SEQUENCE ALIGNMENT ALGORITHMS

ALGORITMI PER L’ALLINEAMENTO DI SEQUENZE

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Part of ALiBio project
http://www.alibio.org/
Preface

ALiBio Goal

In a few words the goal of ALiBio is to give libraries and efficient fundamental algorithms to be used when developing applications in the bioinformatics field.

The large amount of biological data available nowadays makes the need of well-implemented efficient algorithm really stringent. Various projects (Open Bioinformatics Foundation) are already giving a number of libraries to the developer community.

The focus of those projects has always been on the ease of use, especially for people coming from the Biology field, and efficiency issues have not been the top priority.

On the other hand ALiBio is targetting the developer community with a strong CS background. ALiBio is suited for developing highly optimized applications where efficiency is paramount.

Two more issues that will receive a lot of attention in this project are testing and documentation. All libraries and algorithms that will be included in ALiBio must have an extensive suite of regression tests and must be clearly documented. In fact we require the use of noweb[10] for developing code to be included in ALiBio.

The advantages of Free Software (cfr.2.1) are well-known. ALiBio is a free library, at the same time it can be used also for developing non-free software. While we encourage the development of free software, we do not require it in order to use ALiBio.
Sequence Alignment Algorithms

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# Contents

1 Introduction ........................................... 3  
   1.1 Objective of the stage ........................... 3  
   1.2 Sequence Alignment ............................... 3  
   1.3 Specifications of the project .................... 4  
   1.4 Phases of development ............................ 6  

2 Sequence alignment algorithms ...................... 9  
   2.1 Definitions ........................................ 9  
   2.2 Problem formalization ............................. 11  
   2.3 Classic algorithms ................................ 12  
      2.3.1 Dynamic Programming ........................ 12  
      2.3.2 Needleman-Wunsch algorithm ................... 12  
      2.3.3 Smith-Waterman algorithm ..................... 13  
   2.4 Affine gap penalties .............................. 13  
   2.5 Linear space alignment ............................ 14  
      2.5.1 Linear space cost-only alignment .............. 14  
      2.5.2 Track back the alignment ...................... 14  
      2.5.3 Time and space analysis of the algorithm ....... 15  

3 Implementation ....................................... 16  
   3.1 Library structure .................................. 16  
   3.2 The licence ........................................ 17  
   3.3 Included libraries ................................. 17
CONTENTS

3.4 The `score_matrix` class .............................................. 18
   3.4.1 `score_matrix` interface ..................................... 19
   3.4.2 `score_matrix` implementation ............................... 21
3.5 Sequence alignment class ........................................... 27
   3.5.1 `seq_alignment` interface .................................... 27
   3.5.2 `seq_alignment` implementation ............................. 30
   3.5.3 Classic algorithms implementation ......................... 34
   3.5.4 Aligning using affine gap penalties ....................... 37
   3.5.5 Aligning in linear space .................................... 45

4 Tests ............................................................................. 57
   4.1 Regression tests ..................................................... 57
      4.1.1 Structure of the test program ............................. 58
      4.1.2 Structure of the test file ................................. 59
      4.1.3 Test program for `score_matrix` class .................. 60
      4.1.4 Test program for `seq_alignment` class .................. 65
   4.2 Memory leaks test ................................................... 73
   4.3 Random tests ........................................................ 73

5 Examples ........................................................................ 78

6 Performance analysis ...................................................... 81
   6.1 Generating data .................................................... 81
   6.2 Results ..................................................................... 87
   6.3 Performance comparison ............................................. 89
      6.3.1 Results .......................................................... 92

7 Conclusions .................................................................... 95
   7.1 Possible future improvements ................................... 95

8 Quick Methods reference ................................................ 96
   8.1 `score_matrix` Class ............................................... 96
   8.2 `seq_alignment` Class .............................................. 97
Chapter 1

Introduction

1.1 Objective of the stage

The purpose of this stage is to realize an efficient implementation of algorithms for optimal resolution of the Peerwise Sequence Alignment problem which has a great biological interest being a key to finding important regions in the genome, determining its functions and uncovering evolutionary forces. This particular stage is part of the ALiBio project[10] which aim is to provide a collection of useful libraries for the development of bioinformatics programs, with special regard to the efficiency of the implementation.

When I started the stage, the data structures, useful to represent biological information such as sequences of DNA, RNA and proteins, were already implemented. I will base my algorithm implementation upon these data structures.

1.2 Sequence Alignment

In detail, I will implement algorithms for global alignment (cfr.2.1) local alignment (cfr.2.1) and semiglobal alignment (cfr.2.1) following the specifications of the project. This algorithms are designed to find the best possible alignment (cfr.2.1) between two given sequences of symbols. Alignment algorithms are used to compare the sequence homology between two protein or DNA sequences. These programs find the best match between the two sequences. Occasionally gaps need to be introduced to make the two sequences align.

Global alignment algorithms attempt to match sequences from end to end, even though parts of the alignment are not very convincing. for example:

    CATTTAGATT-C
    X|| |X|| X
    --GTT--GTTTAT

On the other hand local alignment algorithms search for segments of the two sequences that match well. Using the same sequences as above, we could get:
1.3 Specifications of the project

Developing the stage I followed the guidelines defined by the AliBio project.

Literate Programming

The first requisite is to produce the job using a tool for literate programming\cite{9} called noweb\cite{10}. Literate programming is a methodology that combines a programming language with a documentation language, thereby making programs more robust, more portable, more easily maintained, and arguably more fun to write than programs that are written only in a high-level language. The main idea is to treat a program as a piece of literature, addressed to human beings rather than to a computer. The program is also viewed as a hypertext document, rather like the World Wide Web.

Languages

As documentation language ALiBio uses the \LaTeX\ writing system, which produces high quality scientific and mathematical documentation.

ALiBio project is implemented with the C++ programming language which, along with C, is probably the most used by developers of scientific applications all over the world. This choice guarantees a high portability with all the compilers that respect the ISO standard, and therefore a high compatibility of the code with any hardware platform.

Coding Conventions

To obtain a sufficient uniformity in the produced code developers should comply with the following conventions:

- Filenames: All in lower case and no spaces
- C++ files suffix: .cpp (programs), .hpp (libraries)
1.3 Specifications of the project

- Code lines length: All code lines must have 80 character at most
- The code: Must be written in lower case, using underscore as word separator
- Variable names: Descriptive names are required for global variables (eg. `string name;`), short names for local vars (eg. `string s;`)
- Functions names should be active verbs (eg. `get_seq()`, not `seq()`)  
- Chunk names must not contain a name between double square brackets
- Printing functions: Generally, the output should not begin with any newline, but should end with a newline.
- Indenting: Four spaces with respect to the block. Four blanks, must be used. tab is not allowed
- Indentations must be used for iterations (while, do), conditional instructions (if), methods and declarations (public, private, protected): Indentations must not be used for code included inside namespaces or classes;
- Block structures: The left brace `{` must be on the same line as the first instruction of the block, separated by a space; The right brace `}` must be on a new line, in the same column as the beginning of the block;
- Comments should not be used frequently, and only used for improving code readability. The real documentation must be in the documentation chunks
- Whenever possible, all assertions should be placed between the declarations and the algorithmic sections
- Blank lines should be used for readability reasons
- Namespaces must be specified for all instructions and not-built-in types (eg. `std::cout`, not `cout`)
- ALiBio’s namespace is alibio. All produced code must be within such namespace
- In order to avoid possible multiple inclusion, all libraries must check inclusions using opportune preprocessor instructions:
- The template construct (along with the type parameters), should be placed on the line before the class definition
- Each member functions implementation should be preceded by the template line too. Typically, when dealing with a complex return value, that too should be placed on a separate line. Moreover, the function name should be preceded by the class name(and type parameters), on the same line and separated by the scope operator.
Programming Paradigms

Besides exploiting the fundamental characteristics of C++ such as object orientation, the project also embraces the concept of Generic Programming as offered by the Standard Template Library\textsuperscript{12} that widely used all over the project. Generic programming is a datatype-independent way of programming computers. This means that the same source code can be used regardless of the datatype the code will be instantiated with or passed as parameters. For example the type \texttt{vector} of the STL allows to define a vector of elements, whose type can be freely chosen at compile time, and at the same time it offers a whole set of efficient functions for its management. This fact guarantees a high flexibility using the libraries.

Useful libraries

To develop the project another important library has been used: the Boost Library\textsuperscript{13} which offers a series of optimized tools for managing data structures as multidimensional matrices, lists, graphs and trees but also mathematical functions, input/output functions and many more.

Licence

From a legal and distributive point of view, ALiBio adopts a free software licence named LGPL, making the code and the documentation accessible to everyone.

External developers who want to contribute to ALiBio are wellcome, anyway to maintain the necessary uniformity of the final work, everyone should follow the project specifications and the chosen conventions for code and documentation writing.

Anyone can freely include the AliBio libraries inside its own program or freely redistributes the libraries under the terms of the GNU Lesser General Public License.

1.4 Phases of development

Learning the tools

Before being able to proceed with the coding phase, I dedicated a period of time studying the necessary tools for the realization of the project. All the software used (noweb, \LaTeX, Gnu C++ Compiler, CVS, GAWK) is freely available for the Linux operating system, so I had the opportunity to deepen my knowledge of Linux as a development platform and of the GNU development toolchain, in particular the Make program.

Although I already knew \LaTeX\textsuperscript{14}, I was completely new to noweb\textsuperscript{10} and the literate programming in general. I also needed to improve my knowledge about the CVS\textsuperscript{15} versioning system.
Concerning the programming part of the work, I already had some C++ knowledge but I had to study the principles on which the Standard Library has been developed in particular the Templates mechanism.

In my work I used some functions taken form the Boost Library[13] so I had to study their on-line documentation.

I also studied the parts of ALiBio that had been already realized, In this way I learned not only how to use the library I needed but also the peculiar coding and documenting style adopted in the project.

Study of the algorithms

Once learned the necessary tools for the realization of the project, I went on studying the algorithms that I had to implement using the original articles written by their authors and a lot of other informations found on the World Wide Web.

Algorithm implementation

To implement the alignment algorithms, I needed to implement a specific class named \( \text{score\_matrix} \) to contain the scoring schema needed for the alignment calculation. The object instantiated from this class will contain a value for each possible couple of elements of the used alphabet in a bi-dimensional matrix and the costs for inserting and extending gaps in the alignment. The class also implements the needed functions to set or modify these values.

The alignment class \( \text{seq\_alignment} \) contains the functions that implement the alignment algorithms and a function to print the calculated alignment. An alignment object is constructed from a \( \text{score\_matrix} \) object and two sequences in various possible formats, the alignment functions (which are the heart of the algorithm) actually calculate the alignment and prepare the output lines that will be printed by \( \text{print\_alignment} \).

Realization of example programs

To facilitate the job of the developers who want to use the library, I wrote an example program which show how it is possible to include and to exploit the functionalities of the library in their programs. The program shows how to create the needed \( \text{score\_matrix} \) object and modify its scoring system, then we show how to instantiate a \( \text{seq\_alignment} \) object with two arbitrary sequences and calculate their different alignments. The program also show how to print the obtained result on standard output.

Testing

In the project a lot of importance is given to the tests phase. Once the code has been written, it’s necessary to write some special tests able to verify its correctness. The adopted technique
is the automatic regression tests that allows to compare the actual output of the test with the expected output or the output of a previous version of the implementation. The project adopts some conventions to write these tests. The purpose is to check the input and the output attended for every single test and to automate their execution accomplished by a dedicated script. Respecting these conventions, the program is written to develop the tests on every single element of the library. The tests are performed in a particular order trying to avoid, as far as possible, the test of a function that calls a function not yet tested.

In my studies, two test programs have been implemented: one for testing the correctness and functionalities of the \( \text{score\_matrix} \), the other one to verify the correct execution of the \( \text{seq\_alignment} \) (which implements the alignment algorithms). The files used as input of the script which executes the test programs contain the input for each test and the expected output. This will be compared with the actual output by the script which establishes the success rather than the failure of the execution.

The implemented algorithms make widely use of low-level access to memory, C++ doesn’t guarantee from memory leaks so we used a specific tool to detect eventual improper memory allocation and access.

To ensure the correctness of the algorithms over a large set of inputs I developed a randomized test that calls the alignment functions over random sequences and verifies the output. These test revealed a series of error in the implementation that weren’t visible previously.

The testing phase of the work took a great amount of time, comparable to the actual development phase.

**Analysing performances**

Other libraries already exist to solve the peerwise alignment problem so we could test our implementations against EMBOSS[16] (one of the most widely used) to compare results and performances.

**Writing the documentation**

The documentation of the job (written in English) is constituted by:

1. an introductory part that describes the global project,
2. a section containing the necessary definitions for a better understanding of the code,
3. a wide description of the code to explain its functionality,
4. the description of the example programs and the tests,
5. the description of the performance analysis carried out
6. the synopsis tables containing, for each function, a brief description, the necessary prerequisites and the effect of its execution.
Chapter 2

Sequence alignment algorithms

2.1 Definitions

alphabet

An alphabet $\Sigma$ is a non-empty finite set of characters.

sequence

A sequence is a list of elements of a given alphabet arranged in a "linear" fashion, such that the order of the members is well defined and significant.

alignment

A one-to-one matching of two sequences so that each character in a pair of sequences is associated with a single character of the other sequence or with a gap.

alignment score

A numerical value that describes the overall quality of an alignment. Higher numbers correspond to higher similarity.

match

In sequence alignment, the existence of the same character in a homologous position in both sequences.
mismatch

In sequence alignment, the existence of different characters in a homologous position in the two sequences.

gap

An insertion or a deletion. In sequence alignment, a pair containing a special null character "]-]."

gap opening penalty

The length-independent cost of opening a gap in a sequence alignment.

gap extension penalty

The cost of extending by one character a pre-existing gap in a sequence alignment.

linear gap penalty

A scoring system for gaps within alignments that charges a penalty for the existence of a gap proportional to the gap’s length.

affine gap penalty

A scoring system for gaps within alignments that charges a penalty for the existence of a gap and additional penalty proportional to the gap’s length.

global alignment

An optimal alignment that includes all characters from each sequence but may miss short regions of high local similarity.

Global alignments are most useful for closely related sequences of known homology.

local alignment

An optimal alignment that includes the most similar local region but may include only short portions of the sequences that were used to calculate the alignment.

Local alignments are especially useful for distantly related sequences.
semiglobal alignment

An optimal alignment that includes a prefix of one of the two sequences and a suffix of the other, in other words a global alignment where initial end terminal gaps are free.

Free software

Free software is a matter of the users’ freedom to run, copy, distribute, study, change and improve the software. More precisely, it refers to four kinds of freedom, for the users of the software:

1. The freedom to run the program, for any purpose (freedom 0).
2. The freedom to study how the program works, and adapt it to your needs (freedom 1). Access to the source code is a precondition for this.
3. The freedom to redistribute copies so you can help your neighbor (freedom 2).
4. The freedom to improve the program, and release your improvements to the public, so that the whole community benefits (freedom 3). Access to the source code is a precondition for this.

A program is free software if users have all of these freedoms. Thus, you should be free to redistribute copies, either with or without modifications, either gratis or charging a fee for distribution, to anyone anywhere. Being free to do these things means (among other things) that you do not have to ask or pay for permission.

You should also have the freedom to make modifications and use them privately in your own work or play, without even mentioning that they exist. If you do publish your changes, you should not be required to notify anyone in particular, or in any particular way.


2.2 Problem formalization

The Sequence Alignment Problem\cite{1} can be formalized as follows: Let $\Sigma$ be a $k$-letter alphabet, and let $A$ and $B$ be two sequences over $\Sigma$. An alignment of strings $A = a_0, \ldots, a_{m-1}$ and $B = b_0, \ldots, b_{m-1}$ is a $2 \times l$ matrix $M^0$ ($l \geq n,m$), such that the first (second) row of $M^0$ contains the characters of $A$ ($B$) in order interspersed with $l - n$ ($l - m$) spaces. We assume that no column of the alignment matrix contain two spaces.

For every pair of symbols $(a_i, b_j)$ from $\Sigma$ is defined a similarity score, the score of the alignment is defined as the sum of the scores of its columns.

The Sequence Alignment problem is to find the alignment of sequences $A$ and $B$ of maximal score.
2.3 Classic algorithms

In 1970 Saul Needleman and Christian Wunsch proposed[3] an algorithm for solving the SA problem which used a technique called Dynamic Programming and guarantees to find the global alignment (cfr.2.1) with the maximum score.

