r)$. Then there will be a cycle $w \to \cdots \to r \to \cdots \to w$. Removing this cycle leads to a shorter path (no negative cycle). A contradiction.
- This means that the two subproblems are independent: the solution of one subproblem does not affect the solution to another subproblem.

A greedy algorithm exists when posing a stricter limit, i.e., all edges have a positive weight.

We will talk about this in next lectures.