

Module: CMP-5015A Programming 2

**Assignment: C++ Timed Programming Reassessment Exercise** 

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Date set: 5th August 2024

Value: 30%

Date due: 6th August 2024 3pm

**Returned by:** N/A

**Submission:** Blackboard

**Checked by:** 

### **Learning outcomes**

On completing this exercise, you will have demonstrated the ability to write a relatively simple class in the C++ programming language, including encapsulation, the use of constructors, accessor methods, operator overloading (particularly for use in stream-based I/O) etc. You will also have demonstrated your ability to perform unit testing of your class and integrating it into an application written by someone else, simulating the way programs are typically produced in industry — by a team rather than an individual programmer. The other parts of the application however may have a number of errors, so the exercise also tests your ability to understand code written by others and to debug and improve it, which are also key skills for any programmer, as maintenance is an important part of the software life-cycle for most non-trivial programs. The exercise may require the use of components from the C++ standard libraries that have not been explicitly covered in the lectures or seminars. The ability to use API documentation to understand and use library components is another programming skill demonstrated by successful completion of the task)

# **Specification**

## Overview

The aim of this assignment is to develop programming, testing, debugging and refactoring skills, by contributing a class to a C++ application that performs a simple analysis of a database of proteins, and then testing, debugging and improving it.

## Description

Proteins are the most basic components of the mechanisms of our cells, and are composed of a string of elementary building blocks, known as amino acids. The function of proteins in our bodies depends on their shape, which is determined by this sequence. Most proteins naturally fold to form a compact shape, but some don't and are described as "intrinsically disordered proteins", and are of considerable scientific and medical interest, for instance the prion protein that gives rise to Creutzfeldt–Jakob disease. Some amino acids are hydrophilic (like being in water) and others are hydrophobic (dislike being in water), and many proteins fold such that

Table 1: Table showing the hydrophobicity of amino acids, according to the Kyte-Dootittle scale, and the electrical charge.

Amino Acid	Symbol	Hydrophobicity	Charge
Alanine	Α	+1.8	+0
Arginine	R	-4.5	+1
Asparagine	N	-3.5	+0
Aspartic acid	D	-3.5	-1
Cysteine	С	+2.5	+0
Glutamine	Q	-3.5	+0
Glutamic acid	E	-3.5	-1
Glycine	G	-0.4	+0
Histidine	Н	-3.2	+1
Leucine	L	+3.8	+0
Isoleucine	I	+4.5	<b>+</b> 0 €
Lysine	K	-3.9	
Methionine	М	+1.9	+0
Phenylalanine	F	+2.8	+0
Proline	Р	-1.6	+0
Serine	S	-0.8	+0
Threonine	T	Ø.7	+0 1
Tryptophan	W	-0.9	<del>+0</del>
Tyrosine	Υ	-1.3	10
Valine	V	+4.2	+0

the hydrophilic amino acids are on the outside of the protein and the hydrophobic amino acids are hidden inside the folded protein in the watery cytoplasm within the cells of our body. Amino acids are also able to carry an electrical charge and this is also thought to affect the ability of a protein to fold to form a stable snape in solution.

Prilusky et al. [2005] suggest a simple mathematical method, called "FoldIndex", to predict whether a protein is likely to be intrinsically disordered, based on the hydrophobicity and charge of its amno acid sequence:

$$U = 2.785 \times H - R - 1.151,$$

where R is the average (arithmetic mean) hydrophobicity of the amino acids, and R is the absolute magnitude of the difference between the number of positively charged and negatively charged amino acids, divided by the length of the sequence. Table 1 shows the electrical charge and hydrophobicity (according to the Kyte-Doolittle scale) for each amino acid. Note that before the hydrophobicities are used to calculate the FoldIndex, they must be scaled to lie in the range o to a (i.e. such that Arginine has a normalised hydrophobicity of zero and Isoleucine a normalised hydrophobicity of one). Positive values of FoldIndex suggest the protein will be folded, negative values suggest the propensity for intrinsic disorder — the more negative the value, the more disordered the protein is likely to be.

The program is intended to load a database of proteins, update it to include calculated values for the FoldIndex for each protein, and then save the database to another file, sorted in descending order of the propensity for intrinsic disorder. The main part of the prototype application main.cpp and the input database, disprot.csv are provided on BlackBoard. Your main task is to implement a class called Protein, to represent all of the information about a given protein, comprised of:

ID - a unique identifier assigned to each protein (stored as a string).

sequence - a string of upper-case characters describing the sequence of amino acids conprising the protein (c.f. Table 1).

label - a string of the same length, where each character indicates that the corresponding amino acid is either part of a folded region, indicated by a '-', or part of an intrinsically disordered region, indicated by a '#'.

foldIndex - a value of the FoldIndex for that protein (or zero if the value has yet to be computed), stored as a double.