In 1981 Temple Smith and Michael Waterman proposed[4] a modification of the original algorithm to find the optimal local alignment (cfr.2.1).

These algorithms are quadratic in time and space complexity and are the fundamental of all the other algorithms we will discuss in this paper.

2.3.1 Dynamic Programming

Dynamic Programming[2] was invented by the mathematician Richard Bellman in 1953, it is a method for reducing the runtime of algorithms exhibiting the properties of overlapping subproblems and optimal substructure where overlapping subproblems means that the problem can be broken down into subproblems which are reused several times and optimal substructure means that optimal solutions of subproblems can be used to find the optimal solutions of the overall problem.

2.3.2 Needleman-Wunsch algorithm

Every possible combination of chars in two sequences are represented in a 2-dimensional array $M^0$ with an extra row and column added to allow the alignment to begin with a gap of any length in either sequence.

<table>
<thead>
<tr>
<th>$M^0$</th>
<th>-</th>
<th>$b_0$</th>
<th>$b_1$</th>
<th>$b_{j-1}$</th>
<th>$b_j$</th>
<th>$b_{n-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>-</td>
<td>$M^0_{0,0}$</td>
<td>$M^0_{0,1}$</td>
<td>$M^0_{0,j}$</td>
<td>$M^0_{0,j+1}$</td>
<td>$M^0_{0,n}$</td>
<td></td>
</tr>
<tr>
<td>$a_0$</td>
<td>$M^0_{1,0}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$a_{i-1}$</td>
<td>$M^0_{i,0}$</td>
<td>$M^0_{i,j}$</td>
<td>$M^0_{i,j+1}$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$a_i$</td>
<td>$M^0_{i+1,0}$</td>
<td>$M^0_{i+1,j}$</td>
<td>$M^0_{i+1,j+1}$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$a_{m-1}$</td>
<td>$M^0_{m,0}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>$M^0_{m,n}$</td>
</tr>
</tbody>
</table>

Scores are filled in for each square, starting from $M^0_{1,1}$, searching for the maximum possible value using the recurrence below in which $M^0_{i+1,j+1}$ is the score of an optimal alignment between $A_i = a_0, \ldots, a_i$ and $B_j = b_0, \ldots, b_j$

$$M^0_{i+1,j+1} = \max \begin{cases} M^0_{i,j+1} + \text{score}(a_i, -) \\ M^0_{i+1,j} + \text{score}(-, b_j) \\ M^0_{i,j} + \text{score}(a_i, b_j) \end{cases}$$
The first raw and column are initialized with the score of an initial gap, the optimal alignment score can be read in $M_{m,n}^0$.

### 2.3.3 Smith-Waterman algorithm

The local alignment problem correspond to finding the best global alignment between all of the substrings of $A$ and $B$. An efficient way to achieve this is non-considering alignments with negative scores, this is the modified recurrence:

$$M_{i+1,j+1}^0 = \max \begin{cases} 
0 \\
M_{i,j+1}^0 + \text{score}(a_i, -) \\
M_{i+1,j}^0 + \text{score}(-, b_j) \\
M_{i+1,j+1}^0 + \text{score}(a_i, b_j)
\end{cases}$$

The largest value found all over $M^0$ represents the score of the best local alignment between $A$ and $B$.

### 2.4 Affine gap penalties

Mutations are usually manifestations of errors in DNA replications. Nature frequently deletes or inserts entire substrings as a unit, as opposite to deleting or inserting individual nucleotides.

It is natural to assume that the score of a gap consisting of $x$ spaces is not just the sum of scores of $x$ spaces. In the Affine gap penalties scheme, the score for a gap of length $x$ is $-(g + xe)$, where $g > 0$ is the penalty for the introduction of the gap and $e > 0$ is the penalty for adding a symbol to the gap.

To calculate affine gap penalties we need two more matrix: $M^1$ for keeping the best alignment ending with a gap in $A$ and $M^2$ for the best alignment ending with a gap in $B$, we calculate the scores using the following three recursions:

$$M_{i+1,j+1}^1 = \max \begin{cases} 
M_{i,j+1}^1 - g \\
M_{i,j+1}^1 - e
\end{cases}$$

$$M_{i+1,j+1}^2 = \max \begin{cases} 
M_{i+1,j}^1 - g \\
M_{i+1,j+1}^0 - e
\end{cases}$$

$$M_{i+1,j+1}^0 = \max \begin{cases} 
M_{i,j}^0 + \text{score}(a_i, b_j) \\
M_{i+1,j+1}^1 \\
M_{i+1,j+1}^2
\end{cases}$$
2.5 Linear space alignment

When computing optimal sequence alignments time is not the only limiting factor, the algorithms above use a 2dimensional matrix leading to a space complexity $O(mn)$. Aligning genomes of different organisms, each 1 billion pairs long, would far exceed the RAM in current computers.

In 1988 Eugene W. Myers and Webb Miller published an article\[5\] that showed how to use a method proposed\[6\] by Hirshberg in 1975 to develop a linear-space version of the Alignment algorithm with affine gap penalties.

2.5.1 Linear space cost-only alignment

It’s easy to compute $M_{m,n}^0$ in linear space because only the scores from the previous row of the matrix are needed therefore the alignment scores in the rows before $i$ can be discarded while computing alignment scores for row $i+1$. This observation leads to an algorithm for a cost-only alignment with a space complexity $O(n)$.

2.5.2 Track back the alignment

Finding the actual alignment requires backtracking through the entire matrix after all of the rows and columns have been filled. To compute both the optimal score and alignment, one option would be to use a divide et impera approach\[2\].

Let’s call $A^{rev}$ and $B^{rev}$ the reverse of $A$ and $B$. $M_{i,j}^{0\text{rev}}$ is the optimal score of aligning $A^{rev}_i = a_{m-1}, \ldots, a_i$ and $B^{rev}_j = b_{n-1}, \ldots, b_j$.

As explained above the values of $M_{i,j}^{0\text{rev}}$ can be computed in time $O(mn)$ and space $O(n)$.

We can prove true the following equivalence:

$M_{m,n}^0 = \max_{k=0\ldots n} M_{m,2}^0 + M_{2,n-k}^{0\text{rev}}$

It states that the middle point of $A$ has to align to some point in $B$, $k$ is the optimal breakpoint in $B$ obtained by maximizing over all possible ways of bisecting the sequence $B$ aligning the first part with $A$ until the middle point and the second part with $A$ after the middle point.

<table>
<thead>
<tr>
<th>$M^0$</th>
<th>$- ,$</th>
<th>$b_0 ,$</th>
<th>$\ldots ,$</th>
<th>$b_{k-1} ,$</th>
<th>$\ldots ,$</th>
<th>$b_{n-1} ,$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$a_0$</td>
<td>$M_{0,0}^0$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\vdots$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$a_{m-1}$</td>
<td>$M_{m,2}^0$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\vdots$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$a_{m-1}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>$M_{m,n}^0$</td>
</tr>
</tbody>
</table>

The square $M_{m,2}^0$ divides the matrix into four parts; the same algorithms is called recursively over the submatrices $[M_{0,0}^0, M_{0,2}^0]$ and $[M_{2,k}^0, M_{m,n}^0]$. Keeping track of the values assumed by $k$, one can determine also the actual alignment in linear space.
2.5 Linear space alignment

2.5.3 Time and space analysis of the algorithm

To compute the middle point of an alignment between two sequences of size m and n:

- Space: $2n$
- Time: $c \times mn$ for some constant $c$

Then, left and right calls cost $c \times (\frac{m}{2} \times k + \frac{m}{2} \times (n - k)) = \frac{c \times mn}{2}$

All recursive calls cost:

- Space: $O(n + m)$ to store the optimal alignment
- Time: $c \times mn + \frac{c \times mn}{2} + \frac{c \times mn}{4} + \ldots = 2c \times mn$ that is $O(mn)$
Chapter 3

Implementation

3.1 Library structure

To implement sequence alignment algorithms we decided to divide the work in two different classes:

1. \(\textquoteright\text{score\_matrix} 8\) class implementing a scoring system.
2. \(\textquoteright\text{seq\_alignment} 25\) class implementing the alignment algorithms.

Here is the file main structure:

\begin{verbatim}
#include <two_sequence.hpp>

namespace {
    class score_matrix;
    class seq_alignment;
}
\end{verbatim}

Root chunk (not used in this document).
3.2 The licence

As stated before, ALiBio project is published under a free software licence named LGPL, the program begins with the opportune references

```
#include <stdlib.h>
#include <assert.h>
#include <iostream>
#include <new>
```

This code is used in chunks 1, 49, 61, 76, 78, 83, and 91–93.

3.3 Included libraries

For our implementation we used a lot of external libraries: Let’s begin with libraries from the standard c++

```
#include <stdlib.h>
#include <assert.h>
#include <iostream>
#include <new>
```

This definition is continued in chunks 4 and 5.

This code is used in chunk 1.
3.4 The score_matrix class

We also needed some data structures that was already implemented as parts of the ALiBio project to represent biological structures:

```cpp
#include <alibio/sequence.hpp>
#include <alibio/bio_string.hpp>
#include <alibio/empty.hpp>
```

This code is used in chunk 1.

From the boost project we used the Boost.MultiArray library which provides a generic N-dimensional array concept definition and common implementations of that interface.

```cpp
#include <boost/multi_array.hpp>
```

This code is used in chunk 1.

We will use the namespace std

```cpp
using namespace std;
```

This definition is continued in chunk 7.

This code is used in chunk 1.

Every element of this library belongs to the namespace alibio.

```cpp
namespace alibio
```

This code is used in chunk 1.

3.4 The score_matrix class

Let’s separate class interface from implementation

```cpp
(score_matrix 8)≡
(score_matrix interface 9)
(score_matrix implementation 13)
```

This code is used in chunk 1.
3.4 The score_matrix class

3.4.1 score_matrix interface

Here is the declaration of the score_matrix class: it’s a template class in which the parameter T is the type of the costs: we could use a small type (ex. short) to save memory or a floating point type (ex. double) if we need decimal positions. The default type for T has been set to float because of its flexibility.

```
(score_matrix interface 9)≡
  template <class T=float>
  class score_matrix{
  private:
    (private variables of score_matrix 10)
    (private functions of score_matrix 11)
  public:
    (functions exported by score_matrix 12)
  };
```
This code is used in chunk 8.

Private variables of score_matrix

The (score_matrix 8) class relies on the boost::multi_array for representing the actual matrix data, the alphabet is memorized in a vector<char> object according to the alibio::alphabet implementation. It also keeps an int for the alphabet size.

```
(private variables of score_matrix 10)≡
  boost::multi_array<T, 2> *data;
  vector<char> alphabet;
  T gap_cost, extend_gap_cost;
  int lnt;
```
This code is used in chunk 9.

Private functions of score_matrix

The (score_matrix 8) class stores the alphabet in a vector, a recursive private function is needed to find a char’s position in logarithmic time.

```
(private functions of score_matrix 11)≡
  int find_char_pos(char letter, int low, int high) const;
```
This code is used in chunk 9.
3.4 The score_matrix class

**Functions exported by score_matrix**

The ⟨score_matrix⟩ class provides several functions which offer a useful interface to the internal data structures. The following is a brief description of each function:

- ⟨score_matrix constructor⟩: Creates a score matrix for the given alphabet: default scores are 1 for matching and -1 for gaps and substitution of a symbol.
- ⟨score_matrix destructor⟩: frees memory after a deallocation.
- ⟨set_score⟩: Sets the score of switching between two given chars.
- ⟨get_score⟩: Returns the score of switching between two given chars.
- ⟨set_match_score⟩: Sets the score of symbol matches.
- ⟨set_mismatch_score⟩: Sets the score of symbol mismatches.
- ⟨set_gap_cost⟩: Sets the cost of inserting and extending a gap.
- ⟨get_gap_cost⟩: Returns the cost of inserting a gap.
- ⟨get_extend_gap_cost⟩: Returns the cost of extending a gap.
- ⟨print⟩: Prints on the standard output the score matrix.

```cpp
(score_matrix(const alibio::alphabet& alph);
 ~score_matrix();
 void set_score(char a, char b, T val);
 T get_score(char a, char b) const;
 void set_match_score(T val);
 void set_mismatch_score(T val);
 void set_gap_cost(T val);
 void set_gap_cost(T val, T ext_val);
 T get_gap_cost() const;
 T get_extend_gap_cost() const;
 void print() const;
```

This code is used in chunk 9.
3.4 The score_matrix class

3.4.2 score_matrix implementation

Here is the implementation of the declared functions:

\[
\begin{align*}
&\text{score_matrix implementation 13} \\
&\text{score_matrix constructor 14} \\
&\text{score_matrix destructor 15} \\
&\text{find_char_pos 16} \\
&\text{set_score 17} \\
&\text{get_score 18} \\
&\text{set_match_score 19} \\
&\text{set_mismatch_score 20} \\
&\text{set_gap_cost 21} \\
&\text{get_gap_cost 22} \\
&\text{get_extend_gap_cost 23} \\
&\text{print 24}
\end{align*}
\]

This code is used in chunk 8.
3.4 The score_matrix class

Constructors implementation

The constructor implementation has an alibio::alphabet object as parameter. It adds the symbol - to the given alphabet for representing a gap in the aligned sequence and then creates the \(\text{score\_matrix} \) object using the boost::multi_array class. The size of the matrix is taken from the alphabet size and is kept in the \text{lnt} variable. Finally it populates the matrix with some default values: 1 for matching, -1 for symbol switching. Gap cost is initialized to 1.

\(\text{score\_matrix}\) constructor

\[
\text{template <class T>}
\]
\[
\text{score\_matrix<T>::score\_matrix(const alibio::alphabet &alph)} \{\}
\]
\[
\text{\quad vector<alibio::symbol> symbols=alph.get\_all\_symbols();}
\]
\[
\text{\quad lnt=symbols.size();}
\]
\[
\text{\quad alphabet.reserve(lnt);} \}
\]
\[
\text{\quad vector<alibio::symbol>::iterator It;}
\]
\[
\text{\quad for(It=symbols.begin(); It!=symbols.end(); It++)}
\]
\[
\text{\quad \quad alphabet.push\_back((*It).get\_id());} \}
\]
\[
\text{\quad try{}
\text{\quad \quad data=new boost::multi\_array< T, 2>(boost::extents[lnt][lnt]);}
\text{\quad \quad } \}
\]
\[
\text{\quad catch(const bad\_alloc x){}
\text{\quad \quad cerr<< "Out of memory in score\_matrix(alphabet alph): "<<x.what()<<"n";}
\text{\quad \quad abort();} \}
\]
\[
\text{\quad //we sort the vector for fast (logaritmic) searching}
\text{\quad sort(alphabet.begin(),alphabet.end());} \}
\]
\[
\text{\quad vector<char>::iterator It1, It2;}
\]
\[
\text{\quad for (It1=alphabet.begin(); It1!=alphabet.end(); It1++)}
\text{\quad \quad for (It2=alphabet.begin(); It2!=alphabet.end(); It2++)}
\]
\[
\text{\quad \quad if(It1==It2)}
\text{\quad \quad \quad this->set\_score(*It1,*It2,(T)1); //default score for match}
\text{\quad \quad \quad else}
\text{\quad \quad \quad this->set\_score(*It1,*It2,(T)-1); //default score for mismatch}
\text{\quad \quad gap\_cost=(T)1; //default cost for gap opening}
\text{\quad \quad extend\_gap\_cost=(T)1; //default cost for gap extending}
\text{\quad \quad } \}
\]
This code is used in chunk 13.
3.4 The score_matrix class

Destructor implementation

It frees the memory allocated by the constructor

(\texttt{score\_matrix destructor 15}) \equiv
\begin{verbatim}
template <class T>
score\_matrix<T>::\texttt{\~}score\_matrix()
{
  delete data;
}
\end{verbatim}

This code is used in chunk 13.

Finding a char

A function for finding a char in the alphabet with logarithmic cost.

We could get a constant cost with an hash table but alphabets are normally small and calculating the hashes would take long...

