# Lecture Notes for INF281 Basics of Bioinformatics Sequence Analysis

Takaya Saito



This work is licensed under a Creative Commons Attribution 4.0 International License.

# Contents

Ι			1
1	Intr 1.1 1.2 1.3	Introduction to Molecular Biology	5
II			8
2	Glo	bal pairwise alignment	8
	2.1	Pairwise alignment	8
	2.2	Alignment by brute–force	9
	2.3	Table representation of alignment	10
	2.4	Global alignment with DP	12
	2.5	Backtracking	14
	2.6	Needleman-Wunsch algorithm	17

# Part I

# 1 Introduction

# 1.1 Introduction to Molecular Biology

Molecular biology is the study of biology focusing on organisms and cells at the molecular level.

#### Five essential facts about cells

## 1. Two primary types of cells - eukaryotes and prokaryotes

• Eukaryote: animals & plants

• Prokaryote: bacteria & archaea

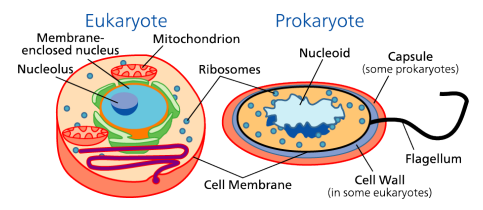


Figure 1.1: Eukaryotic and prokaryotic cells (source: Science Primer, Wikimedia Commons)

## 2. Cell size - around 1 to 100 micrometers

• Cell Size and Scale: http://learn.genetics.utah.edu/content/cells/scale

## 3. The number of cells

• Prokaryotes: 1 cell

• Human: Estimate of 15 trillion cells

## 4. An animal cell and cell organelles



Figure 1.2: An animal cell and organelles (source: Kelvinsong, Wikimedia Commons)

## 5. Cellular processes

- Cell growth, cell development, cell signaling,
- Example: http://www.nature.com/nrg/multimedia/rnai

## Central dogma of molecular biology

It describes the information flow within a cell.

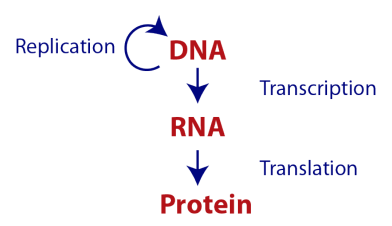


Figure 1.3: Central dogma of molecular biology

## DNA (deoxyribonucleic acid)

DNA stores genetic information. It has four different bases: cytosine (C), guanine (G), adenine (A), and thymine (T).

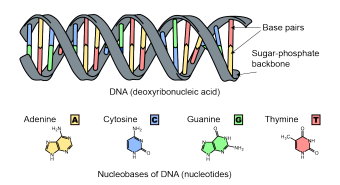


Figure 1.4: DNA double helix and base pairs (modified from the original version by Sponk, Wikimedia Commons)

# Base pair matching (Watson-Crick base pair)

Adenine (A) pairs with thymine (T), whereas cytosine (C) pairs with guanine (G).

DNA strand1: ACGT

 $\square$ 

DNA strand2: TGCA

## RNA (Ribonucleic acid)

RNA has various biological roles and several sub-classes. Messenger RNAs (mRNAs) convey genetic information. It has four different bases: cytosine (C), guanine (G), adenine (A), and uracil (U).

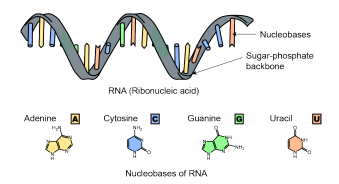


Figure 1.5: Single strand RNA (modified from the original version by Sponk, Wikimedia Commons)

# Transcription: mRNAs are transcribed from DNAs

DNA: ACGT -----> RNA: ACGU Transcription

#### Protein

Proteins are large molecules consisting of amino acids. There are 20 common amino acids.

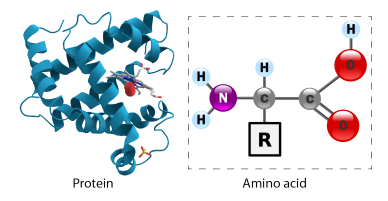


Figure 1.6: Protein 3D structure and amino acids (sources: AzaToth, Wikimedia Commons, YassineMrabet, Wikimedia Commons)

Translation: Amino-acids are translated from mRNAs

mRNA: GUC -----> AA: Valine Translation

## Universal genetic code

A codon consists of three nucleic acids. Single-letter or three-letter names can be used for amino acids.

