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6.047 / 6.878 Computational Biology: Genomes, Networks, Evolution  
Fall 2008

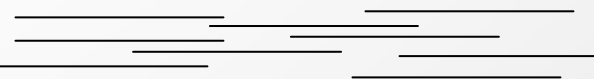
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# Sequence Alignment and Dynamic Programming

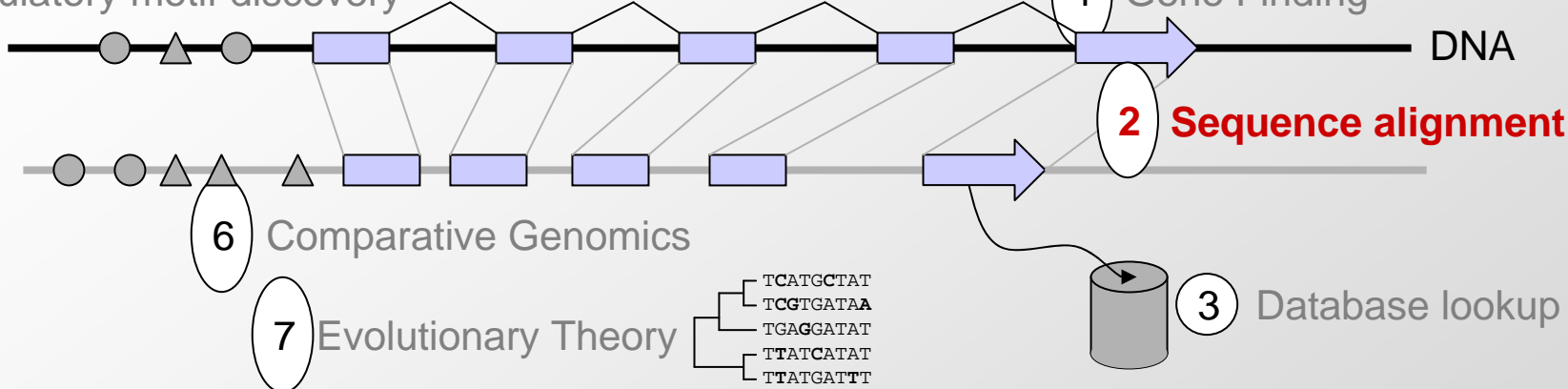
Tue Sept 9, 2008

# Challenges in Computational Biology

4 Genome Assembly

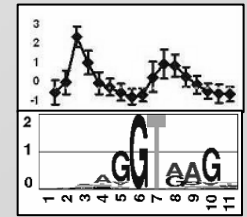
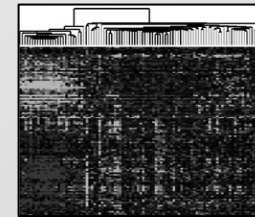


5 Regulatory motif discovery

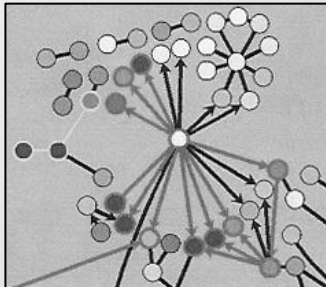


8 Gene expression analysis

RNA transcript



11 Protein network analysis



12 Regulatory network inference

13 Emerging network properties

9 Cluster discovery

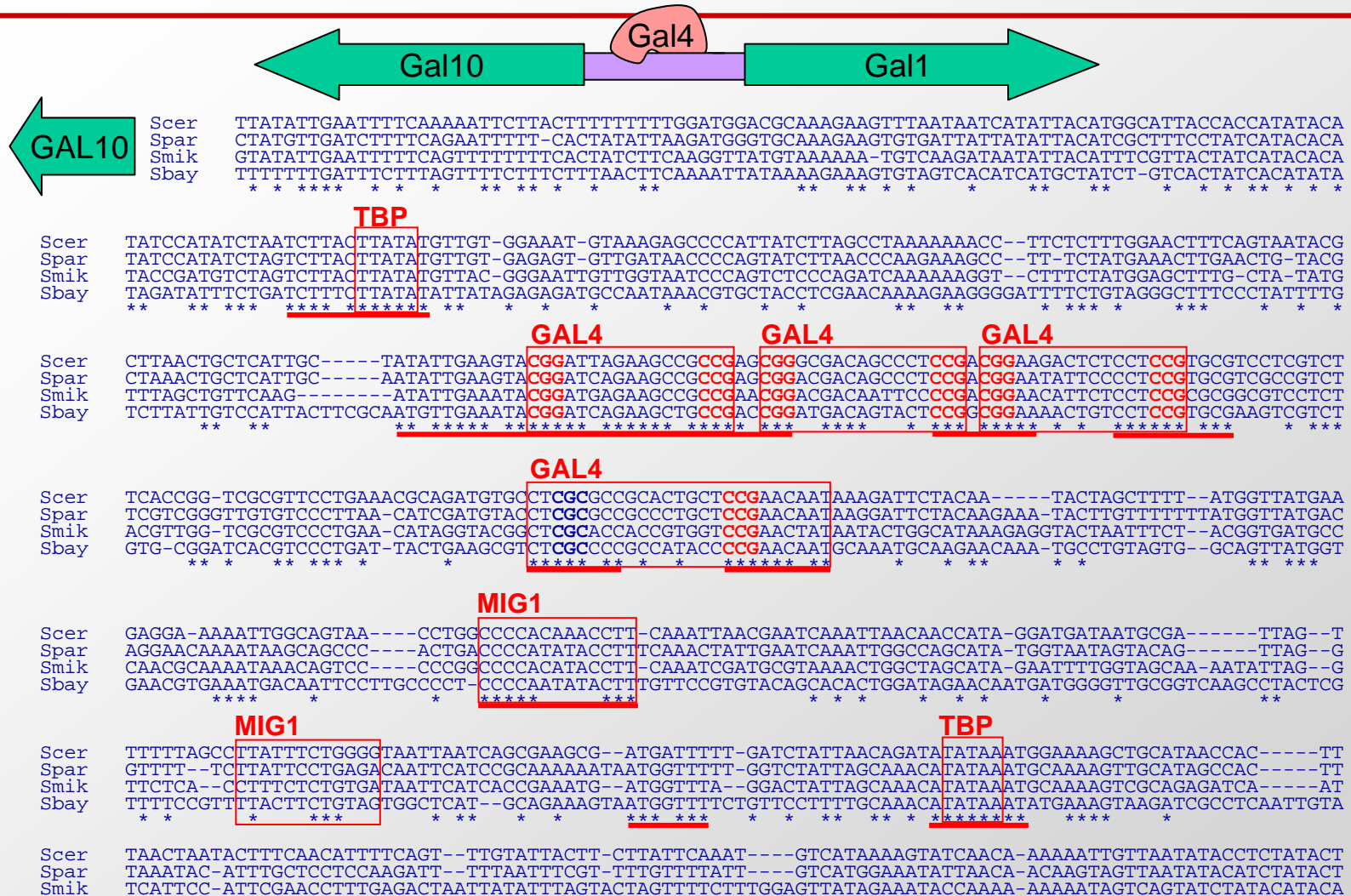
10 Gibbs sampling

# Reminder: Last lecture / recitation

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- Schedule for the term
  - ‘Foundations’ till midterm
  - ‘Frontiers’ lead to final project
  - Duality: basic problems / fundamental techniques
- Biology introduction
  - DNA, RNA, protein, transcription, translation
  - Why computational biology
- Today: Comparative genomics is everywhere!
  - Problem set 1: dating vertebrate whole-genome duplication
  - Problem set 2: discover genes using their conservation properties
  - Problem set 3: discover all motifs across entire yeast genome
  - Problem set 4: reversing human/mouse genome rearrangements

# Evolution preserved functional elements!



We can ‘read’ evolution to reveal functional elements

---

**Today's goal:**

**How do we actually align two genes?**

# Genomes change over time

begin



mutation



deletion



insertion



end



# Goal of alignment: Infer edit operations

---

begin

A	C	G	T	C	A	T	C	A
---	---	---	---	---	---	---	---	---

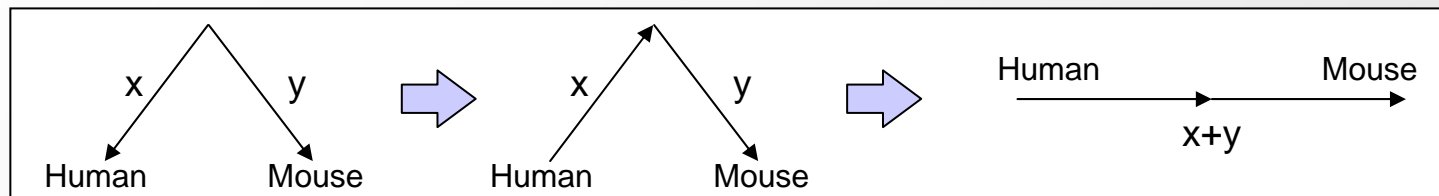


end

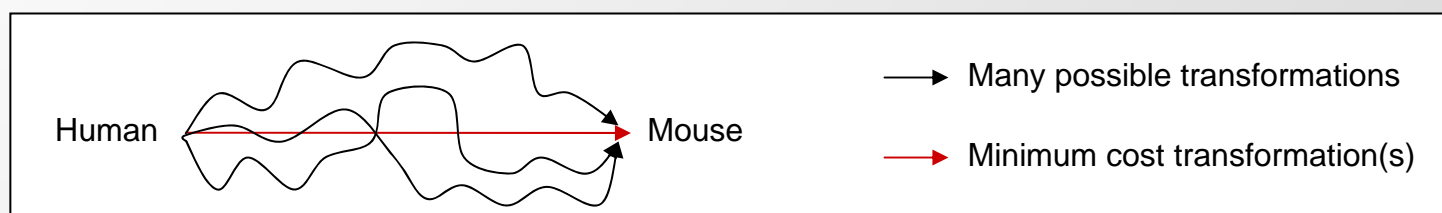
T	A	G	T	G	T	C	A
---	---	---	---	---	---	---	---

# From Bio to CS: Formalizing the problem

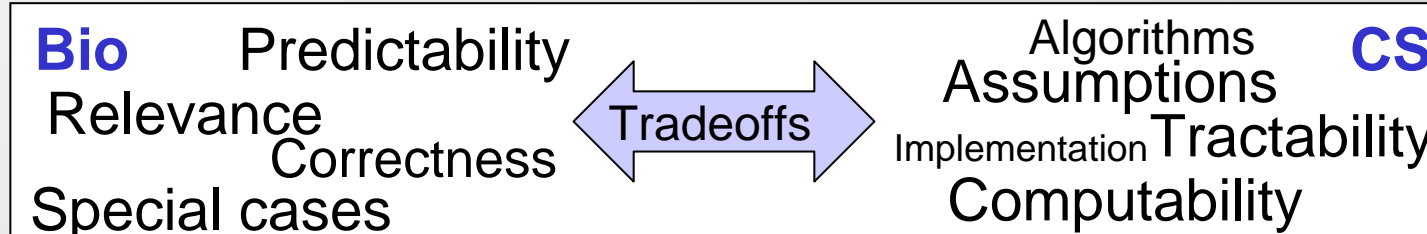
- Define set of evolutionary operations (insertion, deletion, mutation)
  - Symmetric operations allow time reversibility (part of design choice)



- Define optimality criterion (min number, min cost)
  - Impossible to infer exact series of operations (Occam's razor: find min)