(\texttt{find\_char\_pos 16}) \equiv
\begin{verbatim}
template <class T>
int score\_matrix<T>::\texttt{find\_char\_pos}(char letter, int low, int high) const{
  if (high < low) return -1;

  int mid = (high + low) / 2;

  if (alphabet[mid] == letter) return mid;

  if (alphabet[mid] > letter) {
    return find\_char\_pos (letter, low, mid-1);
  } else {
    return find\_char\_pos (letter, mid+1, high);
  }
}
\end{verbatim}

This code is used in chunk 13.
3.4 The score_matrix class

Setting the score

This function permits to set the score of an alignment between the chars a and b to the value val which are its parameters, if the chars don’t belong to the alphabet, the function aborts.

\begin{verbatim}
(set_score 17)≡
//abort if doesn't find the given char
template <class T>
void score_matrix<T>::set_score(char a, char b, T val) {
    int x=find_char_pos(a, 0, lnt);
    int y=find_char_pos(b, 0, lnt);

    if(x==-1) {
        cerr << "Error: char " " not present in the alphabet";
    abort();
    }
    if(y==-1) {
        cerr << "Error: char " " not present in the alphabet";
    abort();
    }

    (*data)[x][y]=val;
    return;
}
\end{verbatim}

This code is used in chunk 13.

Getting the score

This function returns the score of alignment between the chars a and b passed as parameters, if the chars don’t belong to the alphabet, the function aborts.

\begin{verbatim}
(get_score 18)≡
//abort if doesn't find the given char
template <class T>
T score_matrix<T>::get_score(char a, char b) const {
    int x=find_char_pos(a, 0, lnt);
    int y=find_char_pos(b, 0, lnt);

    if(x==-1) {
        cerr << "Error: char " " not present in the alphabet";
    abort();
    }
    if(y==-1) {
        cerr << "Error: char " " not present in the alphabet";
    abort();
    }

    return (*data)[x][y];
}
\end{verbatim}

This code is used in chunk 13.
3.4 The score_matrix class

Setting score for matching

This function permits to set the score of an alignment between any two identical chars to a desired value val.

\[ \text{set_match_score} \]

\[
\begin{align*}
\text{template} & \quad <\text{class } T> \\
\text{void} & \quad \text{score_matrix}<T>::\text{set_match_score}(T \text{ val})\{ \\
& \quad \text{for (int } i=0; i<\text{lnt}; i++) \\
& \quad \quad (*\text{data})[i][i]=\text{val}; \\
& \quad \}
\end{align*}
\]

This code is used in chunk 13.

Setting score for mismatching

This function permits to set the score of an alignment between any two different chars to a desired value val.

\[ \text{set_mismatch_score} \]

\[
\begin{align*}
\text{template} & \quad <\text{class } T> \\
\text{void} & \quad \text{score_matrix}<T>::\text{set_mismatch_score}(T \text{ val})\{ \\
& \quad \text{for (int } j=0; j<i; j++)\{ \\
& \quad \quad (*\text{data})[i][j]=\text{val}; \\
& \quad \quad (*\text{data})[j][i]=\text{val}; \\
& \quad \}
\end{align*}
\]

This code is used in chunk 13.

Setting the cost for opening and extending gaps

These functions permit to set the cost for opening and extending a gap in the alignment, if no extending gap cost is passed as parameter the opening gap cost will be used.

\[ \text{set_gap_cost} \]

\[
\begin{align*}
\text{template} & \quad <\text{class } T> \\
\text{void} & \quad \text{score_matrix}<T>::\text{set_gap_cost}(T \text{ val})\{ \\
& \quad \text{gap_cost}=\text{val}; \\
& \quad \text{extend_gap_cost}=\text{val}; \\
& \}
\end{align*}
\]

\[
\begin{align*}
\text{template} & \quad <\text{class } T> \\
\text{void} & \quad \text{score_matrix}<T>::\text{set_gap_cost}(T \text{ val}, T \text{ ext_val})\{ \\
& \quad \text{gap_cost}=\text{val}; \\
& \quad \text{extend_gap_cost}=\text{ext_val}; \\
& \}
\end{align*}
\]

This code is used in chunk 13.
3.4 The score_matrix class

Getting the cost for opening and extending gaps

```cpp
22 (get_gap_cost 22)\equiv
    template <class T>
    T score_matrix<T>::get_gap_cost() const{
        return gap_cost;
    }
This code is used in chunk 13.

23 (get_extend_gap_cost 23)\equiv
    template <class T>
    T score_matrix<T>::get_extend_gap_cost() const{
        return extend_gap_cost;
    }
This code is used in chunk 13.
```

Printing the score matrix

A function that prints on the standard output a visualizations of the alphabet used, of the score matrix and gap costs

```cpp
24 (print 24)\equiv
    template <class T>
    void score_matrix<T>::print() const{
        //printing the score matrice
        int m=lnt;

        cout << "\t";
        for (int j=0;j<m;j++){
            cout<< alphabet[j] << "\t";
        } cout << \n;

        for (int i=0;i<m;i++){
            cout << alphabet[i] << "\t";
            for (int j=0;j<m;j++){
                cout << (*data)[i][j] << "\t";
            }
            cout << "\n";
        }

        cout << "gap cost=\t" << gap_cost << "\n";
        cout << "gap extension cost=\t" << extend_gap_cost << "\n";
    }//end print()
This code is used in chunk 13.
```
3.5 Sequence alignment class

We need to separate class interface from implementation

\[ \langle \text{seq alignment} \rangle \equiv \langle \text{seq alignment interface} \rangle \langle \text{seq alignment implementation} \rangle \]

This code is used in chunk 1.

3.5.1 seq_alignment interface

Here is the declaration of the seq_alignment class: it’s a template class in which the parameter \( T \) is the type of the costs: default value is float.

\[ \langle \text{seq alignment interface} \rangle \equiv \text{template } \langle \text{class } T=\text{float} \rangle \text{ class seq_alignment}{ \}

private:

\langle \text{private variables of seq_alignment} \rangle

\langle \text{private functions of seq_alignment} \rangle

public:

\langle \text{functions exported by seq_alignment} \rangle

\};

This code is used in chunk 25.
3.5 Sequence alignment class

Private variables of seq_alignment

The \texttt{seq\_alignment} class uses the \texttt{boost::multi\_array} class for storing the matrix needed for its calculations.

\begin{verbatim}
//the score matrix
const score_matrix<T> *c_matrix;
//gap creation and extension costs
T gap_cost, extend_gap_cost;

//the sequences
const vector<char> *A, *B;
//sequence lengths
int m, n;

//Matrix for quadratic-space alignments
boost::multi_array<T, 2> *N;
//Matrix for affine-gap alignments
boost::multi_array<T, 3> *M;
//Second vectors for linear-space alignments
boost::multi_array<T, 3> *M1;

//start and end of the alignment
int min_m, min_n, max_m, max_n; //initialized by constructors
//score of the alignment
T max_score;
//buffers for memorizing the alignment
vector<char> *line1, *line2, *line3;

//infinite negative for type T
T NEG_INFTY;
\end{verbatim}

This code is used in chunk 26.

Private functions of seq_alignment

These three private functions make the actual calculations needed to build the alignments.

\begin{verbatim}
void build_alignment(int i, int j);
void build_affine_alignment(int i, int j);
void build_linear_alignment(int i1, int j1, int i2, int j2,
                           T A_begin_gap_cost, T A_end_gap_cost,
                           T B_begin_gap_cost, T B_end_gap_cost);
\end{verbatim}

This code is used in chunk 26.
3.5 Sequence alignment class

Functions exported by seq\_alignment

(functions exported by seq\_alignment 29)≡

//constructors
seq\_alignment(const score\_matrix<T>& scores,
const vector<char>& fst, const vector<char>& snd);
seq\_alignment(const score\_matrix<T>& scores,
string fst, string snd);
seq\_alignment(const score\_matrix<T>& scores,
alibio::bio\_string fst, alibio::bio\_string snd);

//destructor
~seq\_alignment();

//prints the last calculated alignment
void print\_alignment();

//classic alignment functions
//global - Needleman-Wunsch
void nw\_align();
//local - Smiths-Waterman
void sw\_align();

//alignment functions using affine gap penalties
//global
void nw\_affine\_align();
//local
void sw\_affine\_align();
//semiglobal
void semiglobal\_affine\_align();

//alignment functions using linear space - Myers-Miller
//global
void nw\_linear\_align();
//local
void sw\_linear\_align();
//semiglobal
void semiglobal\_linear\_align();

This code is used in chunk 26.
3.5 Sequence alignment class

3.5.2 seq.alignment implementation

The implementations of the declared functions

\[
\{\text{seq.alignment implementation } 30\} \equiv \\
\{\text{seq.alignment constructors } 31\} \\
\{\text{seq.alignment destructor } 33\}
\]

\[
\{\text{print.alignment } 34\}
\]

\[
\{\text{nw.align } 35\} \\
\{\text{sw.align } 36\} \\
\{\text{build.alignment } 37\}
\]

\[
\{\text{nw.affine.align } 38\} \\
\{\text{sw.affine.align } 39\} \\
\{\text{semiglobal.affine.align } 40\} \\
\{\text{build.affine.alignment } 42\}
\]

\[
\{\text{nw.linear.align } 44\} \\
\{\text{sw.linear.align } 45\} \\
\{\text{semiglobal.linear.align } 46\} \\
\{\text{build.linear.alignment } 43\}
\]

This code is used in chunk 25.
Constructors implementation

The three constructors take a score matrix and two sequences and instantiate the needed memory. The seq_alignment T parameter must be of the same type of the corresponding cost_matrix T parameter, the compiler takes care of point out eventual misuses.

The constructors differs from the type of the sequences passed as parameters, respectively const vector<char>&, string and alibio::bio_string:

\[
\text{seq\_alignment\_constructors } 31
\]

```
template <class T>
seq_alignment<T>::seq_alignment(const score_matrix<T>& scores,
                        const vector<char>& fst, const vector<char>& snd){
    try{
        A=new vector<char>(fst);
        B=new vector<char>(snd);
    }
    catch(const bad_alloc& x){
        cerr << "Out of memory in seq_alignment constructor: "<< x.what()<<"\n";
        abort();
    }
}
```  

```
//end seq_alignment(score_matrix, vector<char>, vector<char>)
```

```
template <class T>
seq_alignment<T>::seq_alignment(const score_matrix<T>& scores,
                        string fst, string snd){
    try{
        A=new vector<char>(fst.begin(),fst.end());
        B=new vector<char>(snd.begin(),snd.end());
    }
    catch(const bad_alloc& x){
        cerr << "Out of memory in seq_alignment constructor: "<<x.what()<<"\n";
        abort();
    }
}
```  

```
//end seq_alignment(score_matrix, string, string)
```

```
template <class T>
seq_alignment<T>::seq_alignment(const score_matrix<T>& scores,
                        alibio::bio_string fst, alibio::bio_string snd){
    try{
        A=new vector<char>(fst.get_sequence().begin(),fst.get_sequence().end());
        B=new vector<char>(snd.get_sequence().begin(),snd.get_sequence().end());
    }
    catch(const bad_alloc& x){
        cerr << "Out of memory in seq_alignment constructor: "<<x.what()<<"\n";
        abort();
    }
}
```
3.5 Sequence alignment class

⟨common initializations 32⟩
} // end seq_alignment(score_matrix, bio_string, bio_string)
This code is used in chunk 30.

All of the constructors need to initialize the same local variables, so we use a single code chunk

⟨common initializations 32⟩≡
c_matrix=&scores;
gap_cost=c_matrix->get_gap_cost();
extend_gap_cost=c_matrix->get_extend_gap_cost();
NEG_INFTY=-numeric_limits<T>::max()/2;

m=A->size(); n=B->size();

//reinizialize min_m min_n max_m max_n max_score
min_m=0; max_m=m; min_n=0; max_n=n;
max_score=NEG_INFTY;

try{
    line1=new vector<char>();
    line2=new vector<char>();
    line3=new vector<char>();
} catch(const bad_alloc x){
    cerr << "Out of memory:" << x.what() << "\n";
    abort();
}

This code is used in chunk 31.

Destructor implementation

⟨seq_alignment destructor 33⟩≡
template <class T>
seq_alignment<T>::~seq_alignment(){
    delete A;
    delete B;
    delete line1;
    delete line2;
    delete line3;
} // ~seq_alignment()

This code is used in chunk 30.
3.5 Sequence alignment class

Printing the alignment

This function prints on the standard output the calculated alignment and its score.

\[
\text{print_alignment} \equiv \\
\text{template } \langle \text{class } T \rangle \\
\text{void seq_alignment}<T>::\text{print_alignment}() \{ \\
\text{vector<char>::iterator It;}
\text{cout }\ll \text{min}_m + 1 \ll \"\t\"; \\
\text{for (It=line1->begin(); It!=line1->end(); It++)} \\
\text{cout }\ll \text{(*It);} \\
\text{cout }\ll \"\t\" \ll \text{max}_m \ll \"\n\"; \\
\text{cout }\ll \"\t\"; \\
\text{for (It=line2->begin(); It!=line2->end(); It++)} \\
\text{cout }\ll \text{(*It);} \\
\text{cout }\ll \"\n\"; \\
\text{cout }\ll \text{min}_n + 1 \ll \"\t\"; \\
\text{for (It=line3->begin(); It!=line3->end(); It++)} \\
\text{cout }\ll \text{(*It);} \\
\text{cout }\ll \"\t\" \ll \text{max}_n \ll \"\n\"; \\
\text{cout }\ll \"\text{Alignment total score:}\" \ll \text{max_score} \ll \"\n\"; \\
\} \text{//end print_alignment()} \\
\] (debug_random 75)

This code is used in chunk 30.
3.5 Sequence alignment class

3.5.3 Classic algorithms implementation

Needleman-Wunsch algorithm implementation

\[
\text{nw_align(T)} \equiv
\begin{align*}
\text{template <class T> void seq_alignment<T>::nw_align()}{ \\
\text{// allocating matrix space} \\
\text{try}{ \\
\text{N= new boost::multi_array< T, 2>(boost::extents[m+1][n+1]);} \\
\text{catch(const bad_alloc &x){} \\
\text{cerr << "Out of memory in nw_align(): " << x.what() << "\n"; abort();}} \\
\text{// reinizialize min_m min_n max_m max_n max_score} \\
\text{min_m=0; max_m=m; min_n=0; max_n=n; max_score=NEG_INFTY;} \\
\text{(*N)[0][0]=(T)0;} \\
\text{for (int j=1;j<=n;j++){} \\
\text{(*N)[0][j]=-j*gap_cost;} \\
\text{for (int i=0;i<m;i++){} \\
\text{int next_i=i+1;} \\
\text{(*N)[next_i][0]=(*N)[i][0]-gap_cost;} \\
\text{for (int j=0;j<n;j++){} \\
\text{int next_j=j+1;} \\
\text{(*N)[next_i][next_j]=max(max((*N)[i][j]+c_matrix->get_score((*A)[i], (*B)[j]), \\
\text{(*N)[i][next_j]-gap_cost)}, \\
\text{(*N)[next_i][j]-gap_cost));} \\
\text{max_score=(*N)[m][n];} \\
\text{// cleaning an eventual precalculated alignment} \\
\text{(*line1).clear();} \\
\text{(*line2).clear();} \\
\text{(*line3).clear();} \\
\text{build_alignment(max_m, max_n);} \\
\text{(memory_line 82)} \\
\text{delete N;} \\
\text{// end nw_align()}
\end{align*}
\]

This code is used in chunk 30.
Smith-Waterman algorithm implementation

```cpp
template <class T>
void seq_alignment<T>::sw_align()
{
    // allocating matrix space
    try{
        N=new boost::multi_array< T, 2>(boost::extents[m+1][n+1]);
    }
    catch(const bad_alloc & x){
        cerr << "Out of memory in sw_align(): " << x.what() << ":\n";
        abort();
    }
    // reinizialize min_m min_n max_m max_n max_score
    min_m=0; max_m=m; min_n=0; max_n=n;
    max_score=0;
    for (int i=0;i<=m;i++)
        (*N)[i][0]=(T)0;
    for (int j=0;j<=n;j++)
        (*N)[0][j]=(T)0;
    for (int i=0;i<m;i++)
    {
        int next_i=i+1;
        for (int j=0;j<n;j++)
        {
            int next_j=j+1;
            (*N)[next_i][next_j]=max(max(max(*N)[i][j]+c_matrix->get_score((*A)[i],
                                    (*B)[j]),
                                    (*N)[next_i][j]-gap_cost),
                                    (*N)[next_i][j]-gap_cost),
                                    (T)0);
            if((*N)[next_i][next_j]>(*N)[max_m][max_n]){
                max_m=next_i;
                max_n=next_j;
            }
        }
    }
    max_score=(*N)[max_m][max_n];
    // cleaning an eventual precalculated alignment
    (*line1).clear();
    (*line2).clear();
    (*line3).clear();
    if (max_score>0)
        build_alignment(max_m, max_n);
    else
        max_score=0;
}
```
delete N;
} // end sw_align()

This code is used in chunk 30.