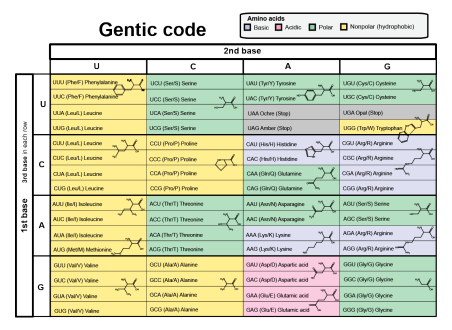


Figure 1.7: Universal genetic code (modified from the original version by Häggström, Wikimedia Commons)

# Cellular functions of proteins

- Enzymes: catalyze chemical reaction
- Cell signaling: hormone (e.g. insulin), antibodies,
- Structural: collagen, cartilage, keratin,

## Exercises 1.1

- 1. Draw a simple diagram of the central dogma of molecular biology and briefly explain the information flow of the molecules.
- 2. What are the DNA sequences of the opposite strand for the following DNA sequences?

Seq1 CCGATT Seq2 TTACGC

Seq3 ACGCGC

- 3. What are the mRNA sequences transcribed from the following DNA sequences?
- 4. What are the polypeptide sequences translated from the following mRNA sequences? Answer them with both one-letter and three letter names.

Seq1 AUGUUUUAA Seq2 GCAGCAAAA

# 1.2 Introduction to Biotechnology

Biotechnology is the use of laboratory techniques to study living organism and cells.

# Applications of biotechnology

Branches of biotechnology can be explained with different colors.

• Red: medical processes

• Green: agricultural processes

• White: industrial processes

• Blue: marine and aquatic applications

## Laboratory tools and equipment



Figure 1.8: Pipette, centrifuge, thermal cycler, and DNA sequencer (sources: Domain, Manske, Rror, RE73 via Wikimedia Commons)

## Human genome project

It was a large-scale international research project to determine the whole DNA sequences of human.

- 1990 2003
- \$2.7 billion

## Next generation sequencing

Sequence technologies have been rapidly advanced since the human genome project. Example: sequence a whole human genome with Illumina HiSeq X Ten.

- One day
- \$1000

## Protein sequencing

Proteins are generally more studied than DNAs and RNAs, but the whole proteome is generally harder to analyze than the whole genome. MS (mass-spectrometry) based technologies are widely used to sequence proteins.



Figure 1.9: Orbitrap mass spectrometer (source: Wiòrkiewicz, Wikimedia Commons)

## 1.3 Bioinformatics

Bioinformatics uses computational approaches to solve problems in life sciences. It is based on computer science.

## Similar or almost equivalent disciplines

- Biostatistics
- Biophysics
- Systems biology
- Computational biology

#### Not much related with bioinformatics

- Health informatics
- Forensic science

## Scope of INF281

We mainly cover the following fields of bioinformatics in this course.

- Pairwise alignment
- Database search
- Statistical evaluation
- Multiple alignment
- Phylogenetic tree
- Scoring scheme
- Sequence patterns

# Popular bioinformatics programs

BLAST and ClustalW are popular tools for sequence analysis.

• BLAST: a program for database search

URL: http://blast.ncbi.nlm.nih.gov

• ClustalW: a program for multiple alignments

URL: http://www.ch.embnet.org/software/ClustalW.html

Rank	Title	Times cited
1	Protein measurement with the folin phenol reagent	305148
2	Cleavage of structural proteins during the assembly of the head of bacterio-	213005
	phage T4	
3	A rapid and sensitive method for the quantitation of microgram quantities of	155530
	protein utilizing the principle of protein-dye binding	
4	DNA sequencing with chain-terminating inhibitors	65335
5	Single-step method of RNA isolation by acid guanidinium thiocyanate-phenol-	60397
	chloroform extraction	
6	Electrophoretic transfer of proteins from polyacrylamide gels to nitrocellulose	53349
	sheets: procedure and some applications	
7	Development of the Colle-Salvetti correlation-energy formula into a functional	46702
	of the electron density	
8	Density-functional thermochemistry. III. The role of exact exchange	46145
9	A simple method for the isolation and purification of total lipides from animal	45131
	tissues	
10	Clustal W: improving the sensitivity of progressive multiple sequence align-	40289
	ment through sequence weighting, position-specific gap penalties and weight	
	matrix choice	
11	Nonparametric estimation from incomplete observations	38600
12	Basic local alignment search tool	38380
13	A short history of SHELX	37978
14	Gapped BLAST and PSI-BLAST: A new generation of protein database	36410
	search programs	
15	A revised medium for rapid growth and bio assays with tobacco tissue cultures	36132

**Table 1.1:** The 15 most cited papers of all time (The top 100 papers, Van Noorden, Maher, and Nuzzo, *Nature*, 2014)

# Part II

# 2 Global pairwise alignment

# 2.1 Pairwise alignment

A pairwise alignment is a basic sequence structure that consists of two sequences. A global alignment stretches to the whole part of two sequences, whereas a local alignment usually contains only part of the sequences.

