



School of Computer Science and Engineering

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The University of New South Wales

Bioinformatics of the Immune System

by

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Abstract

This document describes the requirements to theses submitted for the Bachelor of Engineering in Software Engineering degree at the School of Computer Science and Engineering. Requirements described are that of both of context and layout of the theses. The document is written using the L^AT_EX template provided by the school.

Acknowledgements

This work has been inspired by the labours of numerous academics in the Faculty of Engineering at UNSW who have endeavoured, over the years, to encourage students to present beautiful concepts using beautiful typography.

Further inspiration has come from Donald Knuth who designed T_EX, for typesetting technical (and non-technical) material with elegance and clarity; and from Leslie Lamport who contributed L^AT_EX, which makes T_EX usable by mortal engineers.

John Zaitseff, an honours student in CSE at the time, created the first version of the UNSW Thesis L^AT_EX class and the author of the current version is indebted to his work.

Abbreviations

HMM Hidden Markov Model

BLAST Basic Local Alignment Search Tool

XML Extensible Mark-up Language

NCBI National Centre for Biotechnology Information



Contents

1	Introduction	1
1.1	Immunoglobulin recombination	1
1.2	iHMMuneAlign	3
1.3	Hidden Markov Models	3
2	Analysis	5
2.1	Run Time	5
2.1.1	Interpretation of Results	8
2.2	Profiling	9
2.2.1	CPU Call Graph	9
2.2.2	Threads	11
3	Proposed Solution	13
3.1	Re-implementing From Scratch	13
3.1.1	Code Maintainability	13
3.1.2	Concurrency / Thread Safety	14
3.1.3	Libraries and Language	14
3.1.4	Implementation Stages	15
3.2	Dependencies of iHMMuneAlign	15
3.2.1	BioJava	15

3.2.2	DP Module	17
3.2.3	External Library	17
3.2.4	Custom Implementation	18
3.3	Concurrency	18
3.3.1	Per Sequence	19
3.3.2	Per Processing Stage	20
4	Design & Implementation	21
4.1	Initialization	21
4.1.1	IGHD/IGHJ Repertoires	21
4.1.2	IGHV Repertoire	22
4.1.3	Mutation Ratios	22
4.2	BLAST	23
4.2.1	Invoking Binary	23
4.2.2	Linking Shared Library	24
4.3	Pipeline	26
4.3.1	Parsing Blast Results	26
4.3.2	Calculating A-Score	26
4.3.3	Generating Model	26
4.3.4	Running Viterbi	26
4.3.5	Printing Results	26
4.4	Approach	26
5	Maintenance	28
5.1	Revision Control	28

6	Performance	29
6.1	Blast Lib	29
7	Evaluation	30
7.1	Results	30
7.2	Discussion	30
8	Conclusion	31
8.1	Future Work	31

List of Figures

1.1	The IGHV, IGHD and IGHJ gene recombination process	2
1.2	An example Hidden Markov Model [31]	2
2.1	Total time taken (in seconds) to process the number of sequences (Wall-clock time). n is the number of sequences per invocation of the program	6
2.2	Total time spent doing work by each CPU on behalf of the process during execution (CPU Time). n is the number of sequences per invocation of the program	7
2.3	Time spent by each thread; running () and waiting ()	10
2.4	The call graph generated by jProfiler, for a single run of iHMMuneAlign	10
4.1	An extract of a mutation probabilities file	23
4.2	Overview of the Model Solving ‘Pipeline’	27

Chapter 1

Introduction

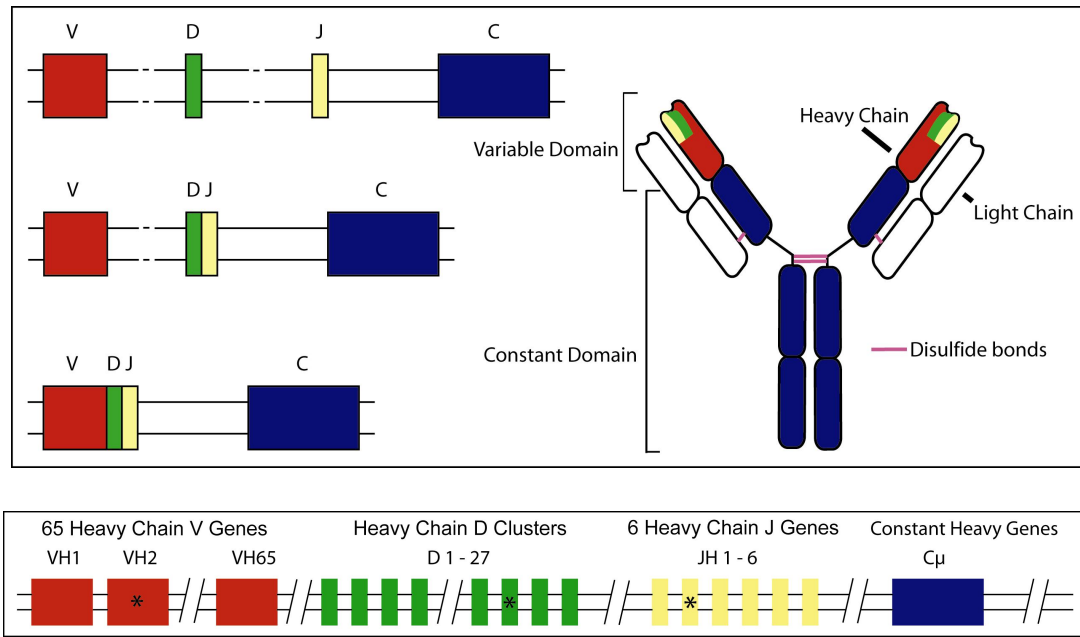
1.1 Immunoglobulin recombination

The human immune system faces many challenges when dealing with foreign pathogens invading the body. One of them is identifying and classifying antigens that are not only vastly diverse, but also constantly mutating and evolving.