Tracking back the alignment

\begin{align*}
\buildalign(37) & \equiv \\
\text{template <class T>}
\text{void seq_alignment<T>::build_alignment(int i, int j)\{}
\text{int prev_i=i-1;}
\text{int prev_j=j-1;}
\text{if((i>0) \&\& (j>0) \&\&}
\text{((N)[i][j] == (N)[prev_i][prev_j] + c_matrix->get_score(A[prev_i],}
\text{B[prev_j])))\{}
\buildalignment(prev_i, prev_j);
\line1->push_back(A[prev_i]);
\line2->push_back('|');
\line3->push_back(B[prev_j]);
\return;
\text{\}}
\text{if((i>0) \&\& ((N)[i][j] == (N)[prev_i][j] - gap_cost)) }
\text{\{}
\buildalignment(prev_i, j);
\line1->push_back(A[prev_i]);
\line2->push_back(' ');
\line3->push_back('-');
\return;
\text{\}}
\text{if((j>0) \&\& ((N)[i][j] == (N)[i][prev_j] - gap_cost)) }
\text{\{}
\buildalignment(i, prev_j);
\line1->push_back('-');
\line2->push_back('-');
\line3->push_back(B[prev_j]);
\return;
\text{\}}
\text{min_m=i;}
\text{min_n=j;}
\} // end build_alignment(int, int)
\end{align*}

This code is used in chunk 30.
Global alignment implementation

```cpp
template <class T>
void seq_alignment<T>::nw_affine_align()
{
    // allocating matrix space
    try{
        M=new boost::multi_array< T, 3>(boost::extents[m+1][n+1][4]);
    }
    catch(const bad_alloc& x){
        cerr << "Out of memory in nw_affine_align(): " << x.what() << "\n";
        abort();
    }
    // reinizialize min_m min_n max_m max_n max_score
    min_m=0; max_m=m; min_n=0; max_n=n;
    max_score=NEG_INFTY;

    // preparing first line of the 4 matrix
    (*M)[0][0][0]=(T)0;
    (*M)[0][0][1]=NEG_INFTY;
    (*M)[0][0][2]=NEG_INFTY;
    (*M)[0][0][3]=(T)3;//STOP HERE
    for (int j=0;j<n;j++){
        int next_j=j+1;
        (*M)[0][next_j][0]=-(gap_cost+j*extend_gap_cost);
        (*M)[0][next_j][1]=NEG_INFTY;
        (*M)[0][next_j][2]=-(gap_cost+j*extend_gap_cost);
        (*M)[0][next_j][3]=(T)2;//LEFT
    }

    for (int i=0;i<m;i++){
        int next_i=i+1;
        // preparing first column of the 4 matrix
        (*M)[next_i][0][0]=-(gap_cost+i*extend_gap_cost);
        (*M)[next_i][0][1]=-(gap_cost+i*extend_gap_cost);
        (*M)[next_i][0][2]=NEG_INFTY;
        (*M)[next_i][0][3]=(T)1;//UP
        for (int j=0;j<n;j++){
            int next_j=j+1;
            // affine elaboration 41
        }
    }
    max_score=(*M)[m][n][0];

    // cleaning an eventual precalculated alignment
```
This code is used in chunk 30.
3.5 Sequence alignment class

Local alignment implementation

(sw_affine_align 39)≡

template <class T>
void seq_alignment<T>::sw_affine_align()
{

    //allocating matrix space
    try{
        M=new boost::multi_array<T, 3>(boost::extents[m+1][n+1][4]);
    }
    catch(const bad_alloc& x){
        cerr << "Out of memory in sw_affine_align(): " << x.what() << "\n"
            abort();
    }

    //reinizialize min_m min_n max_m max_n max_score
    min_m=0; max_m=m; min_n=0; max_n=n;
    max_score=0;

    //preparing first row of the 4 matrix
    (*M)[0][0][0]=(T)0;
    (*M)[0][0][1]=NEG_INFTY;
    (*M)[0][0][2]=NEG_INFTY;
    (*M)[0][0][3]=(T)3;//STOP HERE

    for (int j=0;j<n;j++){
        int next_j=j+1;
        (*M)[0][next_j][0]=(T)0;
        (*M)[0][next_j][1]=NEG_INFTY;
        (*M)[0][next_j][2]=-(gap_cost+j*extend_gap_cost);
        (*M)[0][next_j][3]=(T)3;//STOP HERE
    }

    for (int i=0;i<m;i++){
        int next_i=i+1;

        //preparing first column of the 4 matrix
        (*M)[next_i][0][0]=(T)0;
        (*M)[next_i][0][1]=-(gap_cost+i*extend_gap_cost);
        (*M)[next_i][0][2]=NEG_INFTY;
        (*M)[next_i][0][3]=(T)3;//STOP HERE

        for (int j=0;j<n;j++){
            int next_j=j+1;

            if((T)0>(*M)[next_i][next_j][0]){
                (*M)[next_i][next_j][0]=(T)0;
                (*M)[next_i][next_j][3]=(T)3;//STOP HERE
            }

            if((*M)[next_i][next_j][0]>max_score){
                max_score=(*M)[next_i][next_j][0];
            }
        }
    }
max_m=next_i;
max_n=next_j;
}
}

// cleaning an eventual precalculated alignment
(*line1).clear();
(*line2).clear();
(*line3).clear();
if (max_score>0)
  build_affine_alignment(max_m, max_n);
else
  max_score=0;

(delete M;
} // end sw_affine_align()

This code is used in chunk 30.
Semiglobal alignment implementation

\[ \langle \text{semiglobal\_affine\_align} \; 40 \rangle \equiv \]

\begin{verbatim}
template <class T>
void seq_alignment<T>::semiglobal_affine_align(){
   //allocating matrix space
   try{
      M=new boost::multi_array< T, 3>(boost::extents[m+1][n+1][4]);
   } catch(const bad_alloc& x){
      cerr <<"Out of memory in semiglobal_affine_align(): " <<x.what() << \\
           "\n";
      abort();
   }
   //reinizialize min_m min_n max_m max_n max_score
   min_m=0; max_m=m; min_n=0; max_n=n;
   max_score=0;

   //preparing first row of the 4 matrix
   (*M)[0][0][0]=(T)0;
   (*M)[0][0][1]=NEG_INFTY;
   (*M)[0][0][2]=NEG_INFTY;
   (*M)[0][0][3]=(T)3;
   for (int j=0;j<n;j++){//STOP HERE
      int next_j=j+1;
      (*M)[0][next_j][0]=(T)0;
      (*M)[0][next_j][1]=NEG_INFTY;
      (*M)[0][next_j][2]=-(gap_cost+j*extend_gap_cost);
      (*M)[0][next_j][3]=(T)3;
   }

   for (int i=0;i<m;i++){//STOP HERE
      int next_i=i+1;
      //preparing first column of the 4 matrix
      (*M)[next_i][0][0]=(T)0;
      (*M)[next_i][0][1]=-(gap_cost+i*extend_gap_cost);
      (*M)[next_i][0][2]=NEG_INFTY;
      (*M)[next_i][0][3]=(T)3;
      for (int j=0;j<n;j++){//STOP HERE
         int next_j=j+1;
         //affine elaboration 41
      }
   }
   for (int i=1;i<m;i++)
      if(*M[i][n][0]>max_score){
         max_score=*M[i][n][0];
         max_m=i;
         max_n=n;
      }
\end{verbatim}
for (int j=1;j<=n;j++)
    if ((*M)[m][j][0]>max_score){
        max_score=(*M)[m][j][0];
        max_m=m;
        max_n=j;
    }

    //cleaning an eventual precalculated alignment
    (*line1).clear();
    (*line2).clear();
    (*line3).clear();
    if (max_score>0)
        build_affine_alignment(max_m, max_n);
    else
        max_score=0;

    (memory_line 82)
    delete M;
} //end semiglobal_affine_align()

This code is used in chunk 30.

Matrix elaboration

≡
(*M)[next_i][next_j][0]=(*M)[i][j][0]+c_matrix->get_score((*A)[i],(*B)[j]);
(*M)[next_i][next_j][1]=max((*M)[i][next_j][0]-gap_cost,
                          (*M)[i][next_j][1]-extend_gap_cost);
(*M)[next_i][next_j][2]=max((*M)[next_i][j][0]-gap_cost,
                          (*M)[next_i][j][2]-extend_gap_cost);
(*M)[next_i][next_j][3]=(T)0;//upleft
    if ((*M)[next_i][next_j][1]>=(*M)[next_i][next_j][0] &&
        (*M)[next_i][next_j][1]>(*M)[next_i][next_j][2]){ 
        (*M)[next_i][next_j][0]=(*M)[next_i][next_j][1];
        (*M)[next_i][next_j][3]=(T)1;//up
    } else if ((*M)[next_i][next_j][2]>(*M)[next_i][next_j][0] &&
        (*M)[next_i][next_j][2]>(*M)[next_i][next_j][1]){ 
        (*M)[next_i][next_j][0]=(*M)[next_i][next_j][2];
        (*M)[next_i][next_j][3]=(T)2;//left
}
3.5 Sequence alignment class

Tracking back the alignment

(build_affine_alignment 42) ≡

```cpp
template <class T>
void seq_alignment<T>::build_affine_alignment(int i, int j) {

    if (i>0 && j>0 && (*M)[i][j][3]==(T)0) {
        build_affine_alignment(i-1, j-1);
        line1->push_back((*A)[i-1]);
        if((*A)[i-1]==(*B)[j-1]) // matching
            line2->push_back('|');
        else
            line2->push_back('X'); // mismatch
        line3->push_back((*B)[j-1]);
        return;
    }

    if (i>0 && (*M)[i][j][3]==(T)1) {// it's the beginning of a gap in A
        int k=i;
        while (i>0 && (*M)[i-1][j][1]==(*M)[i][j][1]+extend_gap_cost) {
            i--;//measuring the gap
        }

        if (i>0)
            build_affine_alignment(i-1, j);

        while (i<=k){
            line1->push_back((*A)[i-1]);
            line2->push_back(' ');
            line3->push_back('-');
            i++;
        }

        return;
    }

    if (j>0 && (*M)[i][j][3]==(T)2) {// it's the beginning of a gap in B
        int k=j;
        while (j>0 && (*M)[i][j-1][2]==(*M)[i][j][2]+extend_gap_cost) {
            j--;//measuring the gap
        }

        if (j>0)
            build_affine_alignment(i, j-1);

        while (j<=k){
            line1->push_back('-');
            line2->push_back(' ');
            line3->push_back((*B)[j-1]);
            j++;
        }
    }
}
```
return;
}
if((**M)[i][j][3]==(T)3){
    min_m=i;
    min_n=j;
}
}//end build_affine_alignment(int, int)

This code is used in chunk 30.
3.5 Sequence alignment class

3.5.5 Aligning in linear space

Building the alignment using linear space

\[
\begin{align*}
\langle \text{build_linear_alignment} \rangle \equiv \\
\text{template } <\text{class } T> \ \\
\text{void seq_alignment<T>::build_linear_alignment(int i1, int j1, int i2, int j2,} \\
\text{T A_begin_gap_cost, T A_end_gap_cost,} \\
\text{T B_begin_gap_cost, T B_end_gap_cost)\{} \\
\end{align*}
\]

assert(i2 >= i1 &amp; j2 >= j1);

//best score of the subproblem
T best_score=NEG_INFTY;

//solving the base of the recursion
if (j1==j2){ //only gap in A
    for (int i=i1;i<i2;i++) //gap in the second sequence
        line1->push_back((*A)[i]);
    line2->push_back(' ');
    line3->push_back('-');
}

if((i1==0) &amp;&amp; (j1==0) &amp;&amp; (i2==m) &amp;&amp; (j2==n))
    max_score=-(gap_cost+(m-1)*extend_gap_cost);

return;
}

if (i1==i2){ //only gap in B
    for (int j=j1;j<j2;j++) //gap in the second sequence
        line1->push_back('-');
    line2->push_back(' ');
    line3->push_back((*B)[j]);
}

if((i1==0) &amp;&amp; (j1==0) &amp;&amp; (i2==m) &amp;&amp; (j2==n))
    max_score=-(gap_cost+(n-1)*extend_gap_cost);

return;
}

int mid=(i1+i2)/2; //mid line

//computing maximum path scores form (i1,j1) to (mid,*)
//preparing first line of the 4 matrix (gap in B)
(*M)[i1%2][j1][0]=(T)0;
(*M)[i1%2][j1][1]=NEG_INFTY;
(*M)[i1%2][j1][2]=NEG_INFTY;
(*M)[i1%2][j1][3]=(T)0;

for (int j=j1;j<j2;j++)
    int next_j=j+1;
3.5 Sequence alignment class

(*M)[i1%2][next_j][0]=-(A_begin_gap_cost+(j-j1)*extend_gap_cost);
(*M)[i1%2][next_j][1]=NEG_INFTY;
(*M)[i1%2][next_j][2]=-(A_begin_gap_cost+(j-j1)*extend_gap_cost);
(*M)[i1%2][next_j][3]=(T)2; //left
}

for (int i=i1; i<=mid; i++){

    //precalc i+1 assuring to use only 2 lines
    int next_i=(i+1)%2;
    //calculating the first column (gap in B)
    (*M)[next_i][j1][0]=-(B_begin_gap_cost+(i-i1)*extend_gap_cost);
    (*M)[next_i][j1][1]=-(B_begin_gap_cost+(i-i1)*extend_gap_cost);
    (*M)[next_i][j1][2]=NEG_INFTY;
    (*M)[next_i][j1][3]=(T)1; //up

    for (int j=j1; j<j2; j++){
        int next_j=j+1;
        (*M)[next_i][next_j][0]=(*M)[i%2][j][0]+
c_matrix->get_score(*A)[i],(*B)[j]);
        (*M)[next_i][next_j][1]=max((*M)[i%2][next_j][0]-gap_cost,
        (*M)[i%2][next_j][1]-extend_gap_cost);
        (*M)[next_i][next_j][2]=max((*M)[next_i][j][0]-gap_cost,
        (*M)[next_i][j][2]-extend_gap_cost);
        (*M)[next_i][next_j][3]=(T)0; //upleft

        if((*M)[next_i][next_j][1]>=(*M)[next_i][next_j][0] &&
        (*M)[next_i][next_j][1]>=(*M)[next_i][next_j][2]){
            (*M)[next_i][next_j][0]=(*M)[next_i][next_j][1];
            (*M)[next_i][next_j][3]=(T)1; //up
        } else if((*M)[next_i][next_j][2]>=(*M)[next_i][next_j][0] &&
        (*M)[next_i][next_j][2]>(*M)[next_i][next_j][1]){
            (*M)[next_i][next_j][0]=(*M)[next_i][next_j][2];
            (*M)[next_i][next_j][3]=(T)2; //left
        } else {

        }
    }
    //computing maximum path scores from (i2,j2) down to (mid,*)
    //preparing first line of the 4 matrix (gap in B)
    (*M1)[i2%2][j2][0]=(T)0;
    (*M1)[i2%2][j2][1]=NEG_INFTY;
    (*M1)[i2%2][j2][2]=NEG_INFTY;
    (*M1)[i2%2][j2][3]=(T)0; //STOP HERE

    for (int j=j2; j>j1; j--){
        int prev_j=j-1;
        (*M1)[i2%2][prev_j][0]=-(*A_end_gap_cost+(j2-j)*extend_gap_cost);
        (*M1)[i2%2][prev_j][1]=NEG_INFTY;
    }
for (int i=i2; i>mid; i--){
    // precalc i-1 assuring to use only 2 lines
    int prev_i=(i-1)%2;