## Components of pairwise alignment

We name two sequences as database or d and query or q through this course. They may represent sequences from two different species or organisms.

Identical sequences.

q: ACGT

d: ACGT

One mismatch.

q: ACGT

d: ACGA

The '-' symbol represents a blank. A single or a set of multiple blanks further represents a gap, which is an indication of insertion or deletion in the course of evolution between two organisms.

q: ACGT

d: A-GT

**N.B.** A gap cannot be aligned with another gap.

## Example of a simple scoring scheme

• Match: 1

• Mismatch: 0

• Gap penalty: 1 (use -1 for the actual calculation)

We may use the following notation.

•  $R_{ab} = 1$  for a = b

•  $R_{ab} = 0$  for  $a \neq b$ 

• g = 1

## Exercise 2.1

Use the simple scoring scheme above and calculate the scores of the following two alignments.

Alignment 1 Alignment 2

 q: GCA-GCA
 q: GCA-GCA

 d: GA-TG-A
 d: GA-TG-A

# 2.2 Alignment by brute-force

A brute–force approach finds the alignment with the highest score by simply considering all possible alignments and calculates the score for each of them.

# An example of brute-force approach

We find the optimal alignment for the following sequences by using the scoring scheme below.

Sequences: Scoring scheme:

q: AG, d: ACG 
$$R_{ab} = 1 \text{ for a = b} \\ R_{ab} = 0 \text{ for a } \neq \text{ b} \\ \text{g} = 1$$

# 1. The length of alignment

- Maximum length: length(q) + length(d)
- Minimum length: max(length(q), length(d))

# 2. All possible alignments when length = 5

# 3. All possible alignments when length = 4

# 4. All possible alignments when length = 3

# 5. Alignment with the best score

ACG A-G Score: 1

## Search space size of the brute-force approach

The search space size is the number of all possible alignments. It is 25 (10 + 12 + 3) for the example above.

# Rapid growth of search space size

Example 1 Example 2

q: ACGACG, d: AGAG q: ACGACGACGACG, d: AGAGAGAG

Search space size: 1289 Search space size: 4,673,345

#### Exercise 2.2

Find the alignment with the best score for the sequences. Use the simple scoring scheme below.

Sequences: Scoring scheme:

q: A, d: AC 
$$R_{ab} = 1 \text{ for a } = b$$
 
$$R_{ab} = 0 \text{ for a } \neq b$$
 
$$g = 1$$

1. What are the maximum and minimum lengths of the alignment?

2. What is the search space size when the brute-force approach is used?

3. Identify all possible alignments.

4. What is the best score?

# 2.3 Table representation of alignment

Several data structures can be used to represent an alignment. The table representation is frequently used and also makes the process clear when we combine it with dynamic programming (DP) later.

#### Data structures and algorithms

It is important to consider the following aspects before solving computational problems.

- 1. Identify and analyze the problem you want to solve
- 2. Pick up an algorithm that can efficiently solve the problem
- 3. Decide a data structure that works with the algorithm of your choice

We use a table format (2D array) to solve global alignments by dynamic programming.

# Example of table format

Alignment:

q: -AGd: A-CG

# 1. Initial setup

1. Make a table with the size of (1 + length(q)) by (1 + length(d))

2. Add the database sequence as column labels

3. And the query sequence as row labels

q/d		Α	С	G
	S			
Α				
G				E

## 2. Add arrows

We use three types of arrows to form an alignment.

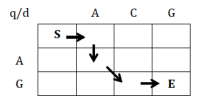
• Move diagonally: add the letters from q and d to the alignment

• Move vertically: add - and the letter from d to the alignment

• Move horizontally: add the letter from q and - to the alignment

Table 2

It should start from S and stops at E.



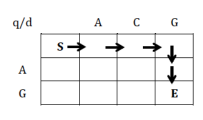
## Exercise 2.3

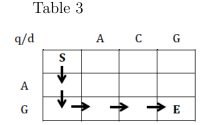
Find the corresponding alignments for Table 1, 2 and 3.

Table 1

q/d A C G

A G F





# 2.4 Global alignment with DP

Dynamic programming (DP) provides a solution for a multi-stage decision process, in which larger decisions recursively nest smaller decisions.