It does so by creating immunoglobulins (also called antibodies) whose job is to identify and bind to certain antigens, effectively marking them for destruction by the rest of the immune system. The process, by necessity is highly selective - a given antibody will only bind to a very specific antigen.

However, to deal with the highly diverse and evolving nature of foreign antigens, antibodies must mutate very rapidly. This allows them to be as diverse as the antigens they face, and adapt to bind to newly encountered pathogens.

The process (essentially randomly) selects 3 germline genes from a repertoire in the person's DNA potentially consisting of between 38 and 45 IGHV genes [13, 14], 23 IGHD genes [12] and 6 IGHJ [22] genes. These selected genes are then mutated (adding, removing or changing nucleotides) to generate further diversity. The antibodies that successfully bind to the antigen emit a chemical signal, which then encourages the



Katherine Jackson

Figure 1.1: The IGHV, IGHD and IGHJ gene recombination process

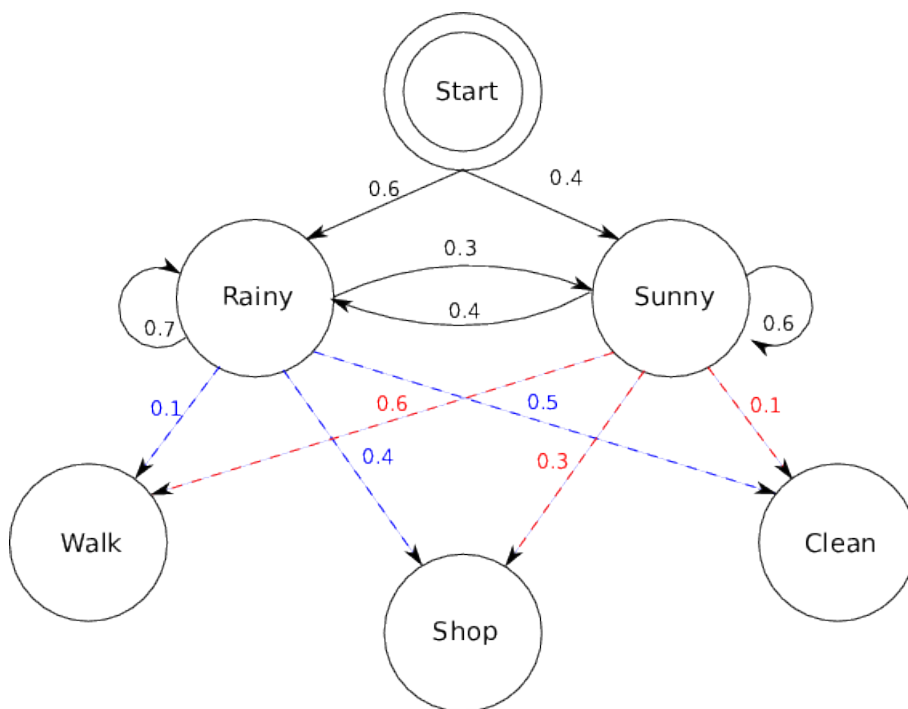


Figure 1.2: An example Hidden Markov Model [31]

production of similar antibodies. This entire process is illustrated in Figure 1.1 and explained in more detail by Gaëta *et al.*[7].

1.2 iHMMuneAlign

As a result of this gene recombination and mutation process, the germline genes often don't provide much information on rearranged genes found in the B-Cells that create the antibodies themselves.

Developed in 2007 by a team of researchers at UNSW, iHMMuneAlign attempts to simulate the process of immunoglobulin gene rearrangement.[7].It represents the recombination process of immunoglobulin heavy chain genes as a hidden Markov model, and then uses it to find the most probable set of germline genes that created the rearranged gene.

First, it selects the most likely IGHV germline gene from its repertoire of possible V genes, using the NCBI's Basic Local Alignment Search Tool (BLAST) [1]. It then constructs a Hidden Markov Model with that IGHV gene, and all the possible IGHD and IGHJ genes, together with other states representing the somatic mutation.

It then uses the Viterbi algorithm to determine the most likely sequence of states (genes and their mutations) that caused the observed emissions (the rearranged sequence given as input).

1.3 Hidden Markov Models

A Hidden Markov Model is effectively a Markov Chain, where the states cannot be observed directly. They are used in a wide variety of machine learning applications, including speech recognition [6].

An example of a HMM is shown in Figure 1.2. In this example, if the sequence of

emissions **Walk** was observed, the probability of it being **Sunny** is

$$I(\text{Sunny}) \times E(\text{Sunny}, \text{Walk}) = 0.4 \times 0.6 = 0.24$$

and the probability of it being **Rainy** is

$$I(\text{Rainy}) \times E(\text{Rainy}, \text{Walk}) = 0.6 \times 0.1 = 0.06$$

where $I(X)$ is the initial probability of X and $E(X, Y)$ is the emission probability of state Y in state X .

If **Walk**, **Clean** were observed. The probability of it having been **Rainy** then **Sunny** is

$$I(\text{Rainy}) \times E(\text{Rainy}, \text{Walk}) \times T(\text{Rainy}, \text{Sunny}) \times E(\text{Sunny}, \text{Clean}) = 0.6 \times 0.1 \times 0.3 \times 0.1 = 0.0018$$

where $T(X, Y)$ is the probability of transitioning from state X to state Y .

The Viterbi algorithm uses dynamic programming to recursively explore these probabilities, and selects the sequence of states with the highest probability at the end.

Chapter 2

Analysis

Since the high level aim of this project is to improve the performance of iHMMuneAlign, it was necessary to analyse the existing implementation in order to gain insight into its structure, as well as to create realistic aims.