    // calculating the first column (gap in B)
    (*M1)[prev_i][j2][0]=-(B_end_gap_cost+(i2-i)*extend_gap_cost);
    (*M1)[prev_i][j2][1]=-(B_start_gap_cost+(i2-i)*extend_gap_cost);
    (*M1)[prev_i][j2][2] = NEG_INFTY;
    (*M1)[prev_i][j2][3]=(T)1; // down

    for (int j=j2; j>j1; j--){
        int prev_j=j-1;
        (*M1)[prev_i][prev_j][0] = (*M1)[i%2][j][0] +
                                    c_matrix->get_score((*A)[i-1],(*B)[prev_j]);
        (*M1)[prev_i][prev_j][1] = max((*M1)[i%2][j][1] - gap_cost,
                                        (*M1)[i%2][j][1] - extend_gap_cost);
        (*M1)[prev_i][prev_j][2] = max((*M1)[prev_i][j][0] - gap_cost,
                                        (*M1)[prev_i][j][2] - extend_gap_cost);

        (*M1)[prev_i][prev_j][3] = T); // downright

        if((*M1)[prev_i][prev_j][1]>=(*M1)[prev_i][prev_j][0] &&
          (*M1)[prev_i][prev_j][1]>(*M1)[prev_i][prev_j][2]){
            (*M1)[prev_i][prev_j][0] = (*M1)[prev_i][prev_j][1];
            (*M1)[prev_i][prev_j][3] = T); // down
        }else if ((*M1)[prev_i][prev_j][2]>=(*M1)[prev_i][prev_j][0] &&
          (*M1)[prev_i][prev_j][2]>(*M1)[prev_i][prev_j][1]){
            (*M1)[prev_i][prev_j][0] = (*M1)[prev_i][prev_j][2];
            (*M1)[prev_i][prev_j][3] = T); // right
        }
    }
}

// finding a midpoint of the global alignment

// position of the best score
int best_score_pos=j1;

for (int j=j1; j<=j2; j++){
    T sum=((*M)[mid%2][j][0]+(*M)[mid%2][j][0]);

    // if 2 gaps are in the same direction we must reduce the penalty
    sum=max(sum,
            (*M)[mid%2][j][1]+(*M)[mid%2][j][1]+gap_cost-extend_gap_cost);
    sum=max(sum,
            (*M)[mid%2][j][2]+(*M)[mid%2][j][2]+gap_cost-extend_gap_cost);

    // find the best j
if(sum>best_score){
    best_score_pos=j;
    best_score=sum;
}
}

// in the first recursion we archive the best global score in max_score
if((i1==0) && (j1==0) && (i2==m) && (j2==n) && (best_score>max_score))
    max_score=best_score;

// only one char left on seq A
if(i2-i1==1){
    if ((*M)[i2%2][j2][3]==(T)0){
        build_linear_alignment(i1, j1, i2-1, j2-1,
                        A_begin_gap_cost,A_end_gap_cost,
                        B_begin_gap_cost,B_end_gap_cost);
        line1->push_back((*A)[i2-1]);
        if((*(A))[i2-1]==*(B)[j2-1]) // matching
            line2->push_back('|');
        else
            line2->push_back('X'); // mismatch
        line3->push_back((*B)[j2-1]);
        return;
    }
    if ((*M)[i2%2][j2][3]==(T)1){ // print the char in A
        build_linear_alignment(i1, j1, i2-1, j2,
                        A_begin_gap_cost,A_end_gap_cost,
                        B_begin_gap_cost,B_end_gap_cost);
        line1->push_back((*(A))[i2-1]);
        line2->push_back(' ');
        line3->push_back('-');
        return;
    }
    if ((*M)[i2%2][j2][3]==(T)2){ // it's the beginning of a gap in A
        int k=j2;
        while(j2>j1 && (*M)[i2%2][j2-1][2]==(*M)[i2%2][j2][2]
            +extend_gap_cost){
            j2--; // measuring the gap
        }
        build_linear_alignment(i1, j1, i2, j2-1,
                        A_begin_gap_cost,A_end_gap_cost,
                        B_begin_gap_cost,B_end_gap_cost);
        while (j2<=k){
            line1->push_back('-');
            line2->push_back(' ');
            line3->push_back((*B)[j2-1]);
        }
    }
}
if (j2>j1)
    build_linear_alignment(i1, j1, i2, j2-1,
                        A_begin_gap_cost,A_end_gap_cost,
                        B_begin_gap_cost,B_end_gap_cost);
while (j2<=k){
    line1->push_back('-');
    line2->push_back(' ');
    line3->push_back((*B)[j2-1]);
}
3.5 Sequence alignment class

```c
    j2++;
    }

    return;
    }

    if(m==1)// A had only 1 char
        max_score=(*M)[i2%2][j2][0];

    return;
}

//operate on the 2 halves
//if the pivot is in the middle of a long gap we divide in 3 parts
//long gap in B
if (best_score==(*M)[mid%2][best_score_pos][1]+
    (*M1)[mid%2][best_score_pos][1]+
gap_cost-extend_gap_cost){

    build_linear_alignment(i1,j1,mid-1,best_score_pos,
        A_begin_gap_cost,A_end_gap_cost,
        B_begin_gap_cost,extend_gap_cost);

    line1->push_back((*A)[mid-1]);
    line2->push_back(' ');
    line3->push_back('-');
    line1->push_back((*A)[mid]);
    line2->push_back(' ');
    line3->push_back('-');

    build_linear_alignment(mid+1,best_score_pos,i2,j2,
        A_begin_gap_cost,extend_gap_cost,
        B_begin_gap_cost,B_end_gap_cost);

    return;
}
//long gap in A
if (best_score==(*M)[mid%2][best_score_pos][2]+
    (*M1)[mid%2][best_score_pos][2]+
gap_cost-extend_gap_cost){

    build_linear_alignment(i1,j1,mid,best_score_pos-1,
        A_begin_gap_cost,extend_gap_cost,A_end_gap_cost,
        B_begin_gap_cost,B_end_gap_cost);

    line1->push_back('-');
    line2->push_back(' ');
    line3->push_back((*B)[best_score_pos-1]);
    line1->push_back('-');
    line2->push_back(' ');
    line3->push_back((*B)[best_score_pos]);

    build_linear_alignment(mid,best_score_pos+1,i2,j2,
        A_begin_gap_cost,extend_gap_cost,B_begin_gap_cost,B_end_gap_cost);

    return;
}
```
This code is used in chunk 30.

Global alignment

\( \langle \text{nw-linear-align} 44 \rangle \equiv \)

```cpp
template <class T>
void seq_alignment<T>::nw_linear_align()
{
    // Allocating needed space
    try{
        M=new boost::multi_array< T, 3>(boost::extents[2][n+1][4]);
        M1=new boost::multi_array< T, 3>(boost::extents[2][n+1][4]);
    }
    catch(const bad_alloc x){
        cerr << "Out of memory in nw_linear_align(): " << x.what() << "\n";
        abort();
    }
    // Reinitalize min_m min_n max_m max_n max_score
    min_m=0; max_m=m; min_n=0; max_n=n;
    max_score=NEG_INFTY;

    // Cleaning an eventual precalculated alignment
    (*line1).clear();
    (*line2).clear();
    (*line3).clear();

    if (m>0||n>0)
        build_linear_alignment(0,0,m,n,gap_cost,gap_cost,gap_cost,gap_cost);
    else max_score=0; // Both sequences empty
}
```

This code is used in chunk 30.
Local alignment

\[ (sw\_linear\_align\ 45) \equiv \]

\[
\text{template <class T>}
\text{void seq_alignment<T>::sw_linear_align()}
\text{
// Allocating needed space}
\text{try}
\text{
M=new boost::multi_array< T, 3>(boost::extents[2][n+1][4]);}
\text{
M1=new boost::multi_array< T, 3>(boost::extents[2][n+1][4]);}
\text{catch(const bad_alloc x){}
\text{
cerr << "Out of memory in sw_linear_align(): " << x.what() << "\n";
\text{abort();}
\text{}}
\text{// reinizialize min_m min_n max_m max_n max_score}
\text{min_m=0; max_m=m; min_n=0; max_n=n;}
\text{max_score=0;}
\text{// determining max_m max_n with an upward cost only alignment}
\text{// preparing first line of the 4 matrix}
\text{(*M)[0][0][0]=(T)0;}
\text{(*M)[0][0][1]=NEG\_INFTY;}
\text{(*M)[0][0][2]=NEG\_INFTY;}
\text{for (int j=0;j<n;j++){
\text{int next_j=j+1;
\text{(*M)[0][next_j][0]=(T)0;}
\text{(*M)[0][next_j][1]=NEG\_INFTY;}
\text{(*M)[0][next_j][2]=-(gap_cost+j*extend_gap_cost);}
\text{}}
\text{for (int i=0; i<m; i++){
\text{int next_i=(i+1)%2;}
\text{// calculating the first column}
\text{(*M)[next_i][0][0]=(T)0;}
\text{(*M)[next_i][0][1]=-(gap_cost+i*extend_gap_cost);}
\text{(*M)[next_i][0][2]=NEG\_INFTY;}
\text{for (int j=0;j<n;j++){
\text{int next_j=j+1;}
\text{(*M)[next_i][next_j][0]=(*M)[i%2][j][0]+}
\text{\hspace{1cm}c_matrix->get_score((*A)[i],(*B)[j]);}
\text{(*M)[next_i][next_j][1]=max((*M)[i%2][next_j][0]-gap_cost,}
\text{\hspace{1cm}(*M)[i%2][next_j][1]-extend_gap_cost);}
\text{(*M)[next_i][next_j][2]=max((*M)[next_i][j][0]-gap_cost,}
\text{\hspace{1cm}(*M)[next_i][j][2]-extend_gap_cost);}
\text{if(((*M)[next_i][next_j][0]>(*M)[next_i][next_j][1])
\text{&& (*M)[next_i][next_j][0]>(*M)[next_i][next_j][2]){
\text{}}
\text{}}
\text{}}
\text{}}\]
3.5 Sequence alignment class

```c
} else if(*M[next_i][next_j][1]>(*M[next_i][next_j][2]){
(*M)[next_i][next_j][0]=(*M)[next_i][next_j][1];
} else {
(*M)[next_i][next_j][0]=(*M)[next_i][next_j][2];
}

if(0>(*M)[next_i][next_j][0]){(*M)[next_i][next_j][0]=(T)0;
}

if(*M)[next_i][next_j][0]>max_score){
    max_m=i+1;
    max_n=next_j;
    max_score=(*M)[next_i][next_j][0];
}
}

//determining min_m min_n with a backward cost only alignment
//preparing first line of the 4 matrix
(*M1)[max_m%2][max_n][0]=(T)0;
(*M1)[max_m%2][max_n][1]=NEG_INFTY;
(*M1)[max_m%2][max_n][2]=NEG_INFTY;

for (int j=max_n;j>0;j--){
    int prev_j=j-1;
    (*M1)[max_m%2][prev_j][0]=(T)0;
    (*M1)[max_m%2][prev_j][1]=NEG_INFTY;
    (*M1)[max_m%2][prev_j][2]=-(gap_cost+(max_n-j)*extend_gap_cost);
}

for (int i=max_m; i>0; i--){
    //precalc i+1
    int prev_i=(i-1)%2;

    //calculating the first column
    (*M1)[prev_i][max_n][0]=(T)0;
    (*M1)[prev_i][max_n][1]=-(gap_cost+(max_m-i)*extend_gap_cost);
    (*M1)[prev_i][max_n][2]=NEG_INFTY;

    for (int j=max_n;j>0;j--){
        int prev_j=j-1;
        (*M1)[prev_i][prev_j][0]=(*M1)[i%2][j][0]+c_matrix->get_score(*A[i-1],*B[prev_j]);
        (*M1)[prev_i][prev_j][1]=max(*M1)[i%2][prev_j][0]-gap_cost,
                        (*M1)[i%2][prev_j][1]-extend_gap_cost);
        (*M1)[prev_i][prev_j][2]=max(*M1)[prev_i][j][0]-gap_cost,
                        (*M1)[prev_i][j][2]-extend_gap_cost);

        if(*M1)[prev_i][prev_j][0]>(*M1)[prev_i][prev_j][1] &&
             (*M1)[prev_i][prev_j][0]>(*M1)[prev_i][prev_j][2]){
```
3.5 Sequence alignment class

```c
} else if((*M1)[prev_i][prev_j][1]>(*M1)[prev_i][prev_j][2]){
    (*M1)[prev_i][prev_j][0]=(*M1)[prev_i][prev_j][1];
} else {
    (*M1)[prev_i][prev_j][0]=(*M1)[prev_i][prev_j][2];
}

if((*M1)[prev_i][prev_j][0]==max_score){
    min_m=i-1;
    min_n=prev_j;
}
}

//cleaning an eventual precalculated alignment
(*line1).clear();
(*line2).clear();
(*line3).clear();
//if first string empty
if (max_score>0)
    build_linear_alignment(min_m,min_n,max_m,max_n,
        gap_cost,gap_cost,gap_cost,gap_cost);
else
    max_score=(T)0;

(memory_line 82)

delete M;
delete M1;
}//end sw_linear_align_align()
```

This code is used in chunk 30.
Semiglobal alignment

\[ \text{(semiglobal\_linear\_align 46)≡} \]

```cpp
template <class T>
void seq_alignment<T>::semiglobal_linear_align()
{
    // Allocating needed space
    try{
        M=new boost::multi_array< T, 3>(boost::extents[2][n+1][4]);
        M1=new boost::multi_array< T, 3>(boost::extents[2][n+1][4]);
    } catch(const bad_alloc& x){
        cerr<< "Out of memory in semiglobal_linear_align(): "<< x.what()<<"\n";
        abort();
    }

    // reinitialize min_m min_n max_m max_n max_score
    min_m=0; max_m=m; min_n=0; max_n=n;
    max_score=0;

    // deternining max_m max_n with an upward, cost only, alignment
    // preparing first line of the 4 matrix
    (*M)[0][0][0]=(T)0;
    (*M)[0][0][1]=NEG_INFTY;
    (*M)[0][0][2]=NEG_INFTY;

    for (int j=0; j<n; j++){
        int next_j=j+1;
        (*M)[0][next_j][0]=(T)0;
        (*M)[0][next_j][1]=NEG_INFTY;
        (*M)[0][next_j][2]=-(gap_cost+j*extend_gap_cost);
    }

    for (int i=0; i<m; i++){
        // precalc i+1
        int next_i=(i+1)%2;

        // calculating the first column
        (*M)[next_i][0][0]=(T)0;
        (*M)[next_i][0][1]=-(gap_cost+i*extend_gap_cost);
        (*M)[next_i][0][2]=NEG_INFTY;

        for (int j=0; j<n; j++){
            int next_j=j+1;
            (*M)[next_i][next_j][0]=(*(M)[i%2][j][0]+
                                      c_matrix->get_score((*A)[i],(*B)[j]));
            (*M)[next_i][next_j][1]=max((*(M)[i%2][next_j][0]-gap_cost,
                                         *(M)[i%2][next_j][1]-extend_gap_cost);
            (*M)[next_i][next_j][2]=max((*(M)[next_i][j][0]-gap_cost,
                                         *(M)[next_i][j][2]-extend_gap_cost);

            if((*(M)[next_i][next_j][1])>(*(M)[next_i][next_j][0] &
                                   *(M)[next_i][next_j][1])>(*(M)[next_i][next_j][2]){
```
3.5 Sequence alignment class

\[
(*M)[next_i][next_j][0] = (*M)[next_i][next_j][1];
\]

\[
\text{else if}((*M)[next_i][next_j][2] >= (*M)[next_i][next_j][0] \&\&
(*M)[next_i][next_j][2] > (*M)[next_i][next_j][1])
\]

\[
(*M)[next_i][next_j][0] = (*M)[next_i][next_j][2];
\]

if((i+1==m || next_j==n) \&\& (*M)[next_i][next_j][0] > max_score)
\[
\begin{align*}
max_m &= i+1; \\
max_n &= next_j; \\
max_score &= (*M)[next_i][next_j][0];
\end{align*}
\]

\]

//determining min_m min_n with a backward, cost only, alignment

//preparing first line of the 4 matrix
\[
(*M_1)[max_m%2][max_n][0] = (T)0; \\
(*M_1)[max_m%2][max_n][1] = NEG_INFTY; \\
(*M_1)[max_m%2][max_n][2] = NEG_INFTY;
\]

for (int j=max_n; j>0; j--)
\[
\begin{align*}
\text{int prev_j} &= j-1; \\
(*M_1)[max_m%2][prev_j][0] &= -(gap_cost + (max_n-j)*extend_gap_cost); \\
(*M_1)[max_m%2][prev_j][1] &= NEG_INFTY; \\
(*M_1)[max_m%2][prev_j][2] &= -(gap_cost + (max_n-j)*extend_gap_cost);
\end{align*}
\]

for (int i=max_m; i>0; i--)
\[
\begin{align*}
\text{int prev_i} &= (i-1)%2; \\
\text{//precalc i+1}
\end{align*}
\]