#### Memorize the best score in a table cell

For global alignment, the core procedure of DP is updating a cell with the highest score from the three different scores calculated from its adjacent cells. DP ends when the entire table is updated.



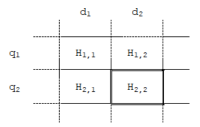
# Table notation and indices

 $H_{i,j}$  represents the score of the cell for the current update.  $H_{i-1,j}$ ,  $H_{i,j-1}$ , and  $H_{i-1,j-1}$  are the scores of the adjacent cells.

Cell  $H_{i,j}$  and its adjacent cells

 $d_{j-1}$   $d_{j}$   $q_{i-1}$   $H_{i-1,j-1}$   $H_{i-1,j}$   $q_{i}$   $H_{i,j-1}$   $H_{i,j}$ 

Example



## Calculation of three candidate scores

 $H_{i,j}^{(0)}$ ,  $H_{i,j}^{(1)}$ , and  $H_{i,j}^{(2)}$  represent the three candidate scores of  $H_{i,j}$ . They are respectively calculated as:

$$H_{i,j}^{(0)} = H_{i-1,j} - g$$
 (vertical)  
 $H_{i,j}^{(1)} = H_{i,j-1} - g$  (horizontal)  
 $H_{i,j}^{(2)} = H_{i-1,j-1} + R_{a,b}$  (diagonal)

#### Exercise 2.4

Calculate the scores of  $H_{4,6}^{(0)}$ ,  $H_{4,6}^{(1)}$ , and  $H_{4,6}^{(2)}$  first and then update  $H_{4,6}$ .

Scoring scheme:

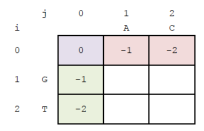
$$R_{ab} = 1 \text{ for } a = b$$
 $R_{ab} = 0 \text{ for } a \neq b$ 
 $R_{ab} = 0 \text{ for } a \neq b$ 
 $R_{ab} = 0 \text{ for } a \neq b$ 

## Initialization

The first row and the first column can be calculated independently from the adjacent cells.

$$H_{0,j}: j*-1*g$$
  
 $H_{i,0}: i*-1*g$ 

Example



## Exercise 2.5

Update all cells of Table 1 and 2. Use the scoring scheme in Exercise 2.4.

Table 1

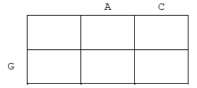
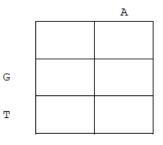


Table 2



## **Sub-solutions**

In DP, larger decisions recursively nest smaller decisions. For instance, Table S is included in Table L.

Table S

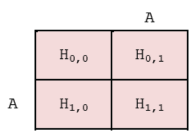


Table L

		A	G
	H <sub>0,0</sub>	H <sub>0,1</sub>	H <sub>0,2</sub>
A	H <sub>1,0</sub>	H <sub>1,1</sub>	H <sub>1,2</sub>
С	H <sub>2,0</sub>	H <sub>2,1</sub>	H <sub>2,2</sub>

## Pseudo-code of updating DP table for global alignment

## Algorithm 2.1: Update dynamic programming table for global alignment

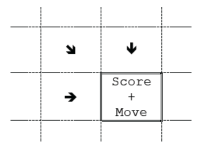
```
\begin{array}{l} \text{H}_{i,j}: \text{Dynamic programming table} \\ \text{R}_{a,b}: \text{Match/mismatch scores} \\ \text{g} : \text{Gap penalty} \\ \\ \text{// Initialization} \\ \text{for } i \leftarrow 0 \text{ to } m \text{ do} \\ \mid \text{H}_{i,0} \leftarrow i * - 1 * g; \\ \text{end} \\ \text{for } j \leftarrow 1 \text{ to } n \text{ do} \\ \mid \text{H}_{0,j} \leftarrow j * - 1 * g; \\ \text{end} \\ \\ \text{// Main loop for table update} \\ \text{for } i \leftarrow 1 \text{ to } m \text{ do} \\ \mid \text{for } j \leftarrow 1 \text{ to } n \text{ do} \\ \mid \text{H}_{i,j} \leftarrow max(\text{H}_{i-1,j} - \text{g}, \text{H}_{i,j-1} - \text{g}, \text{H}_{i-1,j-1} + \text{R}_{a,b}); \\ \mid \text{end} \\ \text{end} \\ \end{array}
```

# 2.5 Backtracking

Backtracking is a post-processing procedure to find the alignments that have yielded the best score.

## Store movement in cells

A table cell can be used for storing the movement.