2.1 Run Time

One of the primary attributes of performance, is run time. We begin by measuring the run time of the existing implementation and -more importantly- how it grows with workload.

The current implementation consists of a Java executable and a shell script, to allow running in batches. The Java executable (which does the actual processing) takes a range of input from the script and attempts to process that many in parallel. It then invokes Java program after breaking up the entire range into batches of a given size.

Since we are interested in the rate the total runtime changes with workload as well with increasing parallelism, both were measured. It was run with a total workload ranging from 1 sequences to 100 sequences, with batch size ranging from 1 to 8. The results are shown in Figure 2.1.

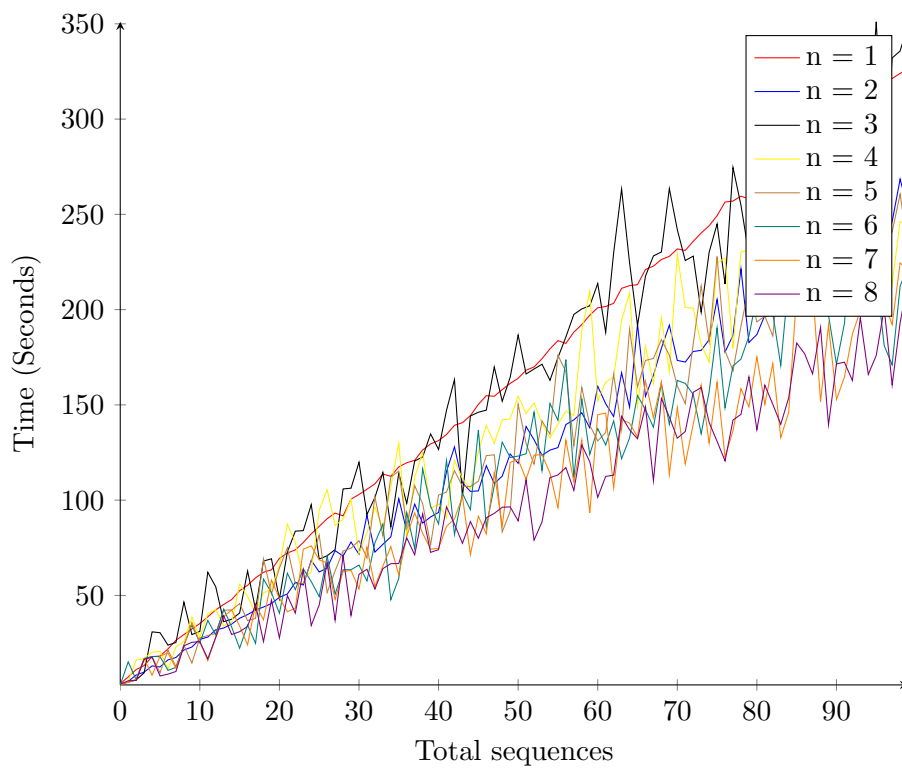


Figure 2.1: Total time taken (in seconds) to process the number of sequences (Wall-clock time). n is the number of sequences per invocation of the program

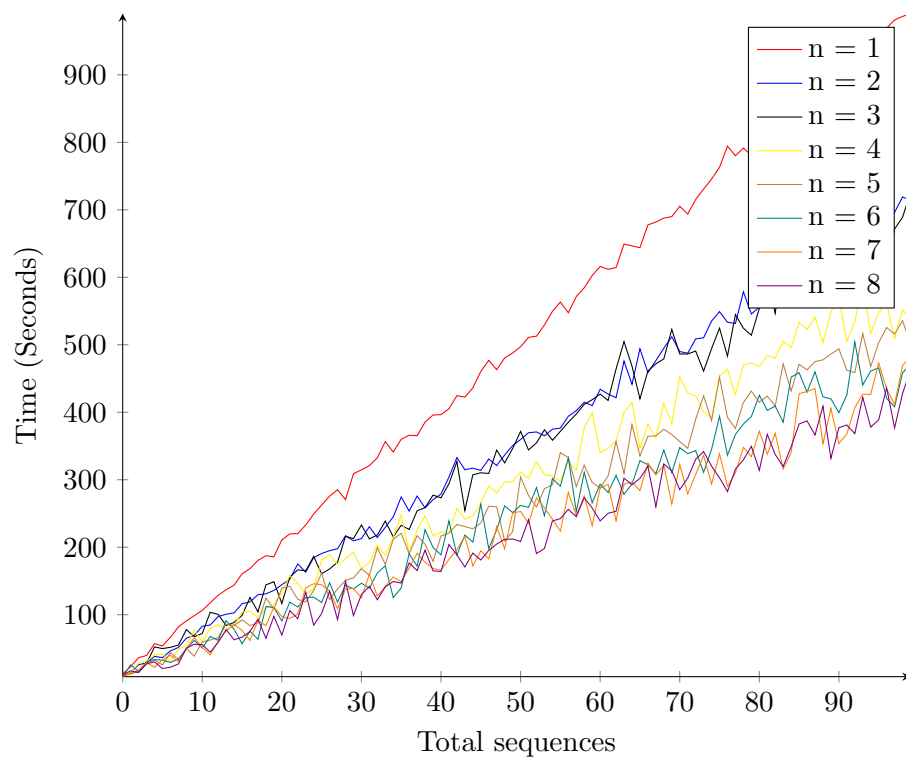


Figure 2.2: Total time spent doing work by each CPU on behalf of the process during execution (CPU Time). n is the number of sequences per invocation of the program

Note that this was performed on a machine with an AMD Phenom II 1090T (6-core) processor, 8GB of RAM and a solid state hard drive.