\]

//calculating the first column
\[
(*M_1)[prev_i][max_n][0] = -(gap_cost + (max_m-i)*extend_gap_cost); \\
(*M_1)[prev_i][max_n][1] = -(gap_cost + (max_m-i)*extend_gap_cost); \\
(*M_1)[prev_i][max_n][2] = NEG_INFTY;
\]

for (int j=max_n; j>0; j--)
\[
\begin{align*}
\text{int prev_j} &= j-1; \\
(*M_1)[prev_i][prev_j][0] &= (*M_1)[i%2][j][0] + \\
\text{c_matrix->get_score}(*A[i-1], *B[prev_j]); \\
(*M_1)[prev_i][prev_j][1] &= max((*M_1)[i%2][prev_j][0] - gap_cost, \\
(*M_1)[i%2][prev_j][1] - extend_gap_cost); \\
(*M_1)[prev_i][prev_j][2] &= max((*M_1)[prev_i][j][0] - gap_cost, \\
(*M_1)[prev_i][j][2] - extend_gap_cost);
\end{align*}
\]

if((*M_1)[prev_i][prev_j][1] >= (*M_1)[prev_i][prev_j][0] \&\&
(*M_1)[prev_i][prev_j][1] >= (*M_1)[prev_i][prev_j][2])
\[
\begin{align*}
(*M_1)[prev_i][prev_j][0] &= (*M_1)[prev_i][prev_j][1];
\end{align*}
\]

} else if((*M_1)[prev_i][prev_j][2] >= (*M_1)[prev_i][prev_j][0] \&\&
(*M_1)[prev_i][prev_j][2] >= (*M_1)[prev_i][prev_j][1])
\[
\begin{align*}
(*M_1)[prev_i][prev_j][0] &= (*M_1)[prev_i][prev_j][2];
\end{align*}
\]
if((i-1==0 || prev_j==0 ) &&
    ((*M1)[prev_i][prev_j][0]==max_score)){
    min_m=i-1;
    min_n=prev_j;
    break; //the first found is OK
}
}

//cleaning an eventual precalculated alignment
(*line1).clear();
(*line2).clear();
(*line3).clear();

if (max_score>0)
    build_linear_alignment(min_m,min_n,max_m,max_n,
                           gap_cost,gap_cost,gap_cost,gap_cost);
else
    max_score=(T)0;

(memory.line 82)

delete M;
delete M1;
}//end semiglobal_linear_align()

This code is used in chunk 30.
Chapter 4

Tests

4.1 Regression tests

In order to test the current and the possible future versions of the produced libraries, we adopt the technique of ”automate regression testing”, which performs a sequence of tests to compare the expected behaviour of the classes with their real behaviour or to compare the new versions of our classes implementation with the old ones. This is done by comparing the output of some appropriate test programs executed on the latest versions of the classes, against the expected output or against the output of the same test programs executed on the old implementation of the classes.

The test requires the following tools:

- A test program (one for each class), which contains a main function implementing tests for all the features of the tested class;
- A behaviour file (one for each class), which contains the tests inputs and the corresponding expected outputs. The inputs provided are supposed to cover all the possible situations (even the unexpected ones) that can rise using the class;
- An output file (one for each class), which contains the actual outputs produced by the test program executed on the latest version of the class;
- A script program that executes the test programs, performs the comparison between the expected and the actual outputs, and reports the possible differences;

Then, the programmer must make sure that the behaviour of the classes hasn’t changed, except in expected ways.
4.1 Regression tests

4.1.1 Structure of the test program

Description of the main function

The test program contains a main function that implements all the suitable tests. Each test is identified by a number code (starting from 1), and can be executed independently from others.

The main function reads a code as parameter and selects the corresponding test. Then it reads the requested test inputs from standard input, executes the test and prints the results on standard output. If incorrect parameters or input values are found, the program reports an error and, if it’s impossible to continue, ends the test and returns the 0 value.

Description of the variables

The main function uses two variables:

- **line**, of **string** data type, used to store each line read from the standard input.
- **code**, of **int** type, used to store the codes of the tests to execute.

47

(main function for test: head 47)≡

```c
#define SUCCESS 0
#define CODE_ERROR 1
#define INPUT_ERROR 2
#define SYNTAX_ERROR 3

int main (int argc, char *argv[]){
    int code;

    //check the correctness of the parameters
    if(argc<2) {
        std::cout << "Usage: " << argv[0] << " TEST_CODE\n";
        return SYNTAX_ERROR;
    }
    code=atoi(argv[1]);
    if(code==0) {
        std::cout << "Error while testing: incorrect code.\n";
        return CODE_ERROR;
    }
}
```

This code is used in chunks 49 and 61.

48

(main function for test: tail 48)≡

```c
    return SUCCESS;
}
```

This code is used in chunks 49 and 61.
4.1 Regression tests

4.1.2 Structure of the test file

The test file must respect the following syntax:

- the first line of each test must begin with the special sequence "====="; everything after that is ignored and can, therefore, be viewed as a comment;
- after that, a test identifier (a numerical code) is expected;
- the third line must begin with the "***** sequence"; everything after that is ignored and can, therefore, be viewed as a comment;
- in the following lines, input data must be specified (each input on a different line);
- after all the input data have been entered, the program expects a line beginning with the "***** sequence"; everything after that is ignored and can, therefore, be viewed as a comment;
- in the following lines, the expected outputs must be specified;
- after all the expected outputs have been entered, either another test or the end of the file is expected.
4.1 Regression tests

Test file example

=====set_score(char, char, T) - 2 chars 1 T=====  
3
*****
G
C
5
*****
5
=====set_match_score(T) - 1 T=====  
4
*****
3
*****

A  C  G  T  
A  3 -1 -1 -1
C  -1  3 -1 -1
G  -1 -1  3 -1
T  -1 -1 -1  3

gap cost=1

gap extension cost=1

4.1.3 Test program for score_matrix class

This program implements all the suitable tests for the ⟨score_matrix⟩ class.

#include <iostream>
#include <stdlib.h>
#include <alibio/two_sequence.hpp>

switch(code) {
    case 1:
        (testing score_matrix(alibio::alphabet) constructor 51)
        break;
    case 2:
        (testing set_score(char, char, T) function 52)
        break;
    case 3:
        (testing get_score(char, char) function 53)
        break;
    case 4:
        (testing set_match_score(T) function 54)
        break;
    case 5:
4.1 Regression tests

```cpp
    (testing set_mismatch_score(T) function 55)
    break;
    case 6:
        (testing set_gap_cost(T) function 56)
        break;
    case 7:
        (testing set_gap_cost(T,T) function 57)
        break;
    case 8:
        (testing get_gap_cost() function 58)
        break;
    case 9:
        (testing get_extend_gap_cost() function 59)
        break;
    case 10:
        (testing print() function 60)
        break;
    default:
        std::cout << "Error while testing: code '" << code << "' not found";
    }
    (main function for test: tail 48)
```

Root chunk (not used in this document).

**Preparing score_matrix object**

50  (preparing score_matrix object 50)≡
    //declaring an alphabet for DNA
    alibio::alphabet dna("DNA");

    //adding symbols to the alphabet
    dna.add_symbol(alibio::symbol('A','Adenine'));
    dna.add_symbol(alibio::symbol('C','Cytosine'));
    dna.add_symbol(alibio::symbol('T','Thymine'));
    dna.add_symbol(alibio::symbol('G','Guanine'));

    //creating the score matrix
    alibio::score_matrix<float> myscores(dna);

This code is used in chunks 51–60 and 62–65.

**Testing score_matrix constructor**

The test creates an alibio::alphabet object and uses it for instantiating a (score_matrix 8) object

51  (testing score_matrix(alibio::alphabet) constructor 51)≡
    {
        (preparing score_matrix object 50)
    }

This code is used in chunk 49.
### 4.1 Regression tests

**Testing set_score function**

The test reads from the input the size of the distance matrix that will be created using the generic constructor `score_matrix(int dim)` and prints it. Then reads a second size and resize the object using `resize(int dim)`. Finally prints the resized object on the standard output using `print()` function.

```cpp
(testing set_score(char, char, T) function 52) ≡

{  
  //preparing score_matrix object 50

  //reading 2 chars from stdin
  std::string line1;
  std::string line2;

  //reading 1 float from stdin
  std::string line3;
  std::getline(std::cin,line1,\n');
  std::getline(std::cin,line2,\n');
  std::getline(std::cin,line3,\n');
  myscores.set_score(line1[0], line2[0], atof(line3.c_str()));
  cout << myscores.get_score(line1[0], line2[0]);
}
```

This code is used in chunk 49.

**Testing get_score function**

The test creates an `alibio::alphabet` and a `score_matrix 8` object, it reads from the standard input two characters (one per line) and prints the score of switching between the first to the second getting the value from the `score_matrix 8` object. If one or both of the chars are not listed in the alphabet, the test prints an error message.

```cpp
(testing get_score(char, char) function 53) ≡

{  
  //preparing score_matrix object 50

  //reading 2 chars from stdin
  std::string line1;
  std::string line2;
  float score;

  std::getline(std::cin,line1,\n');
  std::getline(std::cin,line2,\n');
  score=myscores.get_score(line1[0], line2[0]);
  cout << score;
}
```

This code is used in chunk 49.
Testing \texttt{set\_match\_score} function

The test creates a \texttt{score\_matrix} object
\par \texttt{(testing set\_match\_score(T) function)} \equiv
\par \{ \par \texttt{(preparing score\_matrix object)} \par //reading 1 double from stdin \par std\:\:\:\:string line1; \par std\:\:\:getline\:\:\\:\:cin, line1, '\n'\}; \par myscores.set_match_score(atof(line1.c_str())); \par myscores.print(); \par \} \par This code is used in chunk 49.

Testing \texttt{set\_mismatch\_score} function

The test creates a \texttt{score\_matrix} object
\par \texttt{(testing set\_mismatch\_score(T) function)} \equiv
\par \{ \par \texttt{(preparing score\_matrix object)} \par //reading 1 double from stdin \par std\:\:\:string line1; \par std\:\:\:getline\:\:\\:\:cin, line1, '\n'\}; \par myscores.set_mismatch_score(atof(line1.c_str())); \par myscores.print(); \par \} \par This code is used in chunk 49.

Testing \texttt{set\_gap\_cost} function

The test creates a \texttt{score\_matrix} object.
\par \texttt{(testing set\_gap\_cost(T) function)} \equiv
\par \{ \par \texttt{(preparing score\_matrix object)} \par //reading 1 double from stdin \par std\:\:\:string line1; \par std\:\:\:getline\:\:\\:\:cin, line1, '\n'\}; \par myscores.set_gap_cost(atof(line1.c_str())); \par cout << myscores.get_gap_cost() << endl; \par cout << myscores.get_extend_gap_cost(); \par \} \par This code is used in chunk 49.
4.1 Regression tests

Testing \texttt{get\_gap\_cost} function

The test creates a \textit{score\_matrix} object.

Testing \texttt{get\_extend\_gap\_cost} function

The test creates a \textit{score\_matrix} object.

Testing \texttt{print} function

The test creates a \textit{score\_matrix} object.
4.1 Regression tests

4.1.4 Test program for seq.alignment class

This program implements the tests for the ⟨seq.alignment⟩ class.

```
#include <iostream>
#include <stdlib.h>
#include <alibio/two_sequence.hpp>

(main function for test: head 47)
switch(code) {
    case 1:
        (testing seq.alignment vector constructor 62)
        break;
    case 2:
        (testing seq.alignment string constructor 63)
        break;
    case 3:
        (testing seq.alignment bio_string constructor 64)
        break;
    case 4:
        (testing nw_align() function 66)
        break;
    case 5:
        (testing sw_align() function 67)
        break;
    case 6:
        (testing nw_affine_align() function 68)
        break;
    case 7:
        (testing sw_affine_align() function 69)
        break;
    case 8:
        (testing semiglobal_affine_align() function 70)
        break;
    case 9:
        (testing nw_linear_align() function 71)
        break;
    case 10:
        (testing sw_linear_align() function 72)
        break;
    case 11:
        (testing semiglobal_linear_align() function 73)
        break;
    case 12:
        (testing nw_affine_align() with a different alphabet 74)
        break;
    default:
        std::cout << "\nError while testing: code '" << code << "' not found";
}
(main function for test: tail 48)
```
Testing `seq_alignment` constructors

This test verifies the `(seq_alignment vector constructor 25)` vector constructor using the DNA alphabet and the default score matrix.

```cpp
62  (testing seq_alignment vector constructor 62)≡
{  
   (preparing score_matrix object 50)

   //reading 2 strings from stdin
   std::string line1;
   std::string line2;

   std::getline(std::cin,line1,‘\n’);
   std::getline(std::cin,line2,‘\n’);

   vector<char> A(line1.begin(),line2.end());
   vector<char> B(line1.begin(),line2.end());

   alibio::seq_alignment<float> my_alignment(myscores, A, B);
}
```

This code is used in chunk 61.

This test verifies the `(seq_alignment string constructor 25)` string constructor using the DNA alphabet and the default score matrix.

```cpp
63  (testing seq_alignment string constructor 63)≡
{  
   (preparing score_matrix object 50)

   //reading 2 strings from stdin
   std::string line1;
   std::string line2;

   std::getline(std::cin,line1,‘\n’);
   std::getline(std::cin,line2,‘\n’);

   //creating the alignment matrix
   alibio::seq_alignment<float> my_alignment(myscores, line1, line2);
}
```

This code is used in chunk 61.
This test verifies the `<seq_alignment 25> bio_string constructor using the DNA alphabet and the default score matrix
(testing seq_alignment bio_string constructor 64)>`

```
{                         
  (preparing score_matrix object 50)  

  // declaring 2 sequences with "bio_string" class as primary structure and
  // "empty" class as secondary and tertiary structures
  alibio::sequence<alibio::bio_string,alibio::empty,alibio::empty> seq_one;
  alibio::sequence<alibio::bio_string,alibio::empty,alibio::empty> seq_two;

  // setting the alphabet (working on "seq_one.primary")
  seq_one.primary.set_alphabet(dna);
  seq_two.primary.set_alphabet(dna);

  // reading 2 strings from stdin
  std::string line1;
  std::string line2;
  std::getline(std::cin,line1,'\n');
  std::getline(std::cin,line2,'\n');

  // setting the sequence (working on "seq_one.primary")
  seq_one.primary.set_sequence(line1);

  // setting the sequence (working on "seq_two.primary")
  seq_two.primary.set_sequence(line2);

  // creating the alignment matrix
  alibio::seq_alignment<float> my_alignment(myscores,
                                         seq_one.primary, seq_two.primary);
}
```

This code is used in chunk 61.
4.1 Regression tests

Preparing seq_alignment object

In this section we prepare a \{seq_alignment 25\} object that will be used for all the alignment tests

\(\langle\text{preparing seq_alignment object 65}\rangle\equiv\)

\(\langle\text{preparing score_matrix object 50}\rangle\)

//modifying score matrix
myscores.set_match_score(5);
myscores.set_mismatch_score(-4);
myscores.set_gap_cost(10, 0.5);

//reading 2 strings from stdin
std::string line1;
std::string line2;
std::getline(std::cin,line1,'\n');
std::getline(std::cin,line2,'\n');

//creating the alignment object
alibio::seq_alignment<float> my_alignment(myscores, line1, line2);

This code is used in chunks 66–73.

Testing nw_align function

\(\langle\text{testing nw_align()} function 66\rangle\equiv\)

\{\n\langle\text{preparing seq_alignment object 65}\rangle\nmy_alignment.nw_align();
my_alignment.print_alignment();\n\}

This code is used in chunk 61.

Testing sw_align function

\(\langle\text{testing sw_align()} function 67\rangle\equiv\)

\{\n\langle\text{preparing seq_alignment object 65}\rangle\nmy_alignment.sw_align();
my_alignment.print_alignment();\n\}

This code is used in chunk 61.
4.1 Regression tests

Testing \texttt{nw\_affine\_align} function

\begin{verbatim}
(\texttt{testing nw\_affine\_align}() function 68) ≡
    \begin{verbatim}
    \langle \text{preparing seq\_alignment object 65} \rangle
    my_alignment.nw_affine_align();
    my_alignment.print_alignment();
    \end{verbatim}
\end{verbatim}
This code is used in chunk 61.