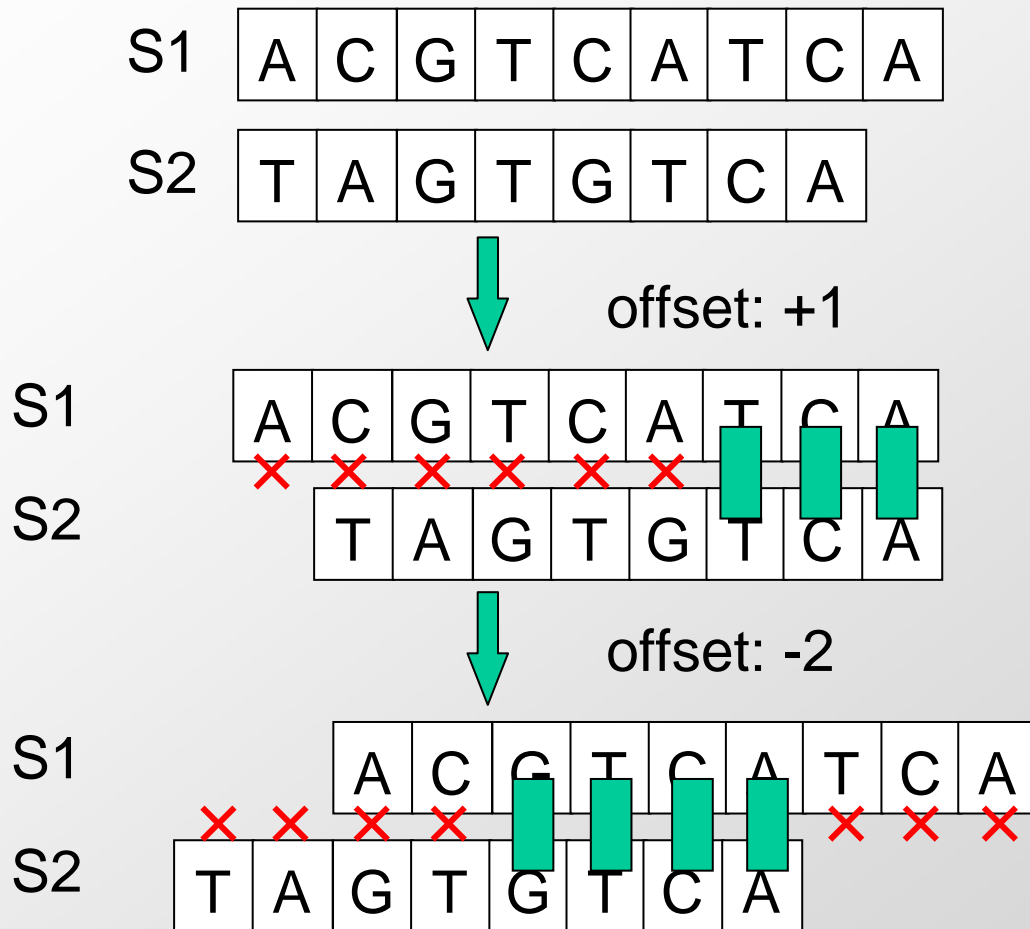
- Design algorithm that achieves that optimality (or approximates it)
  - Tractability of solution depends on assumptions in the formulation



Note: Not all decisions are conflicting (some are both relevant and tractable)  
(e.g. Pevzner vs. Sankoff and directionality in chromosomal inversions)

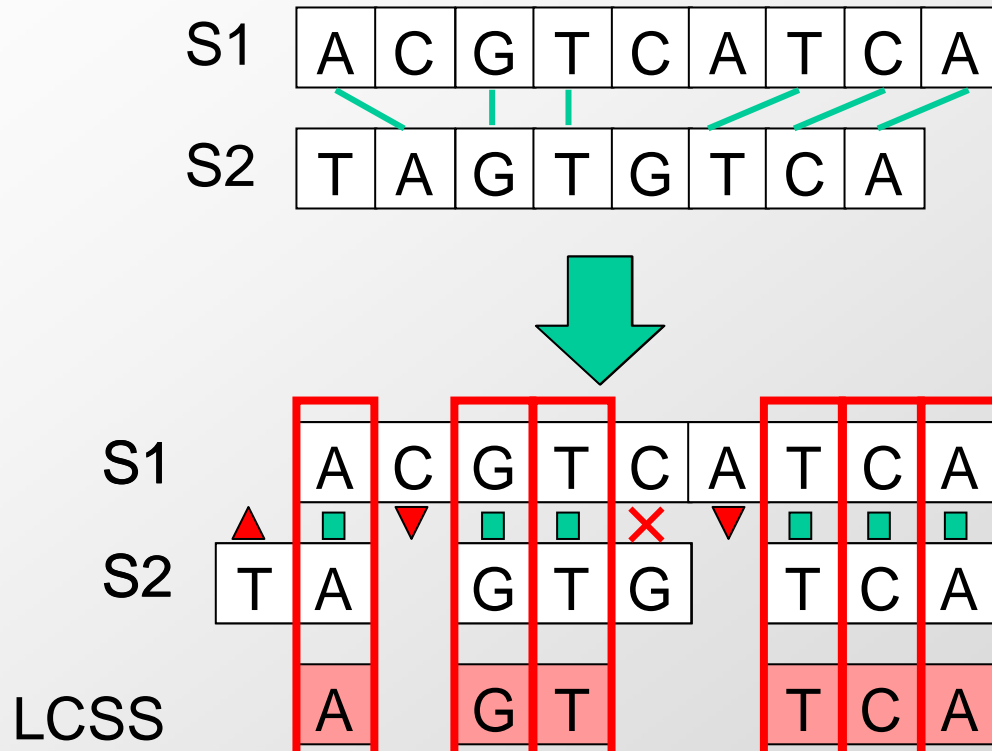
# Formulation 1: Longest common substring

- Given two possibly related strings S1 and S2
  - What is the longest common substring? (no gaps)



## Formulation 2: Longest common subsequence

- Given two possibly related strings S1 and S2
  - What is the longest common subsequence? (gaps allowed)



Edit distance:

- Number of changes needed for  $S1 \rightarrow S2$
- Uniform scoring function

# Formulation 3: Sequence alignment

- Allow gaps (fixed penalty)
  - Insertion & deletion operations
  - Unit cost for each character inserted or deleted
- Varying penalties for edit operations
  - Transitions (Pyrimidine $\leftrightarrow$ Pyrimidine, Purine $\leftrightarrow$ Purine)
  - Transversions (Purine  $\leftrightarrow$  Pyrimidine changes)
  - Polymerase confuses Aw/G and Cw/T more often

Scoring function:

$$\text{Match}(x,x) = +1$$

$$\text{Mismatch}(A,G) = -\frac{1}{2}$$

$$\text{Mismatch}(C,T) = -\frac{1}{2}$$

$$\text{Mismatch}(x,y) = -1$$

	A	G	T	C
A	+1	$-\frac{1}{2}$	-1	-1
G	$-\frac{1}{2}$	+1	-1	-1
T	-1	-1	+1	$-\frac{1}{2}$
C	-1	-1	$-\frac{1}{2}$	+1

purine      pyrimid.

**Transitions:**

$A \leftrightarrow G$ ,  $C \leftrightarrow T$  common  
(lower penalty)

**Transversions:**

All other operations

---

**Etc...**  
**(e.g. varying gap penalties)**

# How can we compute best alignment

---

S1	A	C	G	T	C	A	T	C	A
S2	T	A	G	T	G	T	C	A	

- Given additive scoring function:
  - Cost of mutation (AG, CT, other)
  - Cost of insertion / deletion
  - Reward of match
- Need algorithm for inferring best alignment
  - Enumeration?
  - How would you do it?
  - How many alignments are there?

# Can we simply enumerate all possible alignments?

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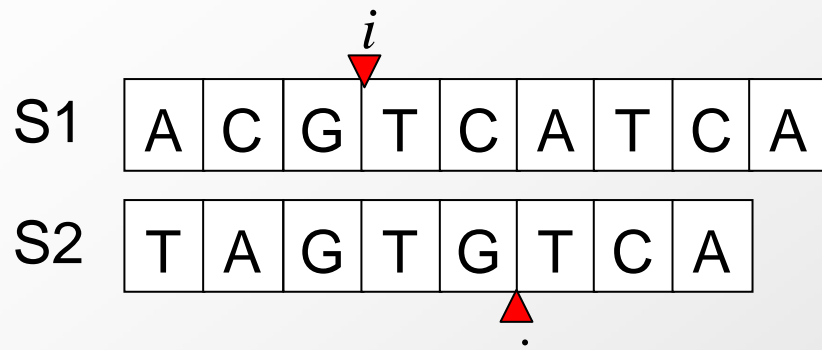
- Ways to align two sequences of length m, n

$$\binom{n+m}{m} = \frac{(m+n)!}{(m!)^2} \approx \frac{2^{m+n}}{\sqrt{\pi \cdot m}}$$