2.1.1 Interpretation of Results

With a batch size of 1, the expected linear increase was present. However, increasing the batch size shows a more sporadic decrease in runtime. There could be a number of causes for this, including the threads being assigned different CPUs by the scheduler on some runs but not others, increasing cache misses. Nonetheless, it wasn't investigated further, due to time constraints, and the fact that we didn't need such high accuracy for this test. However, there is an overall decrease in total execution time, as the batch size increases.

This suggests that the existing implementation is concurrent, and will utilize all the available processing power on the CPU. However, that should have diminishing returns once the amount of threads exceeds the total number of CPU cores on the machine, which is not the case here - execution time continues to decrease at roughly the same pace as n increases.

Looking at the total CPU time spent on the process, rather than total execution time provided a theory (which was later confirmed in Section 2.2.2, on page 11) to explain this behaviour. For each invocation of the Java program, there is a significant amount of overhead. For example, the Java Virtual Machine (JVM) needs to be initialized, it needs to load the program, analyse and perform any JIT compilation (Just-in-Time compilation, to improve execution time [24]). The program also needs to read in the repertoire data files for each gene and process them. On top of all that, any data that are being cached by the operating system on behalf of the program will be invalidated. Since this only needs to be done once per batch, as the batch sizes increases (and the number of batches decrease), this overhead also decreases.

This is clearly apparent in Figure 2.2, which shows that the total work done by the program to process all the sequences is reduced significantly as the n increases.

Comparing the total executable time (wall clock time) and the total CPU time of each run shows, that there was roughly double the CPU time as execution time. This indicates that the program has 2 threads of execution, but that it doesn't increase threads with workload.

2.2 Profiling

In order to determine the most appropriate approaches for improving performance and overcoming the limitations of the original implementation, further analysis is required. Specifically, we are interested in which parts of the program consume the most CPU time and memory, as well as how they increase with the workload.

The method for collecting this data is called profiling. Since the original implementation of iHMMuneAlign was written in Java, we can use various interfaces in the Java Virtual Machine (JVM) to gather this data. The JVM has an interface called Java Native Interface (JNI)[19], that allows external programs and native libraries (usually written in C/C++) to interact with Java objects. Alongside that, from Java version 1.5 onwards, the JVM also provides another interface, called Java Virtual Machine Tool Interface (JVMTI), which allows external tools to be notified as certain events (such as object allocation, method call) occur during execution.

In Figure 2.4, the data for a run of iHMMuneAlign with 6 sequences, instrumented by jProfiler [23] (a profiler implemented with the above method) is shown.

2.2.1 CPU Call Graph

Figure Figure 2.4 shows the time, and the percentage of total execution time spent in each method invocation of a single execution of iHMMuneAlign.

It begins with the equivalent of the `main()` function for a thread using Java's built in threading library - `Thread.run()`, which in turn calls `startAlignment()`. However, it

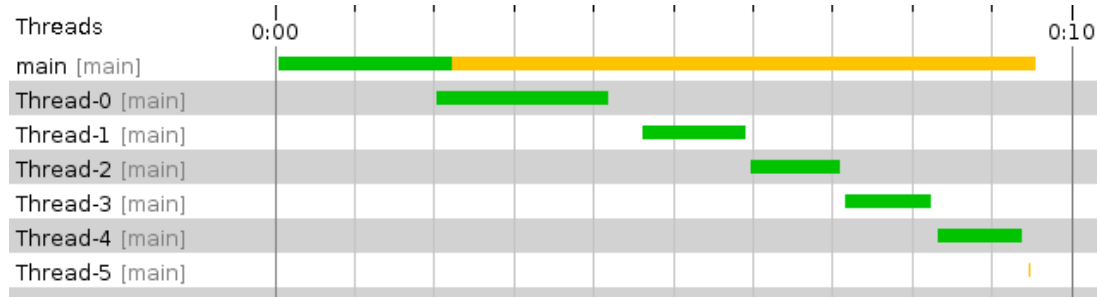


Figure 2.3: Time spent by each thread; running (■) and waiting (■)