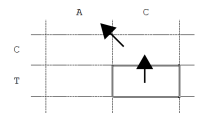


## Example

Cells with scores and directions

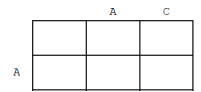
	A	С	
С	Score:1 Move:V	Score:3 Move:D	
Т	Score:0 Move:V	Score:2 Move:V	

Use arrows to indicate backtracking



## Exercise 2.6

Complete the DP table with scores and directions. What is the alignment with the best score?



Scoring scheme:  

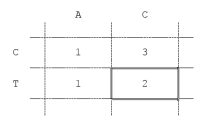
$$R_{ab} = 1 \text{ for } a = b$$
  
 $R_{ab} = 0 \text{ for } a \neq b$   
 $g = 1$ 

## Re-calculate candidate scores

Re-calculating the three candidate scores also reveals the movement.

$$H_{i,j}^{(0)} = H_{i-1,j} - g$$
 (vertical)  
 $H_{i,j}^{(1)} = H_{i,j-1} - g$  (horizontal)  
 $H_{i,j}^{(2)} = H_{i-1,j-1} + R_{a,b}$  (diagonal)

## Example



$$H_{i,j}^{(0)} = 3 - 1 = 2 = H_{i,j}$$
  $\checkmark (vertical)$   
 $H_{i,j}^{(1)} = 1 - 1 = 0 \neq H_{i,j}$  (horizontal)  
 $H_{i,j}^{(2)} = 1 + 0 = 1 \neq H_{i,j}$  (diagonal)

## Common mistake with backtracking

For the re-calculation approach, it is not to find  $max(H_{i-1,j}, H_{i,j-1}, H_{i-1,j-1})$ . You must re-calculate the candidates and then  $max(H_{i,j}^{(0)}, H_{i,j}^{(1)}, H_{i,j}^{(2)})$  to find the actual direction.

#### Implementation with recursive call

Recursive calls are usually used to implement DP backtracking.

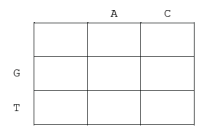
# Algorithm 2.2: DP backtracking

```
S_q: Sequence q
S_d: Sequence d
H<sub>i,i</sub>: Dynamic programming table
R<sub>a,b</sub>: Match/mismatch scores
    : Gap penalty
proc backTrack(i, j, A<sub>q</sub>, A<sub>d</sub>, k)
          : Index of sequence q
          : Index of sequence d
    j
    A_q: q part of alignment (stored in reverse order)
    A<sub>d</sub>: d part of alignment (stored in reverse order)
    k: Index for A_q and A_d
    // Need to implement recursion termination here
    //
                                                                                              // vertical
    if H_{i,j} = H_{i-1,j} - g then
         A_{q,k} \leftarrow S_{q,i};

A_{d,k} \leftarrow -;
         backTrack(i-1, j, A_q, A_d, k+1);
    end
    if H_{i,j} = H_{i,j-1} - g then
                                                                                          // horizontal
         A_{q,k} \leftarrow \ddot{,} - \ddot{,}
         A_{d,k} \leftarrow S_{d,i};
         backTrack(i, j-1, A_q, A_d, k+1);
    end
    if H_{i,j} = H_{i-1,j-1} + R_{S_{q,i},S_{d,i}} then
                                                                                              // diagonal
         \begin{array}{l} \mathbf{A_{q,k}} \leftarrow S_{q,i}; \\ \mathbf{A_{d,k}} \leftarrow S_{d,i}; \end{array}
         backTrack(i-1, j-1, A_q, A_d, k+1);
    end
end
```

## Exercise 2.7

Find the alignment with the best score.



Scoring scheme:  

$$R_{ab} = 1$$
 for  $a = b$   
 $R_{ab} = 0$  for  $a \neq b$   
 $g = 1$ 

# 2.6 Needleman-Wunsch algorithm

The method of using DP to solve global pairwise alignment is called the Needleman-Wunsch algorithm in the field of bioinformatics.

# Complexity

• Time: O(nm)

• Space: O(nm)

## Comparisons with other algorithms

The Needleman-Wunsch algorithm is similar to several algorithms.

## Divide and conquer algorithms

Sub-solutions must be independent with divide and conquer.

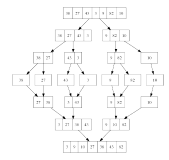


Figure 2.1: Merge sort (source: VineetKumar, Wikimedia Commons)

# Dijkstra's algorithm

Worst-case performance of Dijkstra:  $O(|E| + |V| \log |V|)$ 

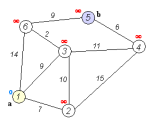


Figure 2.2: Dijkstra's algorithm (source: Ibmua, Wikimedia Commons)