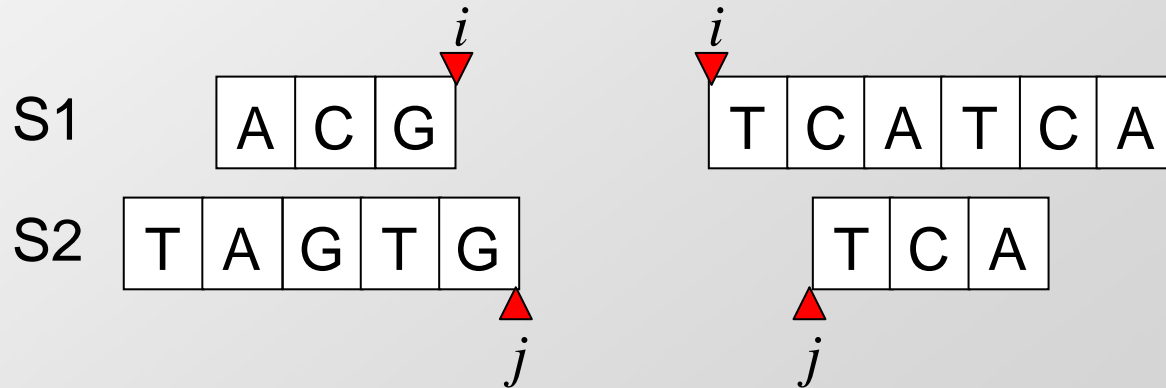
- For two sequences of length n

n	Enumeration	Today's lecture
10	184,756	100
20	1.40E+11	400
100	9.00E+58	10,000

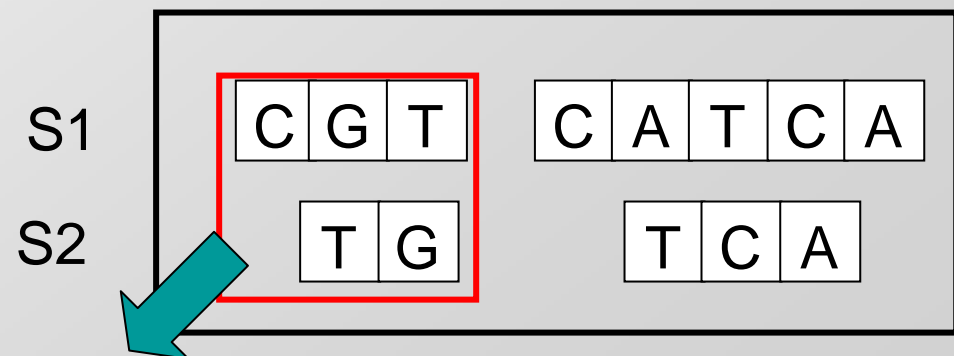
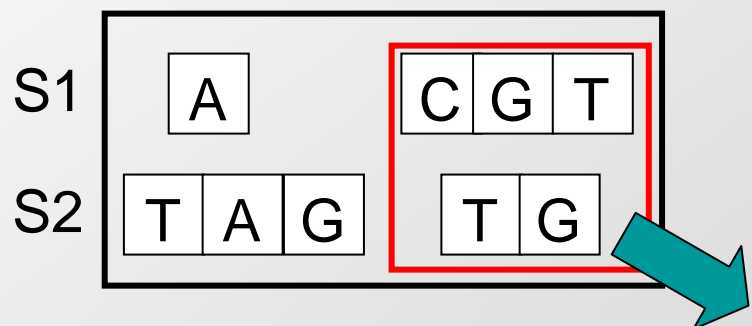
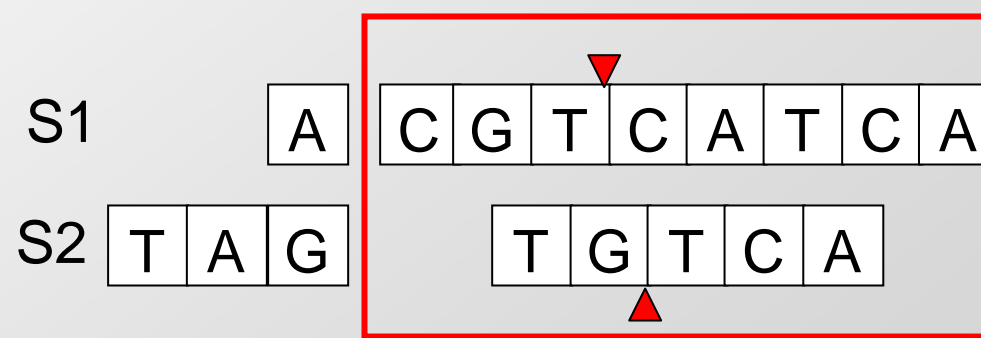
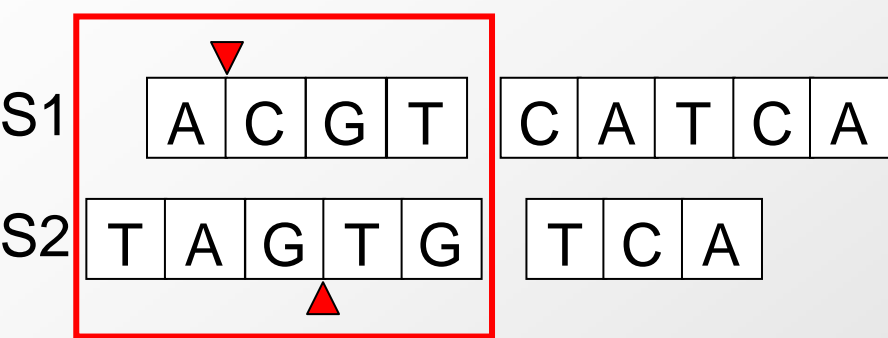
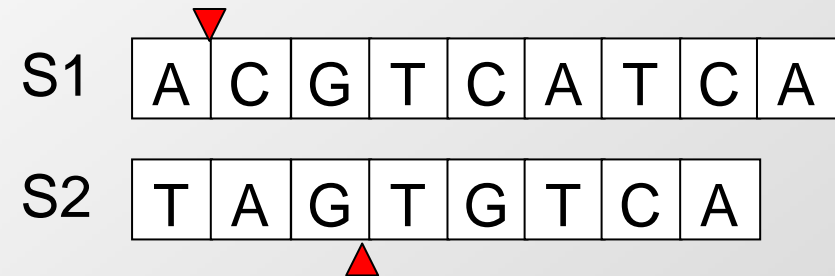
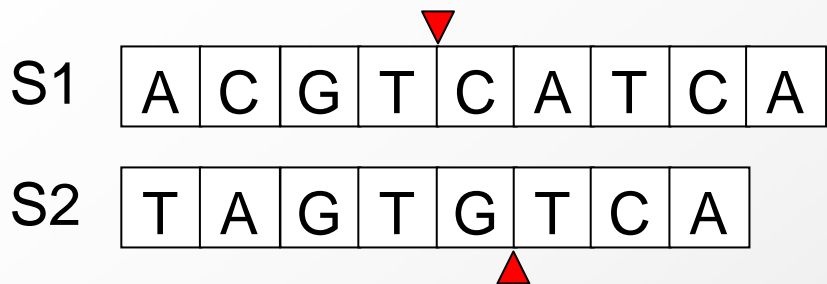
# Key insight: score is additive!



- Compute best alignment recursively
  - For a given aligned pair  $(i, j)$ , the best alignment is:
    - Best alignment of  $S1[1..i]$  and  $S2[1..j]$
    - + Best alignment of  $S1[i..n]$  and  $S2[j..m]$
  - Proof: cut-and-paste argument (see 6.046)



# Key insight: re-use computation



Identical sub-problems! We can reuse our work!

# Solution #1 – Memoization

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- Create a big dictionary, indexed by aligned seqs
  - When you encounter a new pair of sequences
  - If it is in the dictionary:
    - Look up the solution
  - If it is not in the dictionary
    - Compute the solution
    - Insert the solution in the dictionary
- Ensures that there is no duplicated work
  - Only need to compute each sub-alignment once!

**Top down approach**

## Solution #2 – Dynamic programming

---

- Create a big table, indexed by  $(i,j)$ 
  - Fill it in from the beginning all the way till the end
  - You know that you'll need every subpart
  - Guaranteed to explore entire search space
- Ensures that there is no duplicated work
  - Only need to compute each sub-alignment once!
- Very simple computationally!

**Bottom up approach**

# A simple introduction to Dynamic Programming

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- Fibonacci numbers

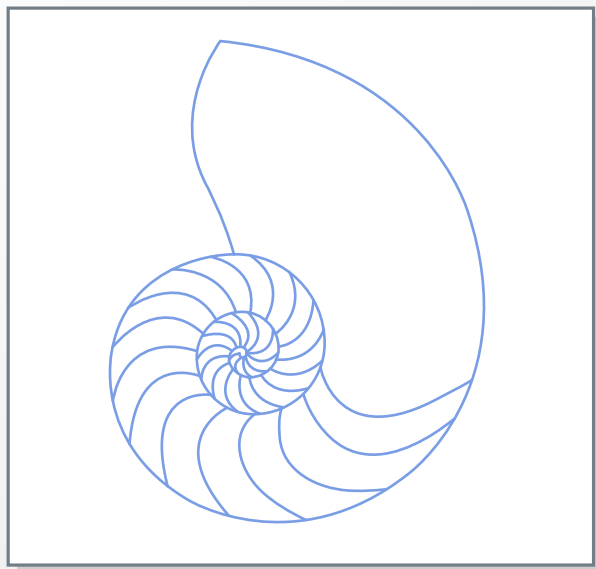
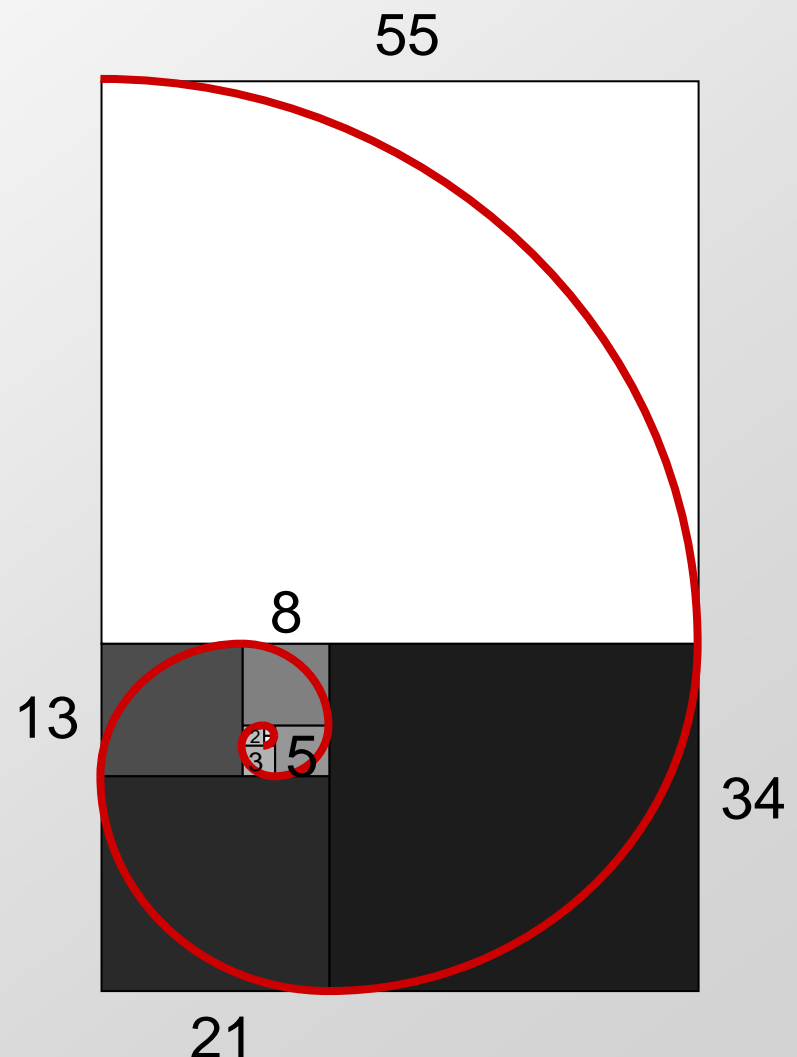
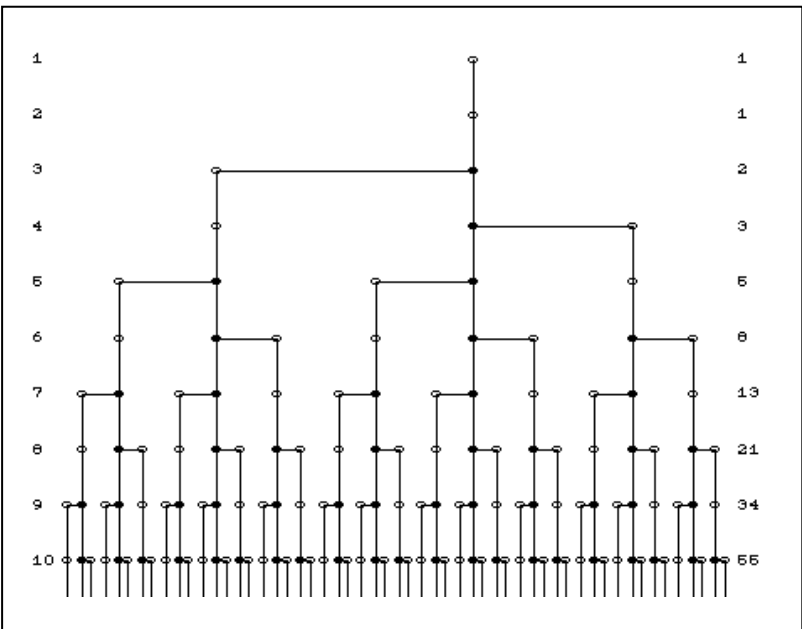


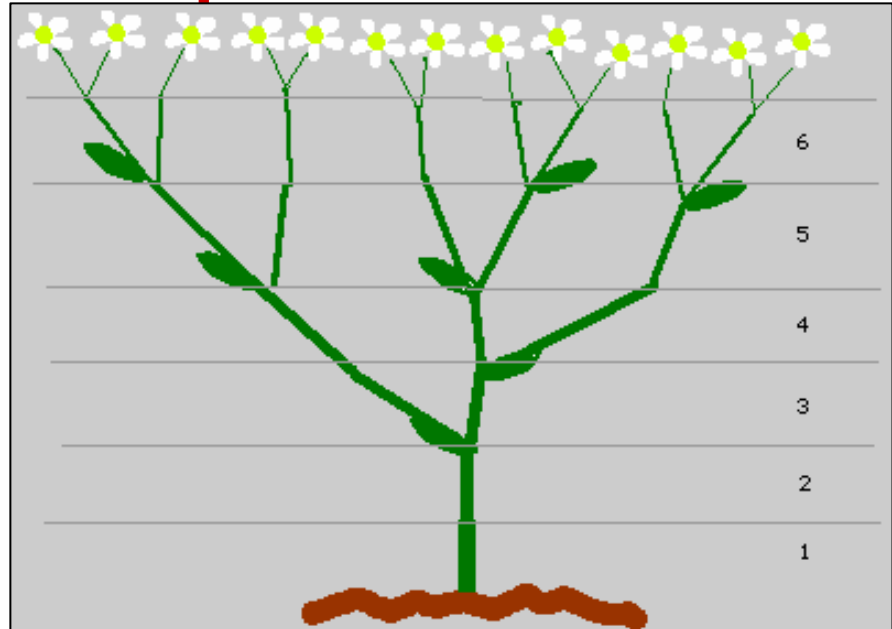
Figure by MIT OpenCourseWare.