The class must provide default and ordinary constructors, an appropriate set of accessor methods, a method called setFoldIndex used to set the value of the FoldIndex for the protein, a full set of relational operators that order Protein objects according to their foldIndex value and operators (<< and >>) for stream-based I/O. The required format for both operators is illustrated by the format of disprot.csv. Note that the format used by both << and >> must be identical, so that I/O operators are mutually compatible (hint: std::quoted). The interfaces for the methods/functions must be compatible with their usage in main.cpp. Note: implementing this class does not require any understanding of protein structure, beyond that given in this assignment brief.

The next step is to write a test harness to perform "unit testing" of the class you have written. This should be implemented in a source code file called ProteinTest.cpp, which should provide a function called testProtein that has no parameters and returns true if the code passes all tests and false otherwise. It should document the steps taken to test your component of the program, and should be easily repeatable so that if the source code is modified, it is easy to check that a regression error had not been introduced. The test harness should not generate large amounts of diagnostic information, especially where the code passes the tests, as this would make testing for regression errors unduly time consuming.

The third step is to integrate your class with the prototype program, provided via Black-Board. Sadly all non-trivial programs contain bugs, and this program is no exception. Test the system as a whole, and identify and correct any bugs that you find. These may prevent the program from compiling, they may cause the program to crash, or they may cause the program to run, but produce incorrect answers.

The last step is to improve the efficiency of the program, by refactoring main.cpp so that it stores a vector of pointers to Protein objects, rather than storing the objects directly (which may result in a lot of unnecessary copying). This sort of modification and improvement of programs is a common activity in the maintenance phase of the software life-cycle.

### References

Jaime Prilusky, Clifford E. Felder, Tzviya Zeev-Ben-Mordehai, Edwin H. Rydberg, Orna Man, Jacques S. Beckmann, Israel Silman, and Joel L. Sussman. FoldIndex©: a simple tool to predict whether a given protein sequence is intrinsically unfolded. *Bioinformatics*, 21(16): 3435–3438, 06 2005. ISSN 1367-4803. doi: 10.1093/bioinformatics/bti537. URL https://doi.org/10.1093/bioinformatics/bti537.

### Relationship to formative assessment

This assignment builds on skills developed through the lab exercises on this module and those on CMP-4008Y - Programming 1.

#### **Deliverables**

Your solution **must** be submitted via Blackboard. Your solution **must** be in the form of a single .pdf file, formatted using PASS, via the PASS server at http://pass.cmp.dea.ac.uk, which can be accessed directly while on campus, and via VPN or the Remote Desirop system from outside the campus (for details, see https://supportwiki.cmp.uea.ac.uk/public/rdpstudent). The PASS program formats your source code, and compiles and runs the program, appending any compiler messages, and the output of your program to the .pdf file. It is your responsibility to check the .pdf file generated by PASS; if there is a problem contact me as soon as possible (mailto:gcc@uea.ac.uk). Do not leave it to the last possible moment to generate the .pdf file — there is a limit to the amount of he(p) am able to give if there is little time remaining until the submission deadline. If you upload your work to the PASS server but do not upload it to the correct subtission point on BlackBoard, it will be considered a non-submission and a mark of zero will be awarded.

Your submission must include at least the following files (but may contain more):

- Protein.h
- Protein.cpp
- ProteinTest.cop
- main.cpp -an initial prototype provided via BlackBoard

PASS is the target environment for the assignment. If your program does not operate correctly using PASS, it doesn't work, even if it gives the correct output on your own machine. The PASS program uses the gcc/g++ compiler from the GNU Compiler Collection tools. It will be set to conform to the C++17 standard. It is *your* responsibility to ensure that your program is compatible with that compiler and that standard.

- The PASS program is not able to provide input to your program via the keyboard, so programs with menu systems, or which expect user input of any kind are not compatible with PASS. Design your program to operate correctly without user input from the keyboard.
- Your code *must* contain only one main function. PASS will compile and link together all of the source files, so if more than one main function is defined, the compiler has no way to determine which one to use and there will be a linker error.