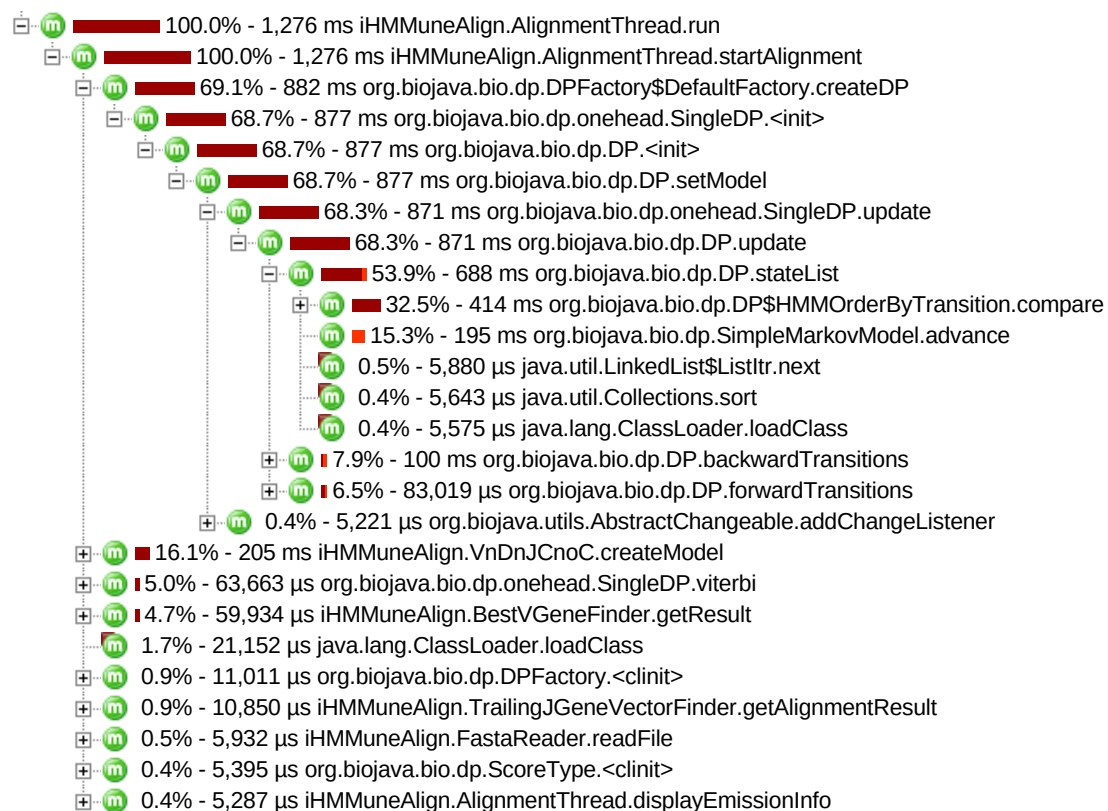


Figure 2.4: The call graph generated by jProfiler, for a single run of iHMMuneAlign

becomes immediately apparent that the majority of execution time from there is spent in the BioJava library.

68% of total execution time is spent constructing a `org.biojava.bio.dp.SingleDP` object, which will allow various algorithms (including Viterbi) to be performed. It takes a `org.biojava.bio.dp.MarkovModel` object, which represents the Markov Model, and converts it to its own internal format. This conversion involves sorting all the states, which is what 53.9% of the whole program's execution time is spent on.

For our purposes, this conversion is unnecessary, as we can generate the states ourselves, and can easily do so in the order necessary to run Viterbi on them. The actual call to the `DP.viterbi()` method, which actually runs the Viterbi algorithm to determine the most likely sequence of states, takes only 5.0% of the total execution time.

2.2.2 Threads

An important aspect of the existing implementation is its threading model - i.e. how it distributes the workload between multiple threads of execution (which can then run on multiple CPUs or CPU cores). Since we already determined in Section 2.1.1 that it does indeed use multiple threads, we need to look more closely at exactly how it does so.

For this, we have our profiler record events in the `java.util.concurrent` package, which include thread creation, destruction and blocking (waiting for another thread to complete). The results of this, shown in Figure 2.3 confirm our earlier hypothesis. The program has two threads active most of the time; one processing the gene, and one (main thread) waiting for it to complete, before starting the next one.

The fact that they use a threading library indicates that the original authors intended to make `iHMMuneAlign` a concurrent program, processing multiple gene sequences in parallel. While their architecture would have been less performance efficient than using a thread pool (as they have to pay the cost of thread creation and destruction for each

sequence), it would still have caused a notable improvement if they allowed the threads to execute in parallel.

Nonetheless, the fact that the main thread doesn't create each thread until its predecessor terminates indicates said efforts of parallelism were abandoned. It is widely regarded in industry that concurrent programming is rather difficult and the original authors most likely deemed completing the project was more prudent than making it concurrent.

No further investigation was done to determine what would need to be done in order to make the current implementation concurrent, as it will most likely be rewritten for this project.

Chapter 3

Proposed Solution

3.1 Re-implementing From Scratch

Based on the above analysis, it is clear that the original codebase has many pitfalls, and limitations that make it difficult to proceed further.

3.1.1 Code Maintainability

Most of the logic for processing a gene sequence is contained in the

```
AlignmentThread.startAlignment()
```

method, which over 2000 lines long. The project doesn't use any revision control systems, and the fact it is littered with commented out code most likely indicates that it didn't do so at any point in the past either.

There are no unit tests that cover most of the program logic, and it would be difficult to write any, since the logic hasn't been split up. This makes it rather difficult to make changes, without introducing any new subtle bugs, or undesired behaviour.

The original authors are no longer available to provide assistance for understanding their codebase, and the style guide they followed (or lack thereof) makes it difficult to read and follow.

It is understandable that the original authors did not prioritize long term code maintainability, as they could have been rushed for time, or they may have deemed it unlikely their code would be used and maintained 7 years later.

3.1.2 Concurrency / Thread Safety

Concurrency must be considered from the very early design decisions onwards, for a program to be efficient and parallel. The fact the original authors abandoned their efforts for parallelism indicates that there may have been concurrency bugs, like race conditions [18] that are very difficult to trace.

Access to shared data must be carefully controlled, and synchronized to avoid data corruption (and race conditions). The potentially high overhead of synchronization should lead to program structure designs where shared data between threads is minimal. In addition, the thread safety of any libraries used must be carefully considered - if a class is not thread safe, it shouldn't be shared between threads; and if a class is not re-entrant, there shouldn't even be separate instances in different threads.

3.1.3 Libraries and Language

Some of the libraries that the original authors used have since been deprecated. The specific libraries are mentioned in more detail below. Considering that some libraries account for most of the performance inefficiencies of the program (as shown in Section 2.2.1), they will certainly need to be replaced. Since the libraries' APIs heavily influenced the structure of the code, replacing them while maintaining the existing code structure will be a difficult task.

There is also the language itself to consider: Java programs tend to be 2-3 times slower

than their equivalent C++ counterparts, and even more so in CPU bound tasks like this [3]. Reimplementing from scratch will allow us to do so in a different language. C++ is a good candidate - it is quite popular for constructing large scale, high performance applications [8]; it contains high concepts from high level languages, while still allowing for low level optimizations, including memory management.

3.1.4 Implementation Stages

The initial version of the new implementation will be loosely based off the already existing implementation. Once the code is rewritten and structured in a manner that lends itself to easy modification, the real optimization (such as threading, and caching computation) can begin.

3.2 Dependencies of iHMMuneAlign

Apart from the codebase, the current implementation has several dependencies that perform significant amounts of computation. This section outlines how the new implementation will replicate their features.