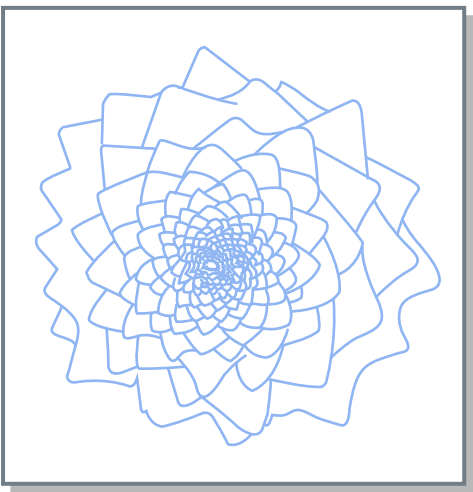
# Fibonacci numbers are ubiquitous in nature



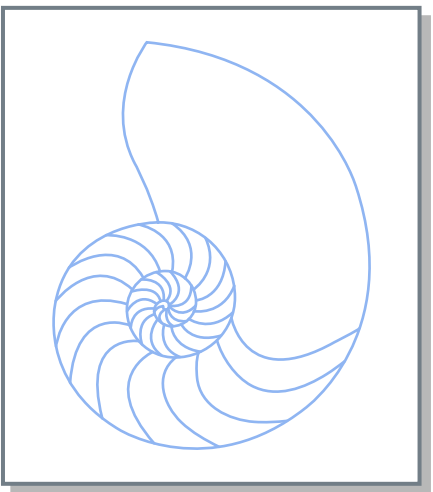
## Rabbits per generation



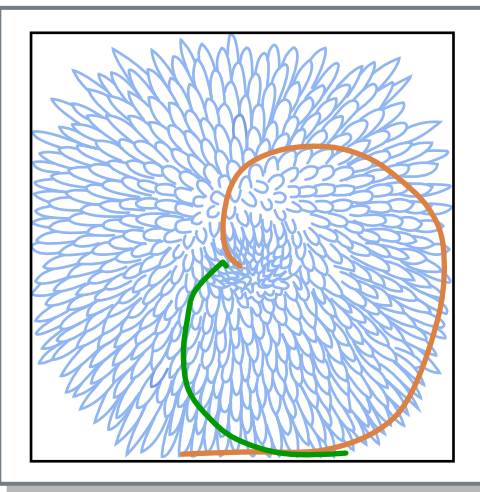
## Leaves per height



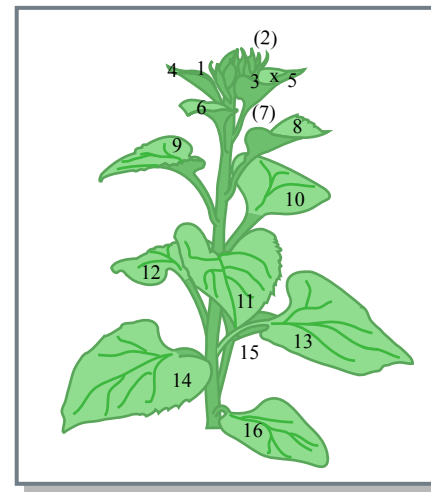
## Romanesque spirals



## Nautilus size



# Coneflower spirals



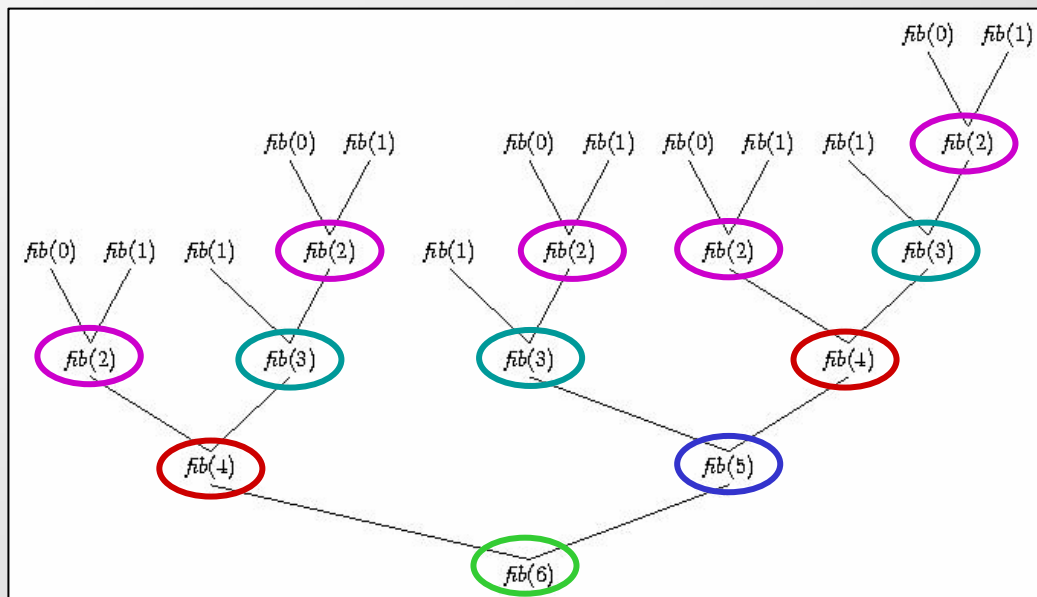
## Leaf ordering

# Computing Fibonacci numbers: Top down

- Fibonacci numbers are defined recursively:
  - Python code

```
def fibonacci(n):  
    if n==1 or n==2: return 1  
    return fibonacci(n-1) + fibonacci(n-2)
```

- Goal: Compute  $n^{\text{th}}$  Fibonacci number.
  - $F(0)=1, F(1)=1, F(n)=F(n-1)+F(n-2)$
  - 1,1,2,3,5,8,13,21,34,55,89,144,233,377,...
- Analysis:
  - $T(n) = T(n-1) + T(n-2) = (\dots) = O(2^n)$



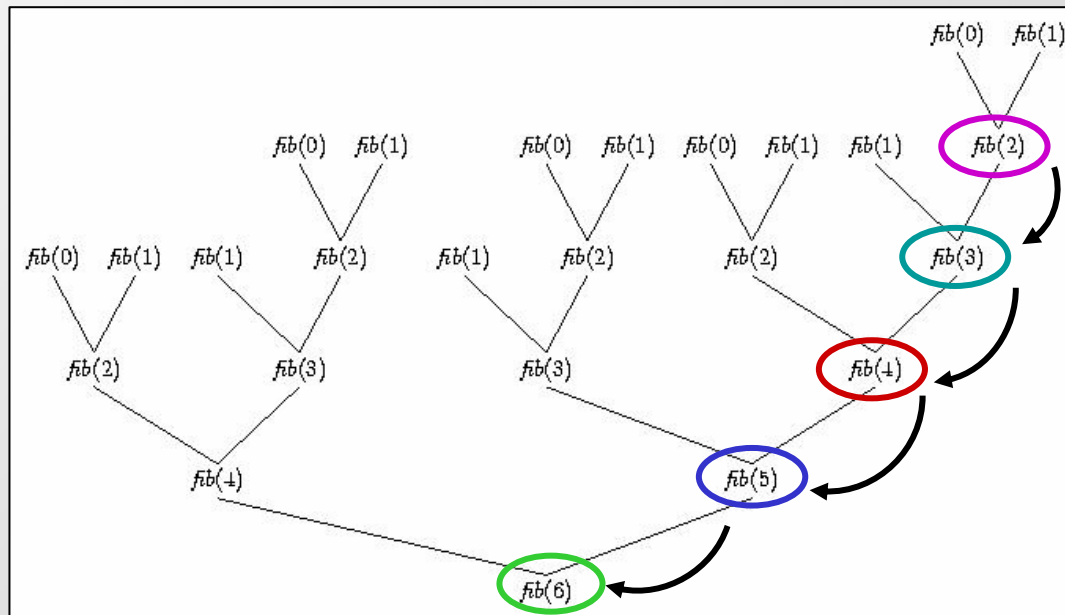
# Computing Fibonacci numbers: Bottom up

- Top-down approach
  - Python code

fib_table	
F[1]	1
F[2]	1
F[3]	2
F[4]	3
F[5]	5
F[6]	8
F[7]	13
F[8]	21
F[9]	34
F[10]	55
F[11]	89
F[12]	?

```
def fibonacci(n):
    fib_table[1] = 1
    fib_table[2] = 1
    for i in range(3,n+1):
        fib_table[i] = fib_table[i-1]+fib_table[i-2]
    return fib_table[n]
```

- Analysis:  $T(n) = O(n)$



# Lessons from iterative Fibonacci algorithm

---

fib_table	
F[1]	1
F[2]	1
F[3]	2
F[4]	3
F[5]	5
F[6]	8
F[7]	13
F[8]	21
F[9]	34
F[10]	55
F[11]	89
F[12]	?

- What did the iterative solution do?
  - Reveal identical sub-problems
  - Order computation to enable result reuse
  - Systematically filled-in table of results
  - Expressed larger problems from their subparts
- Ordering of computations matters
  - Naïve top-down approach very slow
    - results of smaller problems not available
    - repeated work
  - Systematic bottom-up approach successful
    - Systematically solve each sub-problem
    - Fill-in table of sub-problem results in order.
    - Look up solutions instead of recomputing

# Dynamic Programming in Theory

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- Hallmarks of Dynamic Programming
  - **Optimal substructure:** Optimal solution to problem (instance) contains optimal solutions to sub-problems
  - **Overlapping subproblems:** Limited number of distinct subproblems, repeated many many times
- Typically for optimization problems (unlike Fib example)
  - Optimal choice made locally:  $\max(\text{subsolution score})$
  - Score is typically added through the search space
  - Traceback common, find optimal path from indiv. choices
- Middle of the road in range of difficulty
  - Easier: greedy choice possible at each step
  - DynProg: requires a traceback to find that optimal path
  - Harder: no opt. substr., e.g. subproblem dependencies

# Hallmarks of optimization problems

## Greedy algorithms

## Dynamic Programming

### 1. Optimal substructure

*An optimal solution to a problem (instance) contains optimal solutions to subproblems.*

### 2. Overlapping subproblems

*A recursive solution contains a “small” number of distinct subproblems repeated many times.*

