A Toolkit For Reporting On Metagenome Assembly Quality

Final Report for CS39440 Major Project

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Abstract

Metagenomics is the study of environmental samples of genomic data where the contents of the data are potentially unknown and unclear. There is a need for a quality analysis toolkit in order to help researchers avoid analyzing and interpreting bad data, leading to wasting time and money.

The project has designed and produced a prototype toolkit (Khimeta) that aims to meet this need and report on the quality of metagenome assembly data. It provides a user with break downs of the contiguous reads within their data, the GC content of windows of each contiguous read, what Open Reading Frames can be found in them and comparisons between areas of high GC content and Open Reading Frame locations.

It has been tested on artificial and real metagenomic data to determine the usefulness of the prototype, and a number of areas of additional functionality have been identified for future development. This report serves to discuss the analysis, design, implementation, testing and evaluation of the project.

CONTENTS

1	Bacl	ground & Objectives 1
	1.1	Introduction
	1.2	Background
		1.2.1 Metagenomics
		1.2.2 Understanding Quality
		1.2.3 Existing Software
	1.3	Analysis
	1.5	1.3.1 Quality assessment
		1.3.2 Quality report and data input
	1.4	1
	1.4	
		1.4.1 Scrum
		1.4.2 Extreme Programming
		1.4.3 Pomodoro Technique
		1.4.4 Project blog
_	ъ.	
2	Desi	
	2.1	Overall Architecture
		2.1.1 Choice of technologies
		2.1.2 MVC Framework
		2.1.3 Naming Conventions
		2.1.4 User Input
		2.1.5 Directory Structure
		2.1.6 QualitySummary and Results
	2.2	User Interface
		2.2.1 Welcome
		2.2.2 List
		2.2.3 Results
		2.2.4 Error
	2.3	Support Tools
	2.5	2.3.1 Version Control and Continuous Integration
		2.3.1 Version Control and Continuous integration
3	Imp	ementation 34
	3.1	Development Environment
	3.2	Features
		3.2.1 Reading User Input
		3.2.2 Counting GC Content & Percentage
		3.2.3 Displaying GC Content percentage
		3.2.4 Finding Open Reading Frames
		1 7 6
		3.2.6 Superframe Comparisson
		3.2.7 Implementation Review
4	Test	ng 46
-	4.1	
	4.1	
	4.∠	Automated Testing

Annotated Bibliography							
E	Defi	tions of Technical Terms	73				
D	Syst	m Test Table	68				
C	3.1 3.2	ples and Extras Example User Story Breakdown					
В	Ethi	Submission	61				
		1.1.3 Jquery	. 59				
		1.1.2 CSS					
	1.1	Tabs for results					
A		-Party Code and Libraries	57				
Αŗ	pend	es	56				
	5.4	Final Conclusion	. 55				
	5.3	Project Management					
	3.2	5.2.1 Future Work					
	5.1 5.2	Requirements Analysis					
5		ation	52				
		4.2.3 Manual Testing	. 50				
		4.2.2 User Interface Testing					
		4.2.1 Unit Tests	. 46				

LIST OF FIGURES

A very small example of a number of reads being taken into an assembler and the output of a contig where there may be a chimera.	3
A small example of the contents of a FASTA file, with very short contigs. The example is just to show the format of how multiple contigs are stored in the same	
FASTA file	7
A use case diagram demonstrating the expected functionality for a user based on	
the functional requirements laid out in this section.	9
Adding an object to the Model via the Controller to be accessed in the View. The View here is 'list', which is a Thymeleaf template that will dynamically build the	
Including a thymeleaf fragment in a page. You can see how the call is made from	14
· · · · · · · · · · · · · · · · · · ·	15
	10
	15
The MVC framework the application is designed upon. The data flows between	
the Model and the View via the Controller, based on the user requests and interaction.	16
Class diagram for the 'Welcome Page', and what classes are used upon a Request	
for the page	17
	18
	10
	19 22
	22
	23
	23
	24
alter the parameters of the quality assessment inspection	25
If the user clicks to see an explanation of the process, they can click the text that	
says to 'Show explanation', and this text appears	25
	27
*	28
	20
	29
· · · · · · · · · · · · · · · · · · ·	29
	29
· · · · · ·	
finishes, if they wished to inspect the contig themselves using those numbers	30
	output of a contig where there may be a chimera. A small example of the contents of a FASTA file, with very short contigs. The example is just to show the format of how multiple contigs are stored in the same FASTA file. A use case diagram demonstrating the expected functionality for a user based on the functional requirements laid out in this section. Adding an object to the Model via the Controller to be accessed in the View. The View here is 'list', which is a Thymeleaf template that will dynamically build the page using the data from the Model added here. Including a thymeleaf fragment in a page. You can see how the call is made from one 'th:replace' with the name of the html fragment file and then the fragment to be included from that file, in this case 'inspectbox'. Including a thymeleaf fragment in a page. This is the declaration of the fragment being included in the previous figure, declared as such through 'th:fragment'. The MVC framework the application is designed upon. The data flows between the Model and the View via the Controller, based on the user requests and interaction. Class diagram for the 'Welcome Page', and what classes are used upon a Request for the page. Class diagram for the 'List Page', and what classes are used upon a Request for the page. Class diagram for the 'Results Page', and what classes are used upon a Request for the page. Class diagram for the 'Results Page', and what classes are used upon a Request for the page. The layout of the directory structure for the application. The 'welcome' page that greets the user upon requesting the home page of the web service. The 'list' page that is displayed after a user submits their (valid) assembly data in FASTA format. When a user clicks to inspect a contig, this menu appears giving them options to alter the parameters of the quality assessment inspection. If the user clicks to see an explanation of the process, they can click the text that says to 'Show explanation', and this text appears. The GC result tab shown to th