- Do not use absolute path names for files as the PASS server is unable to access files on your machine. If your program is required to load data from a file, or save data to a file, the file is expected to be in the current working directory. If the program is required to load data from a file called rhubarb.txt then the appropriate path name would be just "rhubarb.txt" rather than "C:some\long\chain\of\directories\rhubarb.txt".
- If you develop your solution on a computer other than one of the laboratory machines, make sure that you leave adequate time to test it properly with PASS, in case of any unforeseen portability issues.
- If your program works correctly under CLion on the laboratory machines, but does not
  operate correctly using PASS, then it is likely that the data file you are using has become
  corrupted in someway. PASS downloads a fresh version of the file to test your submission,
  so this is the most likely explanation. Issues relating to the conversion of line terminations
  (line feed and carriage return characters) are a common example of such problems.
- If there are problems, a solution may be found on the PASS Frequently Asked Questions (FAQ) list http://pass.cmp.uea.ac.uk/faq.php.

#### **Resources**

- <a href="https://stackoverflow.com/">https://stackoverflow.com/</a> An excellent site for finding information about specific issues relating to various programming languages, including Java. It is important however not to become too reliant on sites such as StackOverflow, which are great for details, but don't give the "big picture", so it is difficult to get a good understanding of programming in this way. Note that if you re-use or modify code found on-line, then you must provide a comment giving the URL and a brief explanation of what modifications have been made. Re-using code found on line without proper attribution would constitute plagiarism. Obviously it would also be an offerce to ask questions directly relating to this assignment on StackOverflow or other programming forums.
- https://cppreference.com Ar excellent resource for C and C++ programming, providing detailed documentation on both the language and the libraries. This is the site I use most often when an programming (as well as using books).
- It is likely that the fectures and live programming demonstrations for this module provide many examples of e++ programming techniques and idioms that may be useful for this assignment.

# Marking Scheme

Mark will be awarded according to the proportion of specifications successfully implemented, programming style (indentation, good choice of identifiers, commenting etc.), and appropriate use of programming constructs. It is not sufficient that the program generates the correct output (professional programmers are required to produce maintainable code that is easy to understand, easy to debug when (rather than if) bug reports are received and easy to extend. The code needs to be modular, where each module has a well-defined interface to the rest of the program. The function of modules should be made as generic as possible, so that it not only solves the problem specified today, but can easily be modified or extended to implement future requirements without undue cost. The code should also be reasonably efficient. Marks may also be awarded for correct use of more advanced programming constructs not covered in the lectures.

#### Breakdown of marks:

- Implementation of Protein.h and Protein.cpp Marks will be awarded according to the quality and correctness of the code, according to the specifications. [15 marks]
- Unit Testing A file called ProteinTest.cpp must be implemented, demonstrating the steps taken to test Protein.h and Protein.cpp. Marks will be awarded according to the quality and thoroughness of the tests. [5 marks]
- System testing Marks are awarded for identifying and correcting the errors/bugs in the file main.cpp and for fully and accurately describing the nature of the error in the revision history in the block comment at the start of the file. [5 marks]
- Improving efficiency The original prototype is somewhat inefficient as it is based on a vector of Protein objects, which may result in unnecessary copying of objects. Marks will be awarded according to the quality of the refactoring. Note that substantially rewriting main.cpp will not attract a good mark as the new code will require testing and debugging, so adding large amounts of code is likely to reduce the reliability of the program. [5 marks]

# Plagiarism, collusion, and contract cheating

The University takes academic integrity very seriously. You must not commit plagiarism, collusion, or contract cheating in your submitted work. Our Policy on Plagiarism, Collusion, and Contract Cheating explains:

- what is meant by the terms 'plagiarism' 'collusion', and 'contract cheating'
- how to avoid plagiarism, collusion, and contract cheating
- using a proof reader
- what will happen if we suspect that you have breached the policy.

It is essential that you read this policy and you undertake (or refresh your memory of) our school's training on this. You can find the policy and related guidance here:

https://my.lea.ac-uk/departments/learning-and-teaching/students/academic-cycle/regulations-and-discipline/plagiarism-awareness

The policy allows us to make some rules specific to this assessment. Note that:

in this assessment, working with others is *not* permitted. All aspects of your submission, including but not limited to: research, design, development and writing, must be your own work according to your own understanding of topics. Please pay careful attention to the definitions of contract cheating, plagiarism and collusion in the policy and ask your module organiser if you are unsure about anything.