### 3. Greedy choice property

*Locally optimal choices lead to globally optimal solution*

**Greedy Choice is not possible**  
*Globally optimal solution requires trace back through many choices*

# Dynamic Programming in Practice

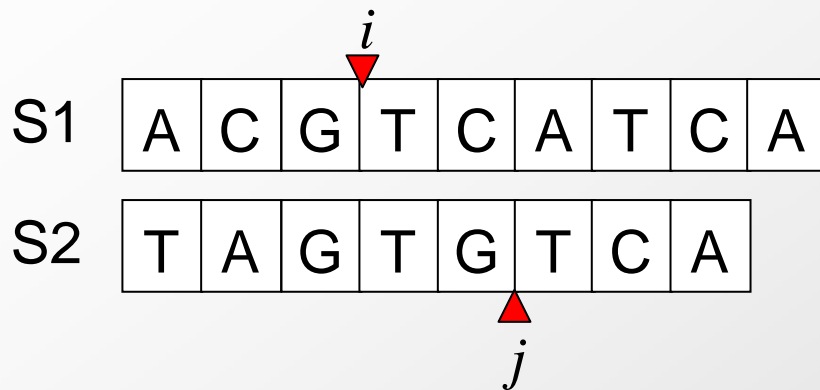
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- Setting up dynamic programming
  1. Find ‘matrix’ parameterization (# dimensions, variables)
  2. Make sure sub-problem space is finite! (not exponential)
    - If not all subproblems are used, better off using memoization
    - If reuse not extensive, perhaps DynProg is not right solution!
  3. Traversal order: sub-results ready when you need them
    - Computation order matters! (bottom-up, but not always obvious)
  4. Recursion formula: larger problems = F(subparts)
  5. Remember choices: typically F() includes min() or max()
    - Need representation for storing pointers, is this polynomial !
- Then start computing
  1. Systematically fill in table of results, find optimal score
  2. Trace-back from optimal score, find optimal solution

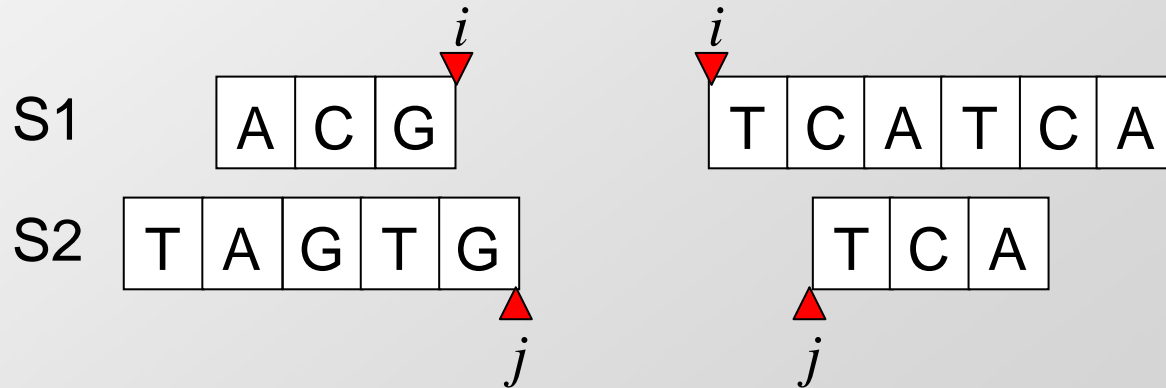
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**How do we apply dynamic programming  
to sequence alignment ?**

# Key insight: score is additive!



- Compute best alignment recursively
  - For a given aligned pair  $(i, j)$ , the best alignment is:
    - Best alignment of  $S1[1..i]$  and  $S2[1..j]$
    - + Best alignment of  $S1[i..n]$  and  $S2[j..m]$



# Dynamic Programming for sequence alignment

---

- Setting up dynamic programming
  1. Find ‘matrix’ parameterization
  2. Make sure sub-problem space is finite! (not exponential)
  3. Traversal order: sub-results ready when you need them
  4. Recursion formula: larger problems =  $F(\text{subparts})$
  5. Remember choices: typically  $F()$  includes  $\min()$  or  $\max()$
- Then start computing
  1. Systematically fill in table of results, find optimal score
  2. Trace-back from optimal score, find optimal solution

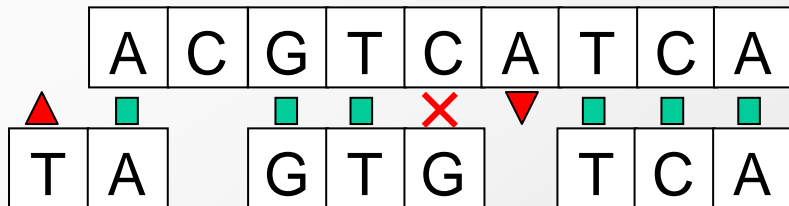
**(1, 2, 3) Store score of aligning (i,j) in matrix M(i,j)**

The diagram illustrates a dynamic programming matrix used for sequence alignment. The matrix is divided into four quadrants by labels at the top and left:

- S[1..i]**: The first  $i$  columns of the matrix, colored light blue.
- i**: The label for the  $i$ -th column, positioned above the boundary between the light blue and grey areas.
- S[i..n]**: The remaining columns of the matrix, colored grey.
- T[1..j]**: The first  $j$  rows of the matrix, colored light blue.
- j**: The label for the  $j$ -th row, positioned to the left of the boundary between the light blue and grey areas.
- T[j..m]**: The remaining rows of the matrix, colored grey.

A red arrow points from a cell in the  $S[1..i]$  column to a cell in the  $S[i..n]$  column, indicating a path through the matrix. A callout box at the bottom right contains the text: "Best alignment  $\Leftrightarrow$  Best path through the matrix".

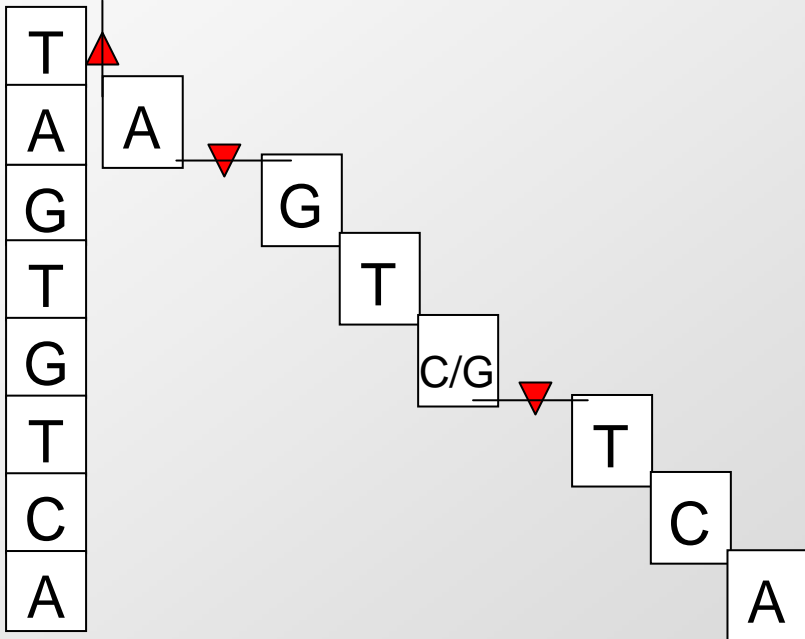
# Duality: seq. alignment $\Leftrightarrow$ path through the matrix



S1

A C G T C A T C A

S2



**Goal:**  
Find best path  
through the matrix

## (4) Filling in the dynamic programming matrix

---

- Local update rules:
  - Compute next alignment based on previous alignment
  - Just like Fibonacci numbers:  $F[i] = F[i-1] + F[i-2]$
  - Table lookup!
- Compute scores for prefixes of increasing length
  - This allows a single recursion (top-left to bottom-right) instead of two recursions (middle-to-outside top-down)
  - Only three possibilities for extending by one nucleotide: a gap in one species, a gap in the other, a (mis)match
  - When you reach bottom right, prefix of length  $n$  is seq S
- Computing the score of a cell from its neighbors
  - $F(i-1, j) - \text{gap}$
  - $- F(i, j) = \max\{ F(i, j) + \text{score}, F(i, j-1) - \text{gap} \}$