Testing \texttt{sw\_affine\_align} function

\begin{verbatim}
(\texttt{testing sw\_affine\_align}() function 69) ≡
    \begin{verbatim}
    \langle \text{preparing seq\_alignment object 65} \rangle
    my_alignment.sw_affine_align();
    my_alignment.print_alignment();
    \end{verbatim}
\end{verbatim}
This code is used in chunk 61.

Testing \texttt{semiglobal\_affine\_align} function

\begin{verbatim}
(\texttt{testing semiglobal\_affine\_align}() function 70) ≡
    \begin{verbatim}
    \langle \text{preparing seq\_alignment object 65} \rangle
    my_alignment.semiglobal_affine_align();
    my_alignment.print_alignment();
    \end{verbatim}
\end{verbatim}
This code is used in chunk 61.

Testing \texttt{nw\_linear\_align} function

\begin{verbatim}
(\texttt{testing nw\_linear\_align}() function 71) ≡
    \begin{verbatim}
    \langle \text{preparing seq\_alignment object 65} \rangle
    my_alignment.nw_linear_align();
    my_alignment.print_alignment();
    \end{verbatim}
\end{verbatim}
This code is used in chunk 61.
4.1 Regression tests

Testing `sw_linear_align` function

\[
\text{(testing `sw_linear_align()` function 72) } \equiv \\
\{ \\
\text{(preparing seq.alignment object 65)} \\
\text{my_alignment.sw_linear_align();} \\
\text{my_alignment.print_alignment();} \\
\} \\
\]
This code is used in chunk 61.

Testing `semiglobal_linear_align` function

\[
\text{(testing `semiglobal_linear_align()` function 73) } \equiv \\
\{ \\
\text{(preparing seq.alignment object 65)} \\
\text{my_alignment.semiglobal_linear_align();} \\
\text{my_alignment.print_alignment();} \\
\} \\
\]
This code is used in chunk 61.
4.1 Regression tests

Testing *nw_affine_align* with a different alphabet

This test runs the *nw_affine_align* using a different alphabet

(testing *nw_affine_align()* with a different alphabet)

```cpp
{ //declaring an alphabet for DNA
  alibio::alphabet prot("Protein");

  //adding symbols to the alphabet
  prot.add_symbol(alibio::symbol('A'));
  prot.add_symbol(alibio::symbol('R'));
  prot.add_symbol(alibio::symbol('N'));
  prot.add_symbol(alibio::symbol('D'));
  prot.add_symbol(alibio::symbol('C'));
  prot.add_symbol(alibio::symbol('Q'));
  prot.add_symbol(alibio::symbol('E'));
  prot.add_symbol(alibio::symbol('G'));
  prot.add_symbol(alibio::symbol('H'));
  prot.add_symbol(alibio::symbol('I'));
  prot.add_symbol(alibio::symbol('L'));
  prot.add_symbol(alibio::symbol('K'));
  prot.add_symbol(alibio::symbol('M'));
  prot.add_symbol(alibio::symbol('F'));
  prot.add_symbol(alibio::symbol('P'));
  prot.add_symbol(alibio::symbol('S'));
  prot.add_symbol(alibio::symbol('T'));
  prot.add_symbol(alibio::symbol('W'));
  prot.add_symbol(alibio::symbol('Y'));
  prot.add_symbol(alibio::symbol('V'));
  prot.add_symbol(alibio::symbol('B'));
  prot.add_symbol(alibio::symbol('Z'));
  prot.add_symbol(alibio::symbol('X'));

  //creating the score matrix for "prot" alphabet
  alibio::score_matrix<float> prot_scores(prot);

  //reading 2 strings from stdin
  std::string line1;
  std::string line2;
  //reading match score from stdin
  std::string line3;
  //reading gap cost from stdin
  std::string line4;
  //reading extending gap cost from stdin
  std::string line5;
  //reading mismatch score from stdin
  std::string line6;

  std::getline(std::cin,line1,'\n');
  std::getline(std::cin,line2,'\n');
```
4.1 Regression tests

```cpp
std::getline(std::cin,line3,'\n');
std::getline(std::cin,line4,'\n');
std::getline(std::cin,line5,'\n');
std::getline(std::cin,line6,'\n');

//modifying score matrix
prot_scores.set_match_score(atof(line3.c_str()));
prot_scores.set_gap_cost(atof(line4.c_str()),atof(line5.c_str()));
prot_scores.set_mismatch_score(atof(line6.c_str()));

//creating the alignment matrix
alibio::seq_alignment<float> prot_alignment(prot_scores, line1, line2);

//printing alignment
prot_alignment.nw_affine_align();
prot_alignment.print_alignment();
}
```

This code is used in chunk 61.
4.2 Memory leaks test

Because of the great amount of operations over arrays that are used in our library we thought
that there was great probabilities of memory leaks in our software so we decided to test it using
a Free Software (cfr.2.1) project called Valgrind that includes a powerful memory analyzer.

Using this software is very easy: we just had to launch our example program \textit{two_sequence_example.cpp} with the command

\$ valgrind --leak-check=yes ./two_sequence_example.bin

4.3 Random tests

We decided to test our functions against random generated sequences hoping to detect errors in
algorithms implementation that didn't appear in our previous tests

We wrote code for calculating alignment scores by printed alignment, this way we could detect
inconsistencies between them and find critical sequences for further tests. Decommenting the
following debugging code we obtain additional output that will be used in \textit{random_test.cpp}.

The code reads the alignment memorized in \texttt{line1}, \texttt{line2} and \texttt{line3} and in the first part calculates the alignment score supposing we did an affine gap penalty (cfr.2.1) alignment, while in the second part it does the same for linear gap penalty (cfr.2.1) alignments.

\begin{verbatim}
75 //decomment for random test
/*
  T true_score=0;
  It=line2->begin();
  for (int i=0;It!=line2->end();i++){
      if (((*line2)[i]==' '){//gap
          if (((*line1)[i]=='-'){//gap in A
              if (i>0 & & ((*line1)[i-1]=='-'))
                  true_score-=extend_gap_cost;
          } else {//gap in B
              if (i>0 & & ((*line3)[i-1]=='-'))
                  true_score-=extend_gap_cost;
          }
      } else true_score+=c_matrix->get_score((*line1)[i],(*line3)[i]);
      It++;
  }

  T lin_score=0;
  It=line2->begin();
  for (int i=0;It!=line2->end();i++){
      if (((*line2)[i]==' ')){//gap
          if (((*line1)[i]=='-'){//gap in A
              if (i>0 & & ((*line1)[i-1]=='-'))
                  true_score-=extend_gap_cost;
          } else {//gap in B
              if (i>0 & & ((*line3)[i-1]=='-'))
                  true_score-=extend_gap_cost;
          }
      } else true_score+=c_matrix->get_score((*line1)[i],(*line3)[i]);
      It++;
  }
\end{verbatim}
lin_score-=gap_cost;
} else {//gap in B
    lin_score-=gap_cost;
}
} else lin_score+=c_matrix->get_score((*line1)[i],(*line3)[i]);
    It++;
}

std::cout<<"True alignment score:"<<true_score<<" score:"<<lin_score << "\n";
//*/
This code is used in chunk 34.
4.3 Random tests

The actual program generates random sequences of random length but not longer than the value in `max_seq_length`. It creates them upon a specified alphabet whose length is given in `alph_size` variable, then it calls all the alignment functions.

The whole procedure is repeated as many times as indicated in `repetitions` variable.

```cpp
#include <iostream>
#include <stdlib.h>
#include <alibio/two_sequence.hpp>

int main(){
    int alph_size=4;
    int repetitions=2000;
    int max_seq_length=30;

    //Getting internal clock time for random
    srand(time(NULL));
    //declaring an alphabet
    alibio::alphabet my_alph("my alphabet");
    //adding symbols to the alphabet
    for (int i=0; i<alph_size; i++){
        my_alph.add_symbol(alibio::symbol('A'+i));
    }
    alibio::score_matrix<> myscores(my_alph);
    //changing scores
    myscores.set_match_score(5);
    myscores.set_mismatch_score(-4);
    myscores.set_gap_cost(10, 0.5);

    for (int n=0;n<repetitions;n++){
        vector<char> fst, snd;
        int n=rand() % max_seq_length;
        int m=rand() % max_seq_length;
        fst.reserve(n);
        snd.reserve(m);

        //generating strings randomly
        for (int i=0; i<n; i++){
            fst.push_back('A'+(rand() % alph_size));
            cout << fst[i];
        } cout << endl;
        for (int j=0; j<m; j++){
            snd.push_back('A'+(rand() % alph_size));
            cout << snd[j];
        } cout << endl;
        //creating the alignment matrix
        alibio::seq_alignment<> my_alignment(myscores, fst, snd);

        //do the alignments
        cout << "Optimal global alignment (linear gap penalty)\n";
    }
}
```
my_alignment.nw_align();
my_alignment.print_alignment();

cout << "Optimal local alignment (linear gap penalty)\n";
my_alignment.sw_align();
my_alignment.print_alignment();

cout << "Optimal global alignment (affine gap penalty)\n";
my_alignment.nw_affine_align();
my_alignment.print_alignment();

cout << "Optimal local alignment (affine gap penalty)\n";
my_alignment.sw_affine_align();
my_alignment.print_alignment();

cout << "Optimal semiglobal alignment (affine gap penalty)\n";
my_alignment.semiglobal_affine_align();
my_alignment.print_alignment();

cout << "Optimal global alignment (affine gap penalty-linear space)\n";
my_alignment.nw_linear_align();
my_alignment.print_alignment();

cout << "Optimal local alignment (affine gap penalty-linear space)\n";
my_alignment.sw_linear_align();
my_alignment.print_alignment();

cout << "Optimal semiglobal alignment (affine gap penalty-linear space)\n";
my_alignment.semiglobal_linear_align();
my_alignment.print_alignment();
}
return 0;
}
4.3 Random tests

To analyze the huge output of our test program we wrote a GAWK script that verifies not only the correctness of reported alignment scores but also that affine alignments gave the same results of the corresponding linear-space ones. The script needs only score values to work, so we need to filter the `random_test.cpp` output with the UNIX command `grep` and then give it as input to the script `random_test.awk`. We did it with the following command assuming we compiled `random_test.cpp` as `random_test.bin`

```
$ ./random_test.bin | grep score | ./random_test.awk
```

The script writes a line for each error found, if no errors are found nothing is printed.

```
#!/usr/bin/awk -f
BEGIN {}
{
    //for each alignment we memorize the alignment score in the variable "old"
    if ((NR%16)%2==1) old=$3;
    if ((NR%16)%2==0){
        //in case of linear gap penalty we compare with the corresponding value
        if (NR%16==2 || NR%16==4){
            if ($4!=old) print old "<-->" $4 " ERROR " NR%16 "\n";
        }
        //in case of affine gap penalty we do the same
        else if($3!=old) print old "<-->" $3 " ERROR " NR%16 "\n";
    }
    else if (NR%16==5) var5=$3;//nw_affine_align score
    else if (NR%16==7) var7=$3;//sw_affine_align score
    else if (NR%16==9) var9=$3;//semiglobal_affine_align score
    else if (NR%16==11) { //comparing nw_affine and nw_linear
        if ($3!=var5) print var5 "<-->" $3 " GLOBAL\n";
    } else if (NR%16==13) { //comparing sw_affine and sw_linear
        if ($3!=var7) print var7 "<-->" $3 " LOCAL\n";
    } else if (NR%16==15) { //comparing semiglobal_affine and semiglobal_linear
        if ($3!=var9) print var9 "<-->" $3 " SEMIGLOBAL\n";
    }
}
END{}
```

This test permitted to discover a number of subtle software bugs that couldn’t be found with the previous tests.
Chapter 5

Examples

The following example program can be used as a quick tutorial on how the alignment functions can be used in conjunction with other ALiBio parts: let’s start declaring an `alibio::alphabet` and adding symbols to it:

```cpp
#include <iostream>
#include <stdlib.h>
#include <alibio/two_sequence.hpp>

int main(){
    // declaring an alphabet for DNA
    alibio::alphabet dna("DNA");

    // adding symbols to the alphabet
    dna.add_symbol(alibio::symbol('a', "Adenine"));
    dna.add_symbol(alibio::symbol('c', "Cytosine"));
    dna.add_symbol(alibio::symbol('t', "Thymine"));
    dna.add_symbol(alibio::symbol('g', "Guanine"));

    This definition is continued in chunks 79–81.
    Root chunk (not used in this document).
```
Then we create two `alibio::sequence` objects and assign to their primary structures the `alibio::alphabet` just created and two strings based on it:

```cpp
//declaring 2 sequences with "bio_string" class as primary structure and
//"empty" class as secondary and tertiary structures
alibio::sequence<alibio::bio_string,alibio::empty,alibio::empty> seq_one;
alibio::sequence<alibio::bio_string,alibio::empty,alibio::empty> seq_two;

//setting the alphabet (working on "seq_one.primary")
seq_one.primary.set_alphabet(dna);
seq_two.primary.set_alphabet(dna);

//setting the sequence (working on "seq_one.primary")
seq_one.primary.set_sequence("ATTAGCATTAGACT");

//setting the sequence (working on "seq_two.primary")
seq_two.primary.set_sequence("GCCGCTCTC");
```

We need to create a `(score_matrix 8)` object constructing it from the same alphabet, now we can modify the scores as we like: by category (match, mismatch, gap) or by specifying a specific char couple:

```cpp
alibio::score_matrix<float> myscores(dna);

//modifying the score_matrix
myscores.set_match_score(5);
myscores.set_mismatch_score(-4);
//modifying the score_matrix
myscores.set_gap_cost(5,0.5);
//modifying the score more selectively
myscores.set_cost('A','T',5);
//printing the score_matrix
myscores.print();

//creating alignment object
alibio::seq_alignment<float> my_alignment(myscores, seq_one.primary, seq_two.primary);
```
Finally we can call the alignment methods and print the generated alignments:

```cpp
cout << "Optimal global alignment (linear gap penalty)\n";
my_alignment.nw_align();
my_alignment.print_alignment();

cout << "Optimal local alignment (linear gap penalty)\n";
my_alignment.sw_align();
my_alignment.print_alignment();

cout << "Optimal global alignment (affine gap penalty)\n";
my_alignment.nw_affine_align();
my_alignment.print_alignment();

cout << "Optimal local alignment (affine gap penalty)\n";
my_alignment.sw_affine_align();
my_alignment.print_alignment();

cout << "Optimal semiglobal alignment (affine gap penalty)\n";
my_alignment.semiglobal_affine_align();
my_alignment.print_alignment();

cout << "Optimal global alignment (affine gap penalty - linear space)\n";
my_alignment.nw_linear_align();
my_alignment.print_alignment();

cout << "Optimal local alignment (affine gap penalty - linear space)\n";
my_alignment.sw_linear_align();
my_alignment.print_alignment();

cout << "Optimal semiglobal alignment (affine gap penalty - linear space)\n";
my_alignment.semiglobal_linear_align();
my_alignment.print_alignment();

return 0;
```
Chapter 6

Performance analysis

This chapter describes how to measure time and space performance of our algorithms, it also includes the scripts we used to produce the graphs presented in section 6.2.

6.1 Generating data

The following c++ program generates the files time.dat and space.dat which contain time and memory used by \textlangle nw\_align 35 \textrangle, \textlangle nw\_offline\_align 38 \textrangle and \textlangle nw\_linear\_align 44 \textrangle functions when operating over sequences of increasing length (from 0 to 3000 chars).

To obtain memory data, we need to decomment the following line which calls the UNIX command ps and transforms the result from KBytes to Bytes before writing it on the file space.dat.

The command supposes we compiled the \textlangle performance.cpp 83 \textrangle source file to performance.bin.

\begin{verbatim}
//decomment for memory tests
/*
system("echo '/bin/ps --no-header -o vsz -C performance.bin'*1024 |\n   bc -l >> space.dat");
//*/
\end{verbatim}

This code is used in chunks 35, 36, 38–40, and 44–46.
The program begins declaring some local variable whose meaning is explained in the code comments.

```cpp
#include <iostream>
#include <fstream>
#include <stdlib.h>
#include <time.h>
#include <alibio/two_sequence.hpp>

int main(){
  //number of symbols in the alphabet
  int alph_size=4;
  //max length of the sequences
  int seq_size=3000;
  //variables for storing time measures
  time_t start,end;
  double dif;
  //output buffer for writing time measures
  ofstream out_time;

  This definition is continued in chunks 84 and 85.
  Root chunk (not used in this document).

  Then the program generates an alphabet of `alph_size` size and deletes the files `time.dat` and `space.dat` if they exist. It also initialize the random number generator with a seed obtained by PC's clock.