2.18	If an error occurs as the user is browsing the web service, i.e. they try to access a page when they do not have the required data submitted, they are presented with this page informing them of an issue	31
2.19	The Jet Brains IntelliJ Integrated Development Environment (IDE) I used for developing my application.	32
2.20	A countdown of the pomodoro tomato timer on http://www.tomato.es/, used for breaking up development time and keeping track of work done over time.	32
	A few of the commits made to my major project repository	33
3.1	Part of the code for reading in a users data when they have pasted it into the text area of the web service. Deals with creating new ContiguousRead objects and adding them to a ContigResult every time it finds a new header for a contig (or reaches the end of the input)	35
3.2	The code for splitting a contig into windows available for calculating the GC content window percentages	36
3.3	The code for calculating the GC content percentages for each window passed into the method	36
3.4	Extracting the data from the Model, provided by the Controller to the View, using Thymeleaf's inline tag to be able to use CDATA to convert the Thymeleaf extracts into Jove Script	37
3.5	into JavaScript	38
3.6	Finding all of the start and stop codons from within the passed frame from the contig.	39
3.7	Extracting each frame and calling to run the ORF Finding process	39
3.8	Getting the base pair characters of a reverse frame is as simple as a switch statement and building the reversed contig from back to front	40
3.9 3.10	'Zipping' together ORF Locations from start and stop codons within the frame. Finding if a click if within an ORF Location is as simple as going through the list of ORF Locations, only checking against those within the same frame as the click, then looking at whether the click is within the start and stop points of an ORF Location within the canvas, based on the size and location of where it was painted (for reverse frames, the start and stop points are swapped, as reverse frames are displayed in the same direction as forward frames, with the indexes in the right	41
3.11	order for a reverse frame)	42
	displaying ORF Location information within the page	43
4.1	An example of a unit test used in my application.	46
4.2 4.3	Results of running the set of unit tests developed for the application	47 47
4.3	The test coverage of my unit tests over the developed application code, broken	
4.5	down for each class	48 48
4.5	YSlow report after running on the Welcome page	49
4.0 17	VSlow report after running on the List page	40

4.8	YSlow report after running on the Results page	49
4.9	Combining two species contigs together at 50% of each of the file (first half one	
	species, second half another species), we see a very obvious split in the GC chart.	50

Chapter 1

Background & Objectives

1.1 Introduction

The project aim was to create an application that could report on the quality of a metagenome assembly provided by the user, presenting them with feedback about the contiguous sequences contained in their data. The requirements for the project topic were very open, as there are multiple techniques for attempting to report the quality of a single species genome assembly, and while some can be used for metagenome assemblies, it was believed that no one tool covered this area yet with numerous techniques. Likewise, the way in which the results could be presented to the user was not established and open to my own interpretation. There are definitions for technical terms in appendix E.

1.2 Background

Before the project began, my knowledge of metagenomics was very limited, close to none. However, I liked the project title and description and thought it would be an interesting and challenging topic to learn new domain knowledge, use different technologies and attempt to implement an application where I had to learn from the ground up. On top of this, I find DNA to be an intriguing topic even with my limited knowledge and I was curious to learn more as I worked on this project.

My first step was trying to understand what exactly it was I was expected to produce at the end of the project. As the requirements of the resulting application were so open, it was up to me to research what metagenomics is, what is meant by 'quality' within the subject, how this quality might be found and reported on, what technologies would be appropriate and what quality techniques could be used.

1.2.1 Metagenomics

Metagenomics is the study of environmental samples of genomic data where the contents of the data are potentially unknown and unclear. It has been described as 'Open-ended sequencing of nucleic acids recovered directly from samples without culture or target-specific enrichment or amplification; usually applies to the study of microbial communities.' [12] It can be used in the

findings of what an animal gut may contain, what viruses are within a sample when looking into outbreaks and finding what microbial communities exist in a sample area.

Metagenomics is a hot and interesting topic in the bioinformatics field, and its uses grow as more is learned, but there is the issue of quality, and how a metagenomic sample should be processed. To help me better understand the project task, I read articles that attempted to provided ways of analyzing metagenomic data to get the best quality results at the end. [26]

1.2.2 Understanding Quality

Considering the nature of metagenomics and the unknowns, it becomes clear quite quickly that when taking a metagenomic sample from sequencing to assembling there can be a number of areas where errors can be introduced. Without the proper tools to quality assess these contiguous sequences a user cannot be sure if what they are creating is an actual thing that could exist in nature, a part of a genome or a chimera.

The process of taking a sample through to sequencing with metagenomic data can be very error prone, leading to misassemblies with duplicate or short reads, or combining reads together to make chimeric contigs. [10] When we take a sample, it gets chopped into short pieces that are then run through a sequencer. The sequencer will output many short 'reads' (between 50-100 base pairs of DNA). Within these reads there many be many overlapping copies, and so when an assembler is used to put the reads together, overlaps are combined to create longer contiguous sequences. A perfect assembly of a single organism could produce a full genome sequence. However, with metageomics it is likely that there will be many contigs, of which some may be genomes and others may be smaller pieces of genomes or even pseudo-genomies made up of chimeric data.

A chimieric contig is an instance where an assembler has put together reads from a sample that it believed were part of the same whole, and yet were in fact of different species/sub-species, and so creates a contig that does not actually exist in nature. It can be understood then that if a user were to try and analyze and use this, unless that is the result that they wanted, it would be unlikely to produce useful or meaningful results. Without good quality assessment tools, how would they know that their assembly data contains chimeric contigs and are not just wasting their time and money?

We can visualize this through the diagram below, where we take a number of reads, put them through an assembler and the output is a number of contigs that the assembler believes it has created correctly, but may in fact be chimeric.

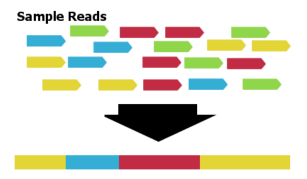


Figure 1.1: A very small example of a number of reads being taken into an assembler and the output of a contig where there may be a chimera.

If we imagine each of the colours representing an individual species (though in reality, there would be no colours, only the characters of data, and potentially hundred to thousands of different species and sub species in a sample) we could, by eye, see how we could extract some interesting genome data, for example taking all of the green in the order they are presented and finding the 'green' genome. As naive as that is, it serves then to demonstrate that if we look at the output of the one contig shown in the figure, we can see how the colours are aligned in a way that they are mixed.

It is possible that the resulting contig is a valid genome that could be found in nature, where the particular sequences in the different coloured sections are shared between each colour 'species' and so this contig could work. However, it is also likely that the different coloured sections are widly different than the natural connecting colours if instead this contig was made up of a single colour (in this figure, all yellow, all blue, etc). This would then be a chimera. This demonstrates that without some quality assessment tools to attempt to report on where these misaligns have occurred, a user may never know if what they have is good quality data or not.

1.2.3 Existing Software

There are a number of tools I discovered in my background research that attempt to aid in the quality assessment of metagenomic data, in particular MEGAN, a 'next-generation metagenomic data, called MEtaGenomeANalyzer', which attempts to do a taxonomical analysis comparison to known reference data [8] and PRINSEQ, a tool which provides 'summary statistics of FASTA (and QUAL) or FASTQ files in tabular and graphical form' [20].

The purpose of MEGAN is to look at user uploaded data sets and compare their data against numerous reference databases of known sequences, and then take the resuling comparison data and display it to the user in an interactive analysis visualisation. it uses a number of techniques to format the users data to check against the reference databases to maximize useful results. The aim of MEGAN is to try to determine what is in the sequences the user provides, if they exist already and how they compare to known reference datasets.

PRINSEQ is a web service (or standalone) tool written in Perl that lets users see statistical analysis data of their metagenome sequences through uploading FASTA files. It allows them to trim, filter and reformat their sequences, and a user can upload files of up to 600MB. Its aim is similar to the task set out for this project, in that it wishes to reduce the amount of time and money

wasted on analyzing and interpreting poor metagenomic data.

When considering what my own application should do, I looked at the techniques used for PRINSEQ most, as these seemed to match up with what I thought would be useful to a user for my own application, and from discussing the topic with my supervisor I found techniques such as the GC Content distribution could be a good place to start.

It was not just tools for metagenomic quality assessment I looked into. I also found the NCBI database and their BLAST tool [6], and kept in mind these may be useful as I progressed through my applications development.

BLAST (Basic Local Alignment Search Tool) allows user to search databases for different sequences and find regions of similarity between biological sequences. A user could upload their own sequence and have it aligned against known sequences to see if it already exists in the reference databases. This could be useful to be used in a metagenome quality toolkit, similar to MEGAN, through letting the user know that their assembly data has some similarities to existing sequences, which might indicate good quality.

Through reading an article that discussed the advantages of k-mer frequency analysis for quality assessment, I looked into the Jellyfish [14] and BFCounter [15] tools for just this role where I could potentially consume the output of their processing, although they took a step back in my mind while I considered what it was I actually wanted my application to be and worked out the requirements.

Jellyfish is a k-mer counting application that runs on the command line and uses the CPUs 'compare and swap' function and encoding with a hash table to increase efficiency. On the other hand, BFCounter uses a Bloom Filter (a probabilistic data structure) to count k-mers, counting those k-mers that appear more than once and ignoring the singleton k-mers, as they are most likely not important to the data. Both take different methods for counting k-mers, and find their own ways for efficiency. If I wished to implement some kind of