## 0. Setting up the scoring matrix

-	A	G	T
-	0		
A			
A			
G			
C			

**Initialization:**

- Top left: 0

**Update Rule:**

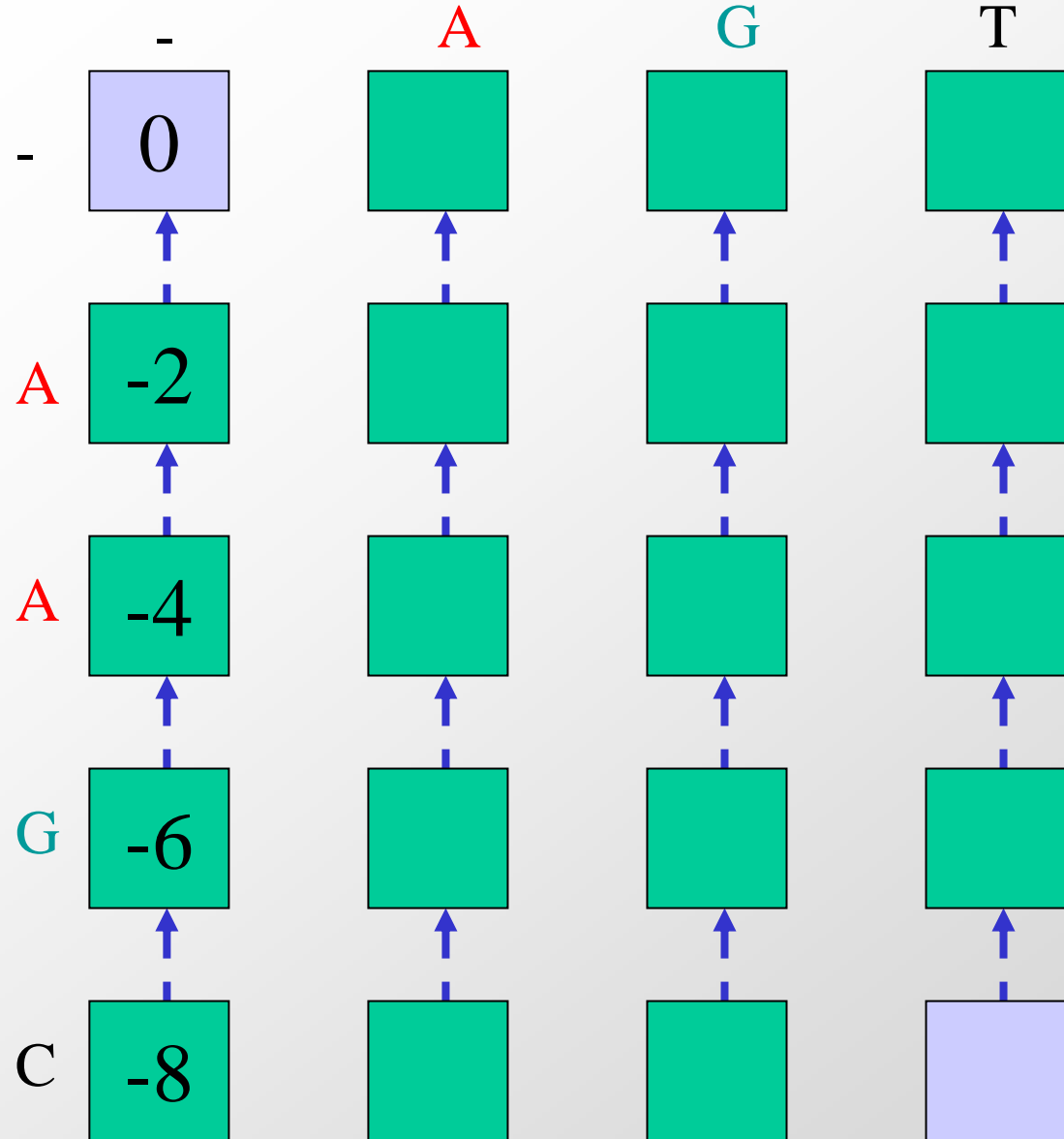
$$A(i,j) = \max\{$$

}

**Termination:**

- Bottom right

# 1. Allowing gaps in s



**Initialization:**

- Top left: 0

**Update Rule:**

$$A(i,j) = \max\{$$

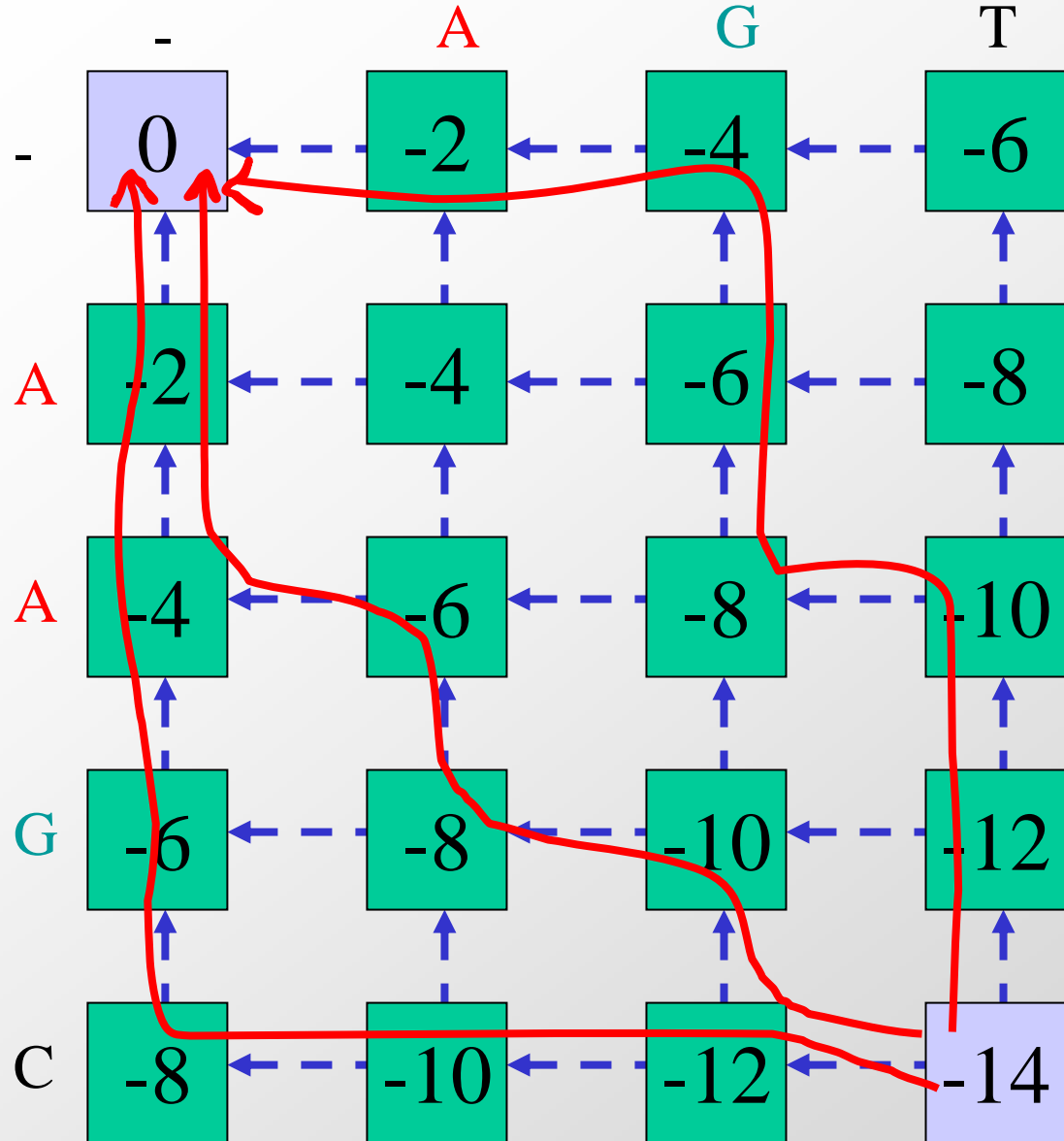
- $A(i-1, j) - 2$

}

**Termination:**

- Bottom right

## 2. Allowing gaps in t



**Initialization:**

- Top left: 0

**Update Rule:**

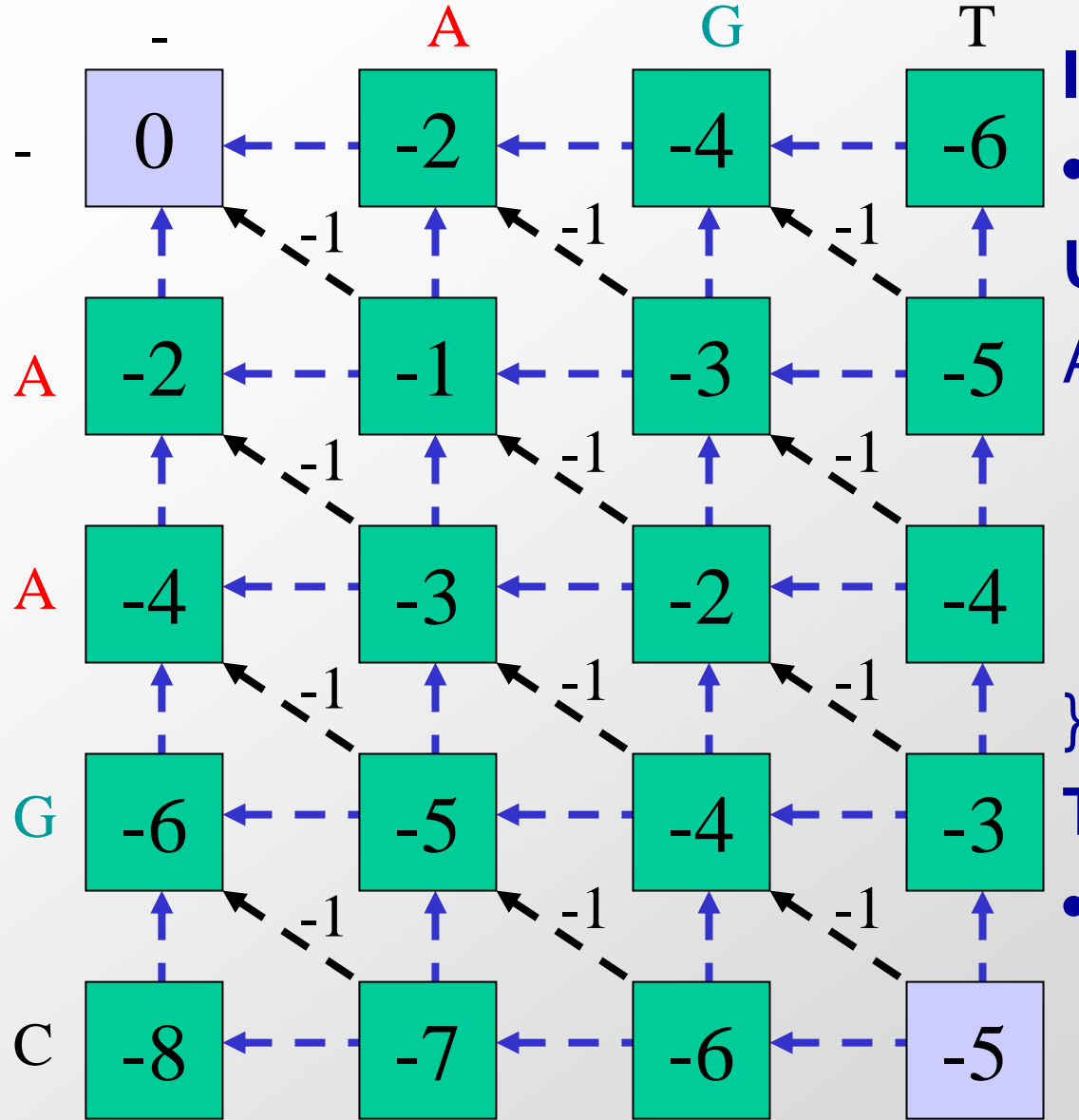
$$A(i,j) = \max \{$$

- $A(i-1, j) - 2$
- $A(i, j-1) - 2$

**Termination:**

- Bottom right

### 3. Allowing mismatches



**Initialization:**

- Top left: 0

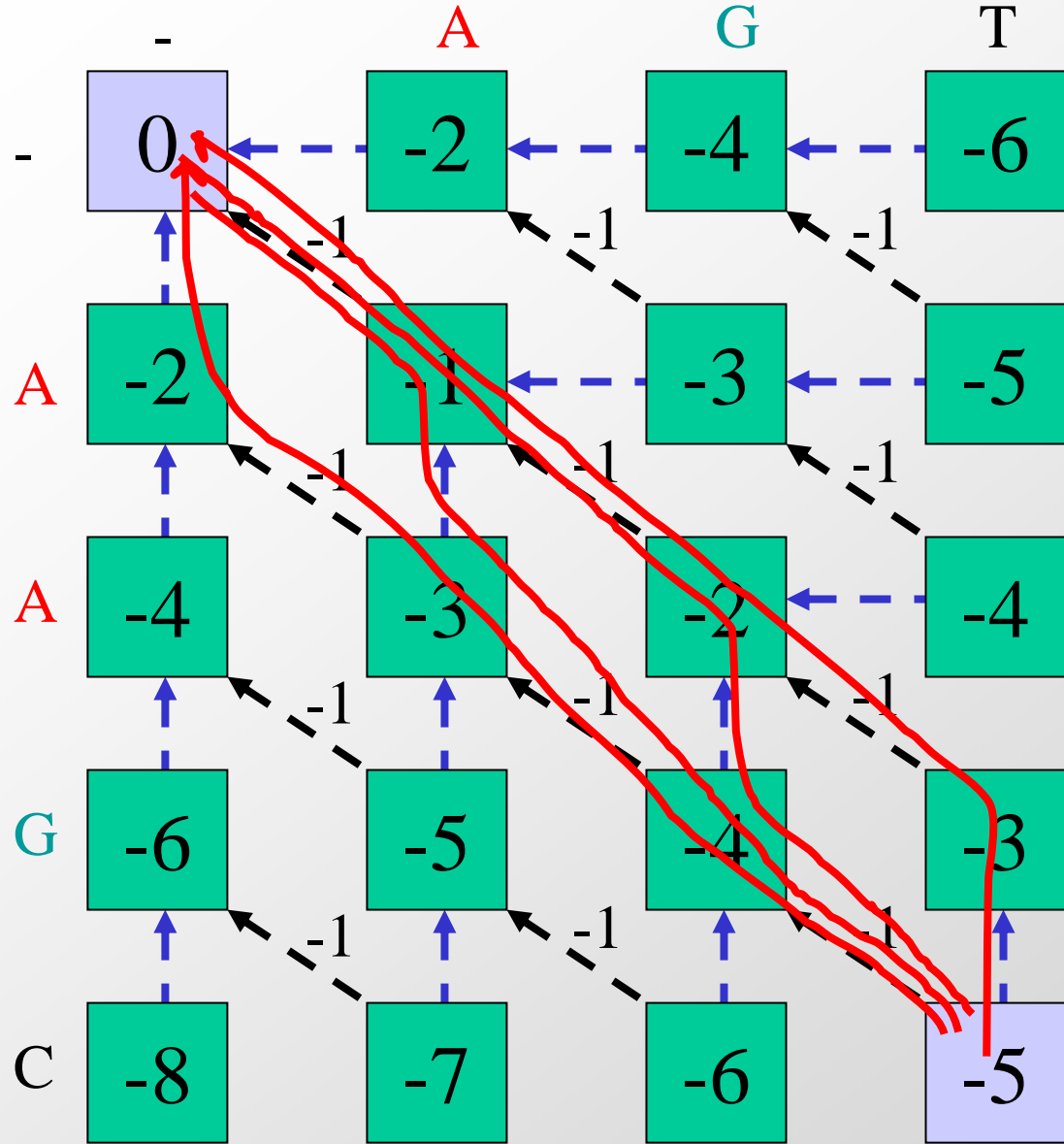
**Update Rule:**

$$A(i,j) = \max \{ \begin{array}{l} \bullet A(i-1, j) - 2 \\ \bullet A(i, j-1) - 2 \\ \bullet A(i-1, j-1) - 1 \end{array} \}$$

**Termination:**

- Bottom right

## 4. Choosing optimal paths



**Initialization:**

- Top left: 0

**Update Rule:**

$$A(i,j) = \max \{$$

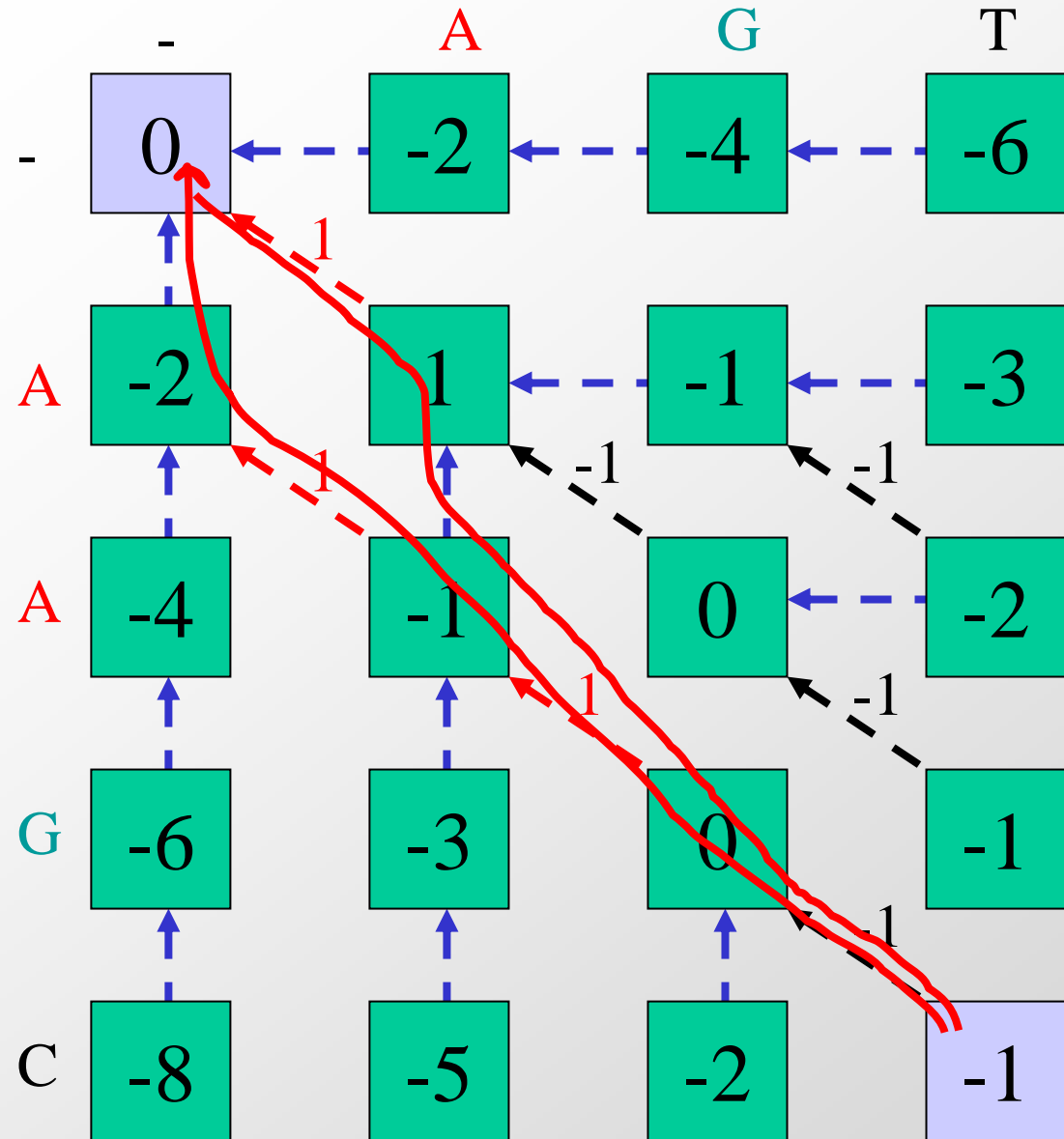
- $A(i-1, j) - 2$
- $A(i, j-1) - 2$
- $A(i-1, j-1) - 1$

$$\}$$

**Termination:**

- Bottom right

## 5. Rewarding matches



**Initialization:**

- Top left: 0

**Update Rule:**

$$A(i,j) = \max \{$$

- $A(i-1, j) - 2$
- $A(i, j-1) - 2$
- $A(i-1, j-1) \pm 1$

$$\}$$

**Termination:**

- Bottom right

# What is missing? (5) Returning the actual path!

- We know how to compute the best score
  - Simply the number at the bottom right entry
- But we need to remember where it came from
  - Pointer to the choice we made at each step
- Retrace path through the matrix
  - Need to remember all the pointers



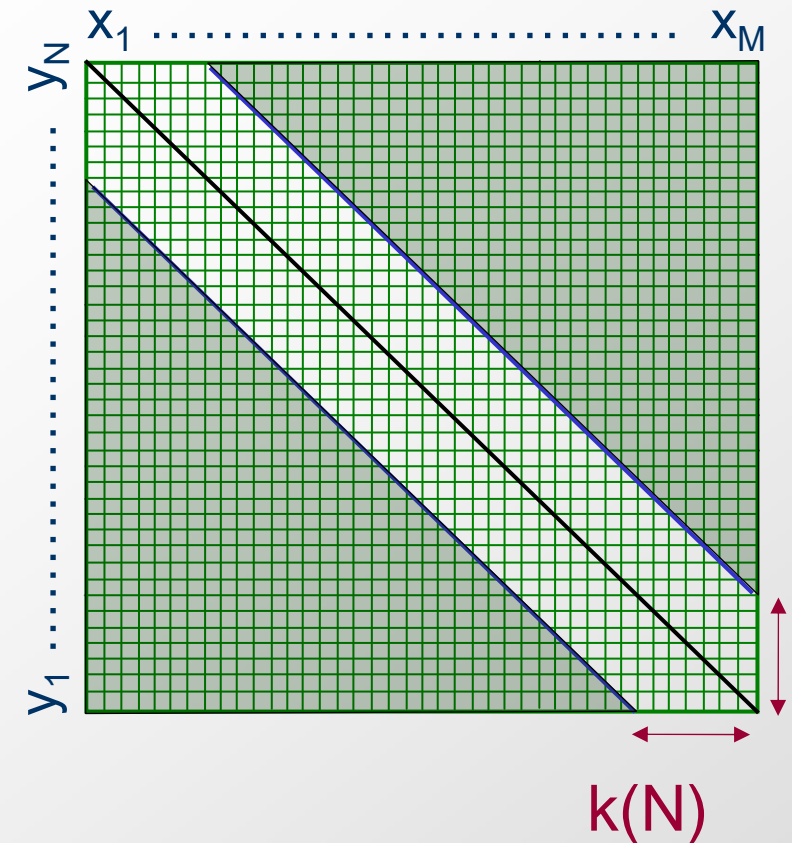
Time needed:  $O(m * n)$   
Space needed:  $O(m * n)$

# Summary

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- Dynamic programming
  - Reuse of computation
  - Order sub-problems. Fill table of sub-problem results
  - Read table instead of repeating work (ex: Fibonacci)
- Sequence alignment
  - Edit distance and scoring functions
  - Dynamic programming matrix
  - Matrix traversal path  $\Leftrightarrow$  Optimal alignment
- Thursday: Variations on sequence alignment
  - Local/global alignment, affine gaps, algo speed-ups
  - Semi-numerical alignment, hashing, database lookup
- Recitation:
  - Dynamic programming applications
  - Probabilistic derivations of alignment scores

# Bounded Dynamic Programming



## Initialization:

$F(i, 0), F(0, j)$  undefined for  $i, j > k$

## Iteration:

For  $i = 1 \dots M$

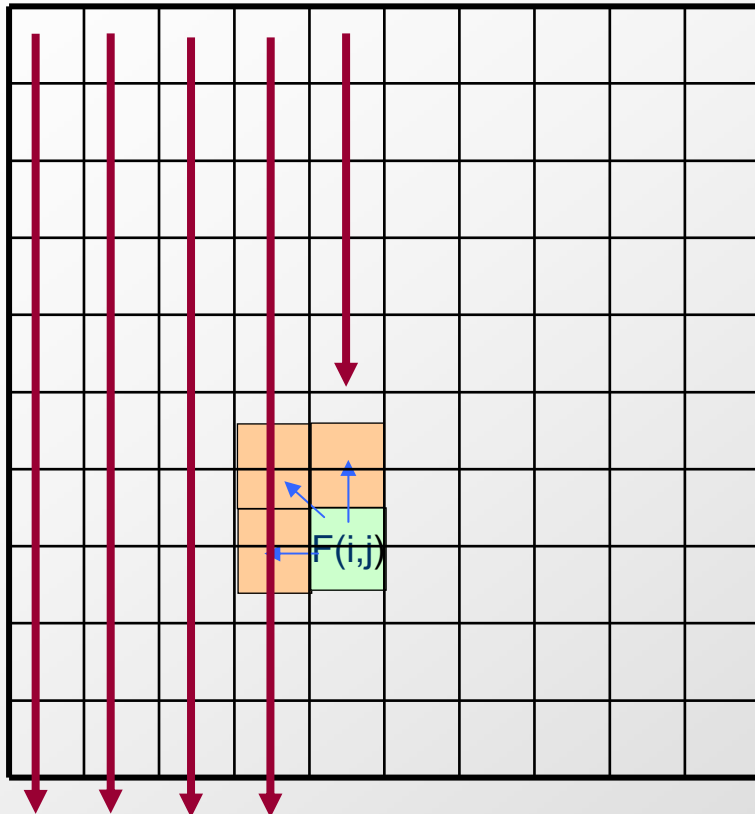
For  $j = \max(1, i - k) \dots \min(N, i + k)$

$$F(i, j) = \max \begin{cases} F(i - 1, j - 1) + s(x_i, y_j) \\ F(i, j - 1) - d, \text{ if } j > i - k(N) \\ F(i - 1, j) - d, \text{ if } j < i + k(N) \end{cases}$$

## Termination: same

# Linear space alignment

It is easy to compute  $F(M, N)$  in linear space



Allocate ( column[1] )

Allocate ( column[2] )

For  $i = 1 \dots M$

If  $i > 1$ , then:

Free( column[i - 2] )

Allocate( column[ i ] )

For  $j = 1 \dots N$

$F(i, j) = \dots$

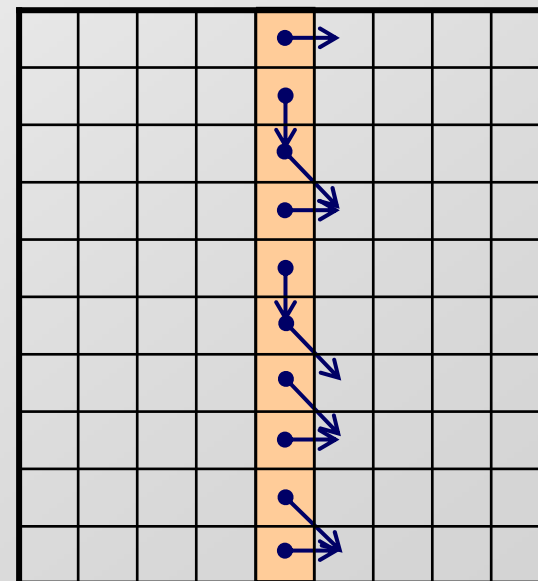
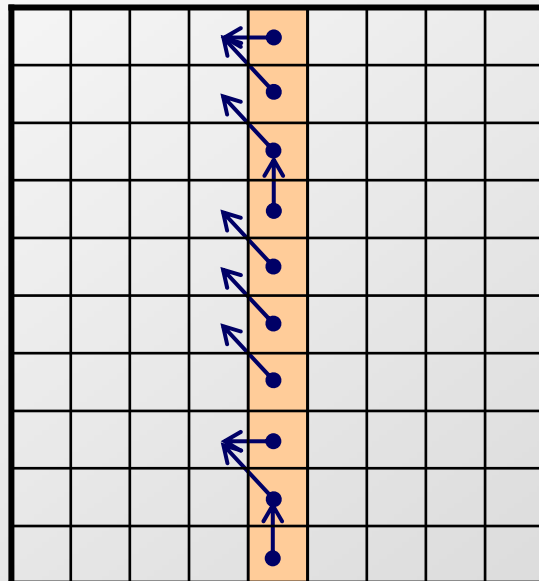
What about the pointers?

# Finding the best back-pointer for current column

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- Now, using 2 columns of space, we can compute for  $k = 1 \dots M$ ,  $F(M/2, k)$ ,  $F^r(M/2, N-k)$

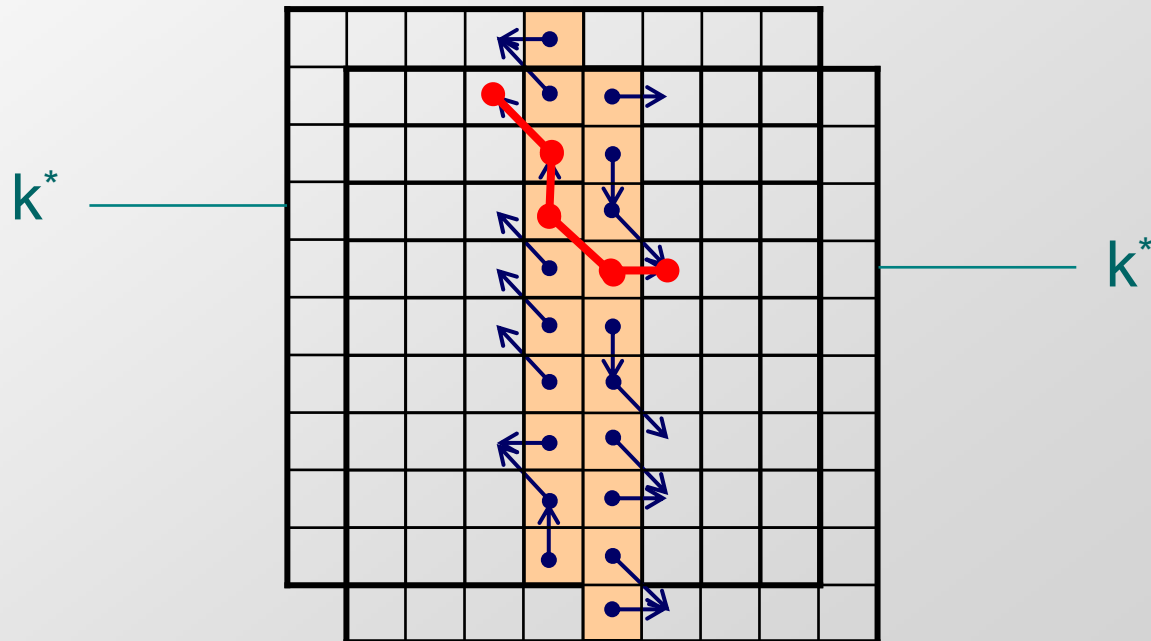
PLUS the backpointers



# Best forward-pointer for current column

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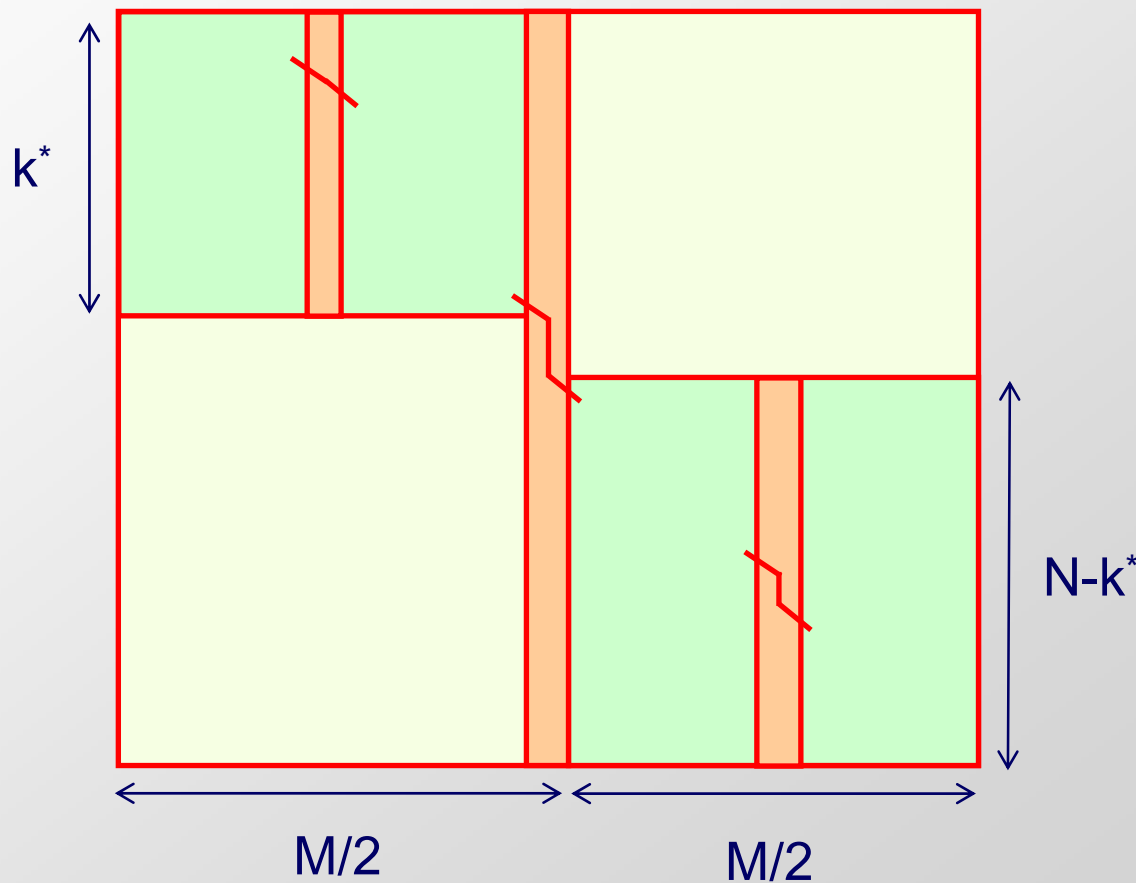
- Now, we can find  $k^*$  maximizing  $F(M/2, k) + F^r(M/2, N-k)$
- Also, we can trace the path exiting column  $M/2$  from  $k^*$



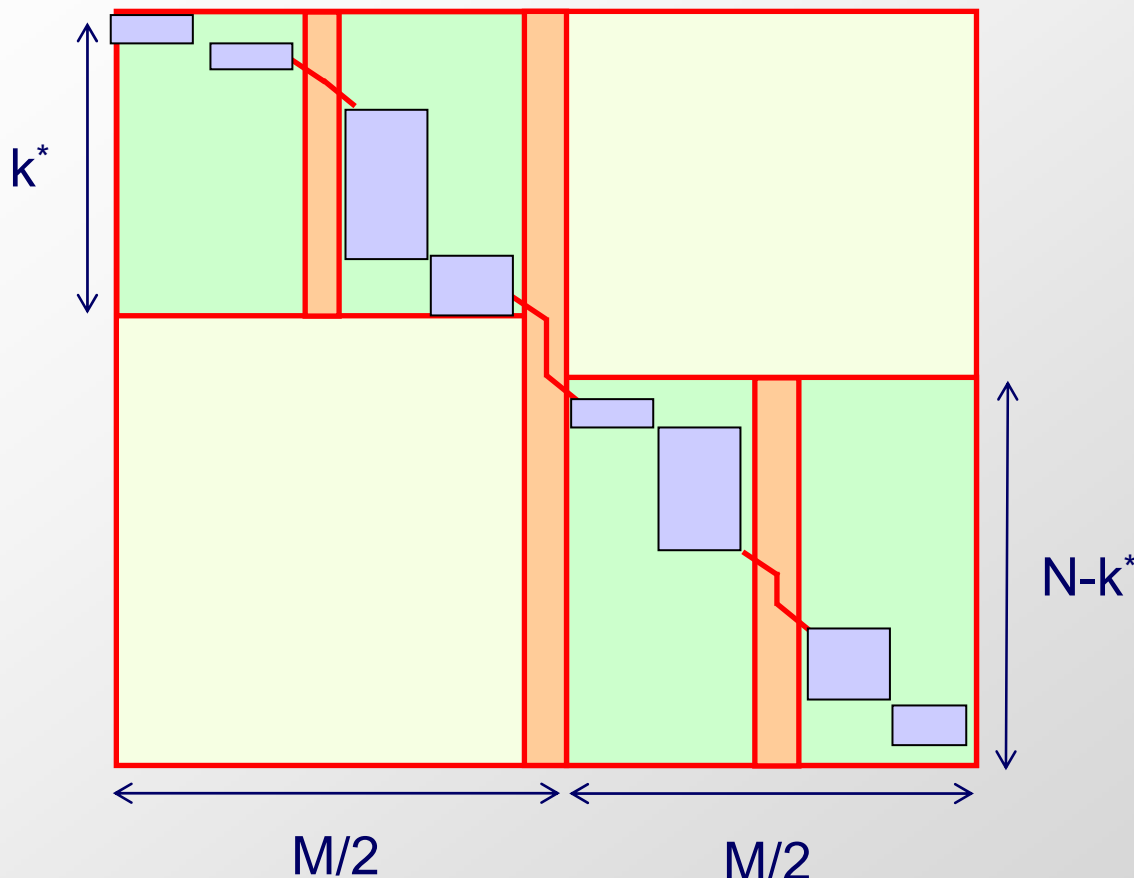
# Recursively find midpoint for left & right

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- Iterate this procedure to the left and right!



# Total time cost of linear-space alignment



**Total Time:**  $cMN + cMN/2 + cMN/4 + \dots = 2cMN = O(MN)$

**Total Space:**  $O(N)$  for computation,  
 $O(N+M)$  to store the optimal alignment

## Formulation 4: Varying gap cost models (next time)

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(still) Varying penalties for edit operations

Now allow gaps of varying penalty:

1. Linear gap penalty
  - Same as before,
2. Affine gap penalty
  - Big initial cost for starting or ending a gap
  - Small incremental cost for each additional character
3. General gap penalty
  - Any cost function
  - No longer computable using the same model
4. Seek duplicated regions, rearrangements, ...