3.2.1 BioJava

BioJava [11] is a general purpose Bioinformatics library, that contains several useful features for iHMMuneAlign. However, since iHMMuneAlign was written, BioJava has undergone a complete restructuring [21], and most -if not all- the modules the original implementation uses have been deprecated.

Fasta Reader

FASTA was originally the format used by a program that bore its name to store databases of DNA sequences [20]. These days, it is commonly used in Bioinformat-

ics to represent large quantities of gene sequences within a file. BioJava could parse files FASTA format and convert them to its in-memory representation, which were then used by iHMMuneAlign. Since we no longer intend to use BioJava, an alternate method was required to parse the inputs and repertoire files for each genes was required. Since the FASTA format is a very simple format, a custom parser could be built quite easily. In addition, the NCBI C++ Toolkit can also be used to trivially parse FASTA files.

Bioinformatic Data Structures

BioJava contains a large collection of pre-built data structures to represent commonly occurring biological objects and concepts (E.g. RichSequence, FiniteAlphabet). Here we have several viable solutions.

Bio++ is a set of general purpose Bioinformatics libraries written in C++ [5]. Much like BioJava, it also provides various pre built data structures that represent various biological entities, like gene sequences. However, how well they correspond to the used data structures from BioJava and how useful they are when re-implementing is yet to be seen.

Qt is a C++ framework, originally created by Trolltech and now maintained by Digia [32]. It has various components that assist in the rapid building of user interfaces, as well complex systems and algorithms. While they are less specific to biology, they are designed for speed and efficiency. Once the initial version is built using Bio++, if Bio++'s data structures are deemed a significant performance bottleneck, new ones will be custom built using Qt's support structure.

STL The Standard Template Library (STL) that accompanies C++ also contains a series of very high performance, generic data structures and algorithms [17]. However, it is widely considered to be less user friendly than its alternatives, like Qt. For this

reason, Qt's data structures were selected as the primary candidate for a custom implementation, and if they cause a significant performance limitation, they will be replaced by their STL counterparts.

Boost is another, highly comprehensive set of libraries that extend the functionality of the STL [4]. Select features from Boost often end up in subsequent revisions of the STL. Should functionality of the STL be insufficient to replace the Qt implementation once it's been deemed a performance bottleneck, Boost will be used to compensate.

3.2.2 DP Module

We have determined in Section 2.2.1 that the DP module (in the `org.biojava.bio.dp` package) accounts for more than two thirds of the total runtime, with half on a data conversion that is ultimately unnecessary.

Since the `iHMMuneAlign` was written, BioJava has undergone a major overhaul [21] and the entire module is now deprecated. For replacing the parts of its functionality that could have been useful moving forward (namely, the implementation of Viterbi algorithm), we have two choices:

3.2.3 External Library

Considering Hidden Markov Models are a very popular for a variety of applications [6], there are several C++ implementations available for use. For this, two of them were evaluated

MLPack is a scalable C++ machine learning library [2], by the Georgia Institute of Technology. It has a module to deal with Hidden Markov Models, which includes an implementation of Viterbi. However, the implementation entertains the possibility of

every state transitioning to every other state, and as such performed too slow in our tests.

StochHMM is a library and set of tools dealing with Hidden Markov Models, by the University of California [15]. Its evaluation was done by feeding in the model as a text file to its Viterbi solver, which crashed. The cause of the crash was not determined, as we moved on to other solutions.

3.2.4 Custom Implementation

After the failures of the other approaches, the possibility of implementing the algorithm ourselves was evaluated. A proof of concept implementation was created, tested with a roughly equivalent workload, and showed an order of magnitude improvement over the original solver from BioJava.

Later on in the project, specifically when the model generation code is implemented, its performance can be evaluated more accurately. The final decision regarding its use will be made at the time, depending on the performance of the alternatives.

3.3 Concurrency

Since the advent of multi-core and multi-processor machines, concurrency has become essential for high performance applications [25]. Since clock speeds have stagnated, yet the parallelism available in modern hardware continues to rise designing our implementation to be concurrent from the start will ensure that it can take advantage of the speed improvements granted by future generations processors.

The nature of our problem presents us two obvious ways of splitting it up to distribute over multiple threads of execution.

3.3.1 Per Sequence

The fundamental idea is to have each processing a different input gene sequence at the same time. This is the approach the authors of the original implementation attempted, although their implementation created a new thread for each sequence.

Creating a thread requires at least one system call (which is very expensive [29]) and destroying requires another. Additionally, the operating system has to do more book-keeping, as well as preparing a bunch of extra resources for the thread, which then have to be cleaned up - causing a significant performance cost [30].

A much better performing design would be to create a pool of worker threads at the start, and a queue of sequences for them to process. Each thread will then pop the sequence at the top of the queue, process it, write out the result, pop the sequence that is now at the head of the queue and repeat. In this case, the primary limitation for parallelism would come from the queue itself - by virtue of being a shared data structure, it would require synchronization to maintain consistency [9].

If the synchronization overhead of the queue is found to be a significant limitation on parallelism, it can be skipped altogether. Since all the inputs sequences are passed to the program in a file, the file itself can be partitioned - then each thread would read, parse and process all the sequences within a partition.

Doing so has the advantage that it will reduce the synchronization necessary between threads. On the other hand, if the sequences in some partitions require significantly less work than others, some threads would finish much quicker than others and be idling instead of doing useful work. This can be mitigated somewhat, by using a technique called work stealing [10], where threads that have finished all their jobs take pending jobs from other threads.

3.3.2 Per Processing Stage

The other way, is to consider each sequence as a series of tasks, and perform each one in a different thread. It's infeasible to speed up the processing of an individual sequence using this method, as most (if not all) of these tasks would depend on the results of the previous, and initializing a thread could take longer than some of the tasks.

Considering that the goal of the project is to scale well with increasing numbers of sequences, the following approach makes more sense; tasks can be done in batches. I.e. have n threads process the first stage of $n \times k$ sequences (where k is an integer constant), and then have all n threads process the second stage and so on. This has the primary advantage of having much greater data locality (decreasing the working set, page faults, cache misses) and increasing performance significantly [28].

Per contra, this approach will be heavily reliant on multiple work queues to distribute the tasks of different sequences between threads. As explained in the above section, the synchronization overhead of the queues may become a serious bottleneck on parallelism - even more so, due to the much heavier reliance on it.

Chapter 4

Design & Implementation

4.1 Initialization

Since iHMMuneAlign is rather configurable, it requires a large amount of input data for each run that cannot be embedded in the code directly. This data has to be read during program initialisation and stored in memory such that it can be efficiently used in their required of the algorithm.

4.1.1 IGHD/IGHJ Repertoires

The use-case for the germline IGHD and IGHJ repertoires are fairly straight forward. For each nucleotide in each gene, a state needs to be created that represents the probability that gene caused it to be there.

Since this needs to be done for each gene/nucleotide in order, there's no need for random access, or efficient searching. They're stored in a contiguous block of memory in a dynamic (variable size) array. This provides good cache/memory locality [**cache-locality**], while being easy to use.

4.1.2 IGHV Repertoire

Unlike the other two genes, not every IGHV gene needs to be in the model. Instead, a BLAST search is performed, and (for now, at least) only the most likely hit –the closest germline IGHV gene– is used.

However, the BLAST results do not in fact give the sequence in its output; It only presents the relevant matching/aligned section of the sequence. Fortunately, it also gives the (unique) name of the gene, as specified in the repertoire file it was given.

Consequently, an index must be built from the gene repertoire that maps each gene name to its corresponding sequence. Considering this lookup needs to be done for each model, it should be as efficient as possible. This index is currently stored as a hash-table in memory, allowing for efficient, constant time lookup [16].

4.1.3 Mutation Ratios

A large part of what iHMMuneAlign attempts to model is mutation of various germline genes, during the recombination process. While it is known that some nucleotides are more likely to mutate to a certain one than others, determining this is out of scope for iHMMuneAlign [7]. As a result, iHMMuneAlign takes in these probabilities as input.

A simple map from a nucleotide to nucleotide would not be sufficient, as it depends on the two surrounding nucleotides as well. These ratios are given in a format specific to the previous implementation of iHMMuneAlign; an example can be seen in Figure 4.1.3.

The probabilities are parsed into double-precision floating point numbers; the string on the left is split up, with all the unnecessary characters (arrow and brackets) dropped. Doing so allowed for more efficient hashing when put into a hash-table in memory, effectively forming an index much like the IGHV Repertoire.

```

A(A->C)A  0.200711181
A(A->C)C  0.234567901
A(A->C)G  0.090360982
A(A->C)T  0.249740516
A(A->G)A  0.698044251
A(A->G)C  0.679012346
A(A->G)G  0.798111707
A(A->G)T  0.533465028
A(A->T)A  0.101244567
A(A->T)C  0.086419753

```

Figure 4.1: An extract of a mutation probabilities file

4.2 BLAST

The first step is to determine the most likely germline V gene is in the target sequence. Fortunately, this can be determined quite efficiently by aligning the input sequence with all known V genes [7]. For that we opted to use the excellent alignment tool by NCBI: BLAST [1], like the original implementation.

BLAST is distributed as source code, and can be used in two ways:

4.2.1 Invoking Binary

There are binaries for all the common tasks, that take in sequence repertoires as files in FASTA [20] format. The previous implementation used the `blastn` executable, which takes in a sequence in FASTA format, and writes the results (matching sequences in the repertoires) out to an XML file.

The primary advantage of using blast in this manner would be the ease of development. Since the binaries come pre-packaged, very little extra code would be required to invoke the binary from within iHMMuneAlign, and parse the XML output. However, doing so incurs a (potentially significant) performance penalty at run time due to the cost of:

Process Creation The operating system has to create a new process for the blast executable, perform all its bookkeeping operations, and clean up after it is completed.

Reading Input Files The executable would then have to read the database files from disk again, instead of using the copy that already in memory (residing in iHMMuneAlign’s address space).

Serialising Results BLAST, like most high performance programs, uses its own internal intermediate representation for the input and germline genes[1]. Once the alignment search is completed, it will then need to convert that data into a human readable, or standardised format that can be parsed by iHMMuneAlign.

Transferring Results Modern operating systems provide a certain level of isolation, such that a process can not directly access data in a section of memory belonging to another process [26]. As a result, the previous implementation required the BLAST executable to write its results to disk so it could be read.

While IO operations are generally expensive, ones involving disk access tend to be especially so [27]. This is due to the fact that disks are several orders of magnitude slower than main memory (RAM).

As explained in Section 2.1.1 (on page 8), these overheads were partially responsible for large performance inefficiencies in the original implementation.

4.2.2 Linking Shared Library

BLAST+ can also be compiled as a library, that can be statically or dynamically linked the target program using it. This would the alignment search to be performed directly in the process of iHMMuneAlign, effectively eliminating all of the overheads listed in the previous section.

The disadvantage of this approach is that it requires extra work during the development of iHMMuneAlign. This includes the work required for:

Acquiring Libraries The BLAST+ binary executables are quite prevalent in bioinformatics research and as such, they are readily available for most research/development platforms. The libraries however, are much less common, as they are generally used by developers working on production tools relating to BLAST+.

This would require researching the build process used by the BLAST+ libraries. This is a non-trivial task, as the documentation is somewhat lacking, and makes implicit assumptions about the build environment which do not necessarily hold outside of NCBI lab machines.

Programming for Libraries The binaries accept the input data and germline repertoires in FASTA format, and it outputs in XML. Both of these are widely used, standard formats; There are a range of utilities in most languages to deal with them. As a result, very little code would need to be written specifically for it.

Using the data-structures that comprise the internal representations used by the BLAST directly, would be much more complicated. And once again, the documentation regarding were not as comprehensive as they could have been.

Distributing Libraries The binaries are quite widespread, well known, and can be assumed to be already available on machines that iHMMuneAlign would run on. Thus, it would be the users' responsibility to ensure they have it installed on their machine, as it's a required dependency.

That same assumption can not be made for the library, and as a result we must distribute it with the program. Also –and more significantly–, the libraries must be available to the compiler when iHMMuneAlign is being built; This would significantly increase the work required to port iHMMuneAlign to a new platform, as well as making it infeasible to distribute primarily as source code.

Due to the performance considerations described in Section 4.2.1 (on page 23), the we opted to link to the BLAST libraries, and perform the alignment search directly in iHMMuneAlign. However, after spending a significant amount of time attempting to overcome the difficulties detailed in Section 4.2.2 (on page 24), we elected to use the binaries for now.

At the time of writing, the current implementation of iHMMuneAlign assumes the `blastn` binary has already been invoked, and expects to be given its results in XML format. This process can be made into a script easily, and will be revisited at a later date, once more impactful performance optimisations have been exhausted.

4.3 Pipeline

The entire process of creating a HMM to represent immunoglobulin recombination and solving it using Viterbi's algorithm can be split up into several smaller problems. These can be viewed as stages in an assembly line where the result of each one feeds into the next, terminating with the final result. In computing, this is called a pipeline.

4.3.1 Parsing Blast Results

4.3.2 Calculating A-Score

4.3.3 Generating Model

Generating V/D/J States

Generating Transitions

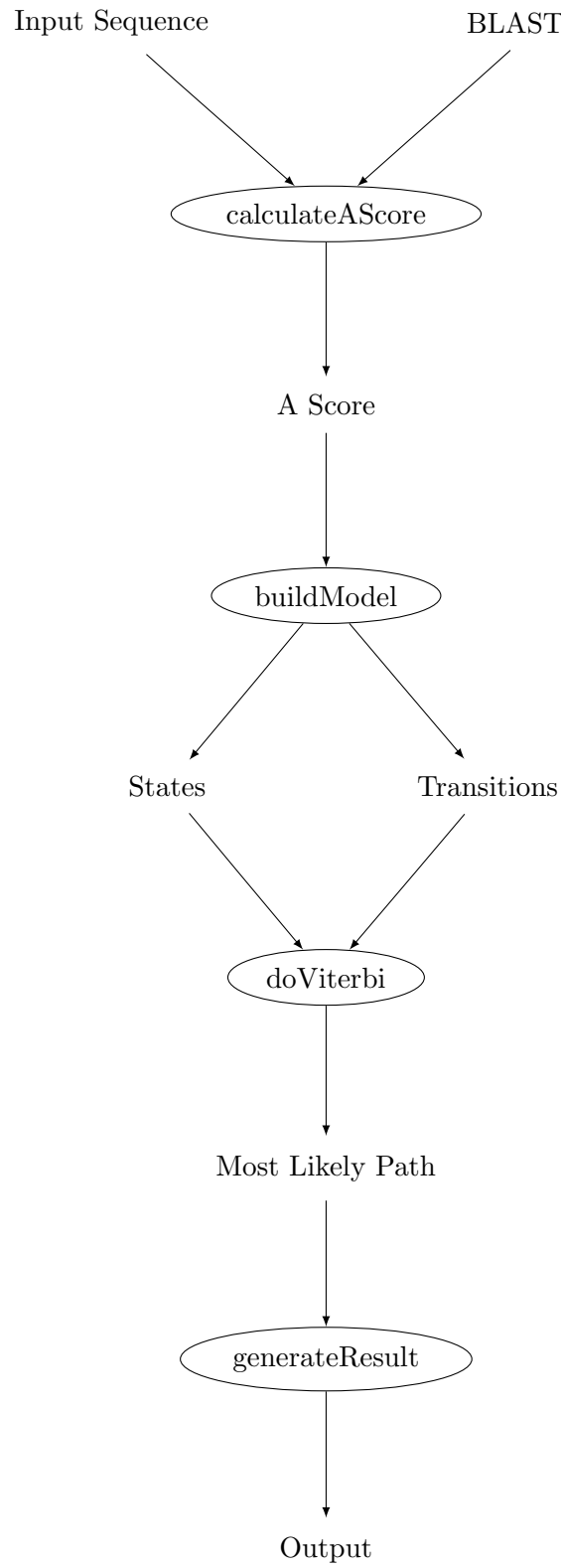
4.3.4 Running Viterbi

4.3.5 Printing Results

4.4 Approach

Talk about agile here.

Figure 4.2: Overview of the Model Solving ‘Pipeline’



Chapter 5

Maintenance

Talk about how I attempted to improve maintainability/readability here

5.1 Revision Control

Explain how revision control will help future maintenance

Chapter 6

Performance

Talk about how I set off to improve performance here

6.1 Blast Lib

Chapter 7

Evaluation

This chapter is mainly provided for the purpose of showing a typical thesis structure. There are no more thesis requirements described.

7.1 Results

The result of this work is the present document, being both a \LaTeX template and a thesis requirement specification.

7.2 Discussion

The Dual function of this document somewhat de-emphasises the primary purpose of the document, namely the thesis requirements. It would be better, if these could be stated on a few concise pages (cf Appendix 1, p??).

Chapter 8

Conclusion

A thesis requirements/template document has been created. This serves the dual purposes of giving students specific requirements to their theses — both style and content related — while providing a typical thesis structure in a \LaTeX template.

8.1 Future Work

Extract the requirements from the template in order to have very concise requirements.

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