```
The program then generates randomly the sequences and calls the alignment functions on them printing the used time on `time.dat` while the decommented `<memory_line 82>` prints memory usage on `space.dat`. Everything is repeated until sequence size equals `seq_size`.

```cpp
(83) //GLOBAL ALIGNMENT - LINEAR GAP PENALTY
time (&start);//reset timer
my_alignment.nw_align();//do the alignment
my_alignment.print_alignment();
//writing time elapsed in time.dat
time (&end);
dif = difftime (end,start);

out_time.open("time.dat", ofstream::out | ofstream::app);
out_time << dif << endl; out_time.close();

//GLOBAL ALIGNMENT - AFFINE GAP PENALTY
 time (&start);//reset timer
my_alignment.nw_affine_align();//do the alignment
my_alignment.print_alignment();
//writing time elapsed in time.dat
time (&end);
dif = difftime (end,start);

out_time.open("time.dat", ofstream::out | ofstream::app);
out_time << dif << endl; out_time.close();

//GLOBAL ALIGNMENT - AFFINE GAP PENALTY - LINEAR SPACE
 time (&start);//reset timer
my_alignment.nw_linear_align();//do the alignment
my_alignment.print_alignment();
//writing time elapsed in time.dat
time (&end);
```
6.1 Generating data

```c

dif = difftime (end, start);
out_time.open("time.dat", ofstream::out | ofstream::app);
out_time << dif << endl; out_time.close();
}
return 0;
}
```

The produced files are then formatted by the following GAWK script creating `gnuplot_time.dat` and `gnuplot_space.dat` with the UNIX commands

```bash
$ performance.awk time.dat >> gnuplot_time.dat
$ performance.awk space.dat >> gnuplot_space.dat
```

```
#!/usr/bin/awk -f
BEGIN {ORS=" "; nchar="0";}
{
    if (NR==1) base=$0;

    if (NR%3==1) {
       print "\n" nchar "\t" $0;
       nchar+=10;
    }
    else print "\t" $0;
}
END{print "\n" ;}
```

Root chunk (not used in this document).
6.1 Generating data

The data files obtained are then plotted with the following Gnuplot script.

```gnuplot
#!/usr/bin/gnuplot
set terminal postscript eps enhanced color
set xlabel "Input size (chars)"

set out 'gnuplot_time.eps'
#set xlabel "Input size (chars)"
set ylabel "Time (sec)"
set title "Time performance"
plot \\
'gnuplot_time.dat' using 1:2 title 'nw align' \ smooth csplines lt 1 linewidth 3, \\
'gnuplot_time.dat' using 1:3 title 'nw affine align' \ smooth csplines lt 2 linewidth 3, \\
'gnuplot_time.dat' using 1:4 title 'nw linear align' \ smooth csplines lt 3 linewidth 3

set out 'gnuplot_space.eps'
#set xlabel "Input size (chars)"
set ylabel "Space (Bytes)"
set format y "%.0s{/Symbol \327}10^{-%S}"
set title "Space performance"
#unset logscale
plot \\
'gnuplot_space.dat' using 1:2 title 'nw align' \ smooth csplines lt 1 linewidth 3, \\
'gnuplot_space.dat' using 1:3 title 'nw affine align' \ smooth csplines lt 2 linewidth 3, \\
'gnuplot_space.dat' using 1:4 title 'nw linear align' \ smooth csplines lt 3 linewidth 3

set logscale xy
set xtics (1000,1500,2000,2500,3000)

set out 'gnuplot_time_compl.eps'
#set xlabel "Input size (chars)"
set ylabel "Time (sec)"
set format y "10^{-%L}"
set title "Time complexity (log axes)"
plot [1000:] \\
'gnuplot_time.dat' using 1:2 title 'nw align' \ smooth csplines lt 1 linewidth 3, \\
'gnuplot_time.dat' using 1:3 title 'nw affine align' \ smooth csplines lt 2 linewidth 3, \\
'gnuplot_time.dat' using 1:4 title 'nw linear align' \ smooth csplines lt 3 linewidth 3, \\
(x**2)/100000 title 'quadratic' smooth csplines lt -1

set out 'gnuplot_space_compl.eps'
#set xlabel "Input size (chars)"
```

The Gnuplot script sets up the environment, defines the input size and output formats, plots the time and space performance, and then plots the time complexity with a log scale.
set ylabel "Space (Bytes)"
set xtics (1000,1500,2000,2500,3000)
set format y "%.0f/\text{Symbol \textbackslash 327}10^{-\{\%}\text{S}}"
set title "Space complexity (log axes)"
plot [1000:3000][:50000000] \
   'gnuplot_space.dat' using 1:2 title 'nw align'\n   smooth csplines lt 1 lw 3,\n   'gnuplot_space.dat' using 1:3 title 'nw affine align' \n   smooth csplines lt 2 lw 3, \n   x**2 title 'quadratic' smooth csplines lt -1

Root chunk (not used in this document).
The following graphs represent time and space performances of our implementations, from them we can obtain informations about which algorithms are faster or memory-hungrier.

We can already notice that the space complexity of \((nw\_linear\_align\ 44)\) is linear while the others are polynomial.
To have a visual indication of the polynomial functions rank we use logarithmic scale for both axes, now polynomials are straight lines whose slope depends on rank: if a function looks parallel to \( x^2 \) we can assure that its implementation is really quadratic. As expected, all the functions are quadratic-time, \( \langle nw\text{-}align 35 \rangle \) and \( \langle nw\text{-}affine\text{-}align 38 \rangle \) are also quadratic-space.
6.3 Performance comparison

In this section we compare the performances of our functions against the equivalent ones obtained from a well known project named EMBOSS[16] which is also Free Software (cfr.2.1). We used a Valgrind[17] tool called Massif which is a memory profiler that generates an easily understandable graph of memory usage and time computation for a given program.

In order to test the program over a realistic input we downloaded two uncorrelated nucleotide sequences from the European Bioinformatics website: embl:AF177870 and embl:L07770.

We also used a very common score matrix which corresponds to the EDNAFULL matrix in the EMBOSS package:

\[
\begin{array}{cccc}
A & T & G & C \\
A & 5 & -4 & -4 & -4 \\
T & -4 & 5 & -4 & -4 \\
G & -4 & -4 & 5 & -4 \\
C & -4 & -4 & -4 & 5 \\
\end{array}
\]

We tested the Affine gap penalties versions of the algorithms using 10 as gap opening cost and 1 as extending cost.

The programs

All of the programs share a common part that takes care of creating the needed \texttt{alibio::seq\_alignment} object:

\begin{verbatim}
  //declaring an alphabet for DNA
  alibio::alphabet dna("DNA");

  //adding symbols to the alphabet
  dna.add_symbol(alibio::symbol('a','Adenine'));
  dna.add_symbol(alibio::symbol('c','Cytosine'));
  dna.add_symbol(alibio::symbol('t','Thymine'));
  dna.add_symbol(alibio::symbol('g','Guanine'));

  //declaring 2 sequences with "bio\_string" class as primary structure and
  //"empty" class as secondary and tertiary structures
  alibio::sequence<alibio::bio_string,alibio::empty,alibio::empty> seq_one;
  alibio::sequence<alibio::bio_string,alibio::empty,alibio::empty> seq_two;

  //setting the alphabet (working on "seq\_one\_primary")
  seq_one.primary.set_alphabet(dna);
  seq_two.primary.set_alphabet(dna);

  //setting the sequence (working on "seq\_one\_primary")

\end{verbatim}
6.3 Performance comparison

```cpp
//setting the sequence (working on "seq_two.primary")
⟨embl:L07770 90⟩

alibio::score_matrix<int> myscores(dna);

//modifying the score_matrix
myscores.set_match_score(5);
myscores.set_mismatch_score(-4);
//modifying the score_matrix
myscores.set_gap_cost(10,1);

//creating alignment object
alibio::seq_alignment<int> my_alignment(myscores,
    seq_one.primary, seq_two.primary);
```

This code is used in chunks 91–93.

The lines of code that assign the downloaded sequences to the object

```cpp
⟨embl:AF177870 89⟩
seq_one.primary.set_sequence("gaacgcgaatgcctc ... aagagtttacag");
```

This code is used in chunk 88.

```cpp
⟨embl:L07770 90⟩
seq_two.primary.set_sequence("ggtagaacagcttca ... aaatttcttgcagt");
```

This code is used in chunk 88.

Here are the actual programs used for graph creation:

```cpp
⟨semiglobal_affine.cpp 91⟩
⟨licence 2⟩
#include <iostream>
#include <stdlib.h>
#include <alibio/two_sequence.hpp>

int main(){
    (object_preparing 88)
    cout << "Optimal semiglobal alignment (affine gap penalty)\n";
    my_alignment.semiglobal_affine_align();
    my_alignment.print_alignment();

    return 0;
}
```

Root chunk (not used in this document).
6.3 Performance comparison

#include <iostream>
#include <stdlib.h>
#include <alibio/two_sequence.hpp>

int main()
{
    //object_preparing 88
    cout << "Optimal local alignment (affine gap penalty)\n";
    my_alignment.sw_affine_align();
    my_alignment.print_alignment();
    return 0;
}

Root chunk (not used in this document).

#include <iostream>
#include <stdlib.h>
#include <alibio/two_sequence.hpp>

int main()
{
    //object_preparing 88
    cout << "Optimal global alignment (affine gap penalty - linear space)\n";
    my_alignment.nw_linear_align();
    my_alignment.print_alignment();
    return 0;
}

Root chunk (not used in this document).
6.3 Performance comparison

6.3.1 Results

The graphs we obtained represent time ($x$ axes) and space occupation ($y$ axes) of the tested functions. We used the following command line where the actual `PROGRAM` tested is indicated at the top of each graph.

```
$ valgrind --tool=massif PROGRAM
```

Semiglobal alignment

![Graph 1](image1.png)

ALiBio (**semiglobal_affine_align 40**) function:
- time = 14 sec
- space = 80 MB
- alignment score = 1102

![Graph 2](image2.png)

EMBOSS needle program:
- time = 22 sec
- space = 40 MB
- alignment score = 1102

While AliBio function is quite faster, EMBOSS program uses half space
6.3 Performance comparison

Local alignment

ALiBio (sw_offine_align 39) function:
time=35sec space=80MB alignment score=1113

EMBOSS water program:
time=30sec space=40MB alignment score=1113

While time performances are quite similar, EMBOSS program uses half space
6.3 Performance comparison

Linear-space global alignment

ALiBio (*nw_linear_align 44*) function:
- time = 35 sec
- space = 140 kB
- alignment score = 668

EMBOSS stretcher program:
- time = 9 sec
- space = 220 kB
- alignment score = 668

ALiBio function uses half space than EMBOSS program but is also 5 times slower.

Different color sections are associated with the different functions that actually reserve memory as reported in the caption.
Chapter 7

Conclusions

Once finished the stage period, the proposed objectives were reached. The requested algorithm implementation was produced, along with its documentation according to the project standard, also the implementation resource occupation is in line with the expected one.

The free software nature of the ALiBio project is an incentive for its improvement. In fact the bioinformatics software developers can freely use ALiBio inside their own programs, but also contribute with their own improvements and their expansions in developing the ALiBio libraries.

7.1 Possible future improvements

- On the performance side some improvements can be made to the given implementations, in particular the memory footprint can probably be furthermore reduced.

- Some modifications for better time performances could be added to the implemented algorithms, in particular a technique known as "Four Russians"\[7\] could reduce the time complexity by a factor of $\log^2 n$.

- In the past years other algorithms have been developed for the peerwise alignment problem, in particular to find sub-optimal solutions; their implementation could be added to the present library.

- The ALiBio library has been thought to be useful for developers of bioinformatics applications, anyway the presented alignment functions could be easily used by end users if a Command Line Interface were developed.

From that CLI, using pre-existing Free Software (cfr.2.1) like Pise[18] and Kaptain, a web interface and a graphical one can be obtained very quickly.
Chapter 8

Quick Methods reference

In this chapter we give a quick reference for all the methods implemented in the alignment library

8.1 score_matrix Class

score_matrix (const alibio::alphabet&) \rightarrow void
Description: \langle score_matrix constructor 14 \rangle creates a new object for the given alibio::alphabet
Precondition: -
Postcondition: - ~score_matrix (void) \rightarrow void
Description: \langle score_matrix destructor 15 \rangle deallocates memory used by the \langle score_matrix 8 \rangle
Precondition: -
Postcondition: - set_score (char;char;T) \rightarrow void
Description: \langle set_score 17 \rangle sets the score of mismatch between the two char to the given T value
Precondition: -
Postcondition: - get_score (char;char) \rightarrow T
Description: \langle get_score 18 \rangle returns the score of a mismatch between the two given char
Precondition: -
Postcondition: - set_match_score (T) \rightarrow void
Description: \langle set_match_score 19 \rangle sets the score for matching chars
Precondition: -
Postcondition: - set_mismatch_score (T) \rightarrow void
Description: \langle set_mismatch_score 20 \rangle sets the score for mismatching chars
Precondition: -
Postcondition: - set_gap_cost (T;T) \rightarrow void
Description: \langle set_gap_cost 21 \rangle sets the score for gap insertions
Precondition: -
Postcondition: - print (void) \rightarrow void
Description: \langle print 24 \rangle prints the score matrix
Precondition: -
Postcondition: -
8.2  seq_alignment Class

seq_alignment (const score_matrix&; const vector<char>&; const vector<char>&) \rightarrow void
Description: \langle seq_alignment constructors 31 \rangle create a new object for the given \langle score_matrix \rangle and vector<char>
Precondition: -
Postcondition: -

seq_alignment (const score_matrix&; biostring; biostring) \rightarrow void
Description: \langle seq_alignment constructors 31 \rangle create a new object for the given \langle score_matrix \rangle and alibio::biostrings
Precondition: -
Postcondition: -

seq_alignment (const score_matrix&; const string&; const string&) \rightarrow void
Description: \langle seq_alignment constructors 31 \rangle create a new object for the given \langle score_matrix \rangle and strings
Precondition: -
Postcondition: -

~seq_alignment (void) \rightarrow void
Description: \langle seq_alignment destructor 33 \rangle deallocates memory used by the \langle seq_alignment 25 \rangle
Precondition: -
Postcondition: -

print_alignment (void) \rightarrow void
Description: \langle print_alignment 34 \rangle prints the last calculated alignment
Precondition: -
Postcondition: -

nw_align (void) \rightarrow void
Description: \langle nw_align 35 \rangle calculates a global alignment using linear gap penalties
Precondition: -
Postcondition: -

sw_align (void) \rightarrow void
Description: \langle sw_align 36 \rangle calculates a local alignment using linear gap penalties
Precondition: -
Postcondition: -

nw_affine_align (void) \rightarrow void
Description: \langle nw_affine_align 38 \rangle calculates a global alignment using affine gap penalties
Precondition: -
Postcondition: -

sw_affine_align (void) \rightarrow void
Description: \langle sw_affine_align 39 \rangle calculates a local alignment using affine gap penalties
Precondition: -
Postcondition: -

semiglobal_affine_align (void) \rightarrow \text{void}
Description: ⟨semiglobal_affine_align 40⟩ calculates a semiglobal alignment using affine gap penalties
Precondition: -
Postcondition: -

nw_linear_align (void) \rightarrow \text{void}
Description: ⟨nw_linear_align 44⟩ calculates a global alignment using affine gap penalties and linear memory
Precondition: -
Postcondition: - sw_linear_align (void) \rightarrow \text{void}
Description: ⟨sw_linear_align 45⟩ calculates a local alignment using affine gap penalties and linear memory
Precondition: -
Postcondition: - semiglobal_linear_align (void) \rightarrow \text{void}
Description: ⟨semiglobal_linear_align 46⟩ calculates a semiglobal alignment using affine gap penalties and linear memory
Precondition: -
Postcondition: -
Bibliography


[8] ALiBio: Algorithm Library for BIOinformatics


[14] LATEX: Tobias Oeitker: The not so short introduction to LATEX, Copyright ©1998 Tobias Oeitker and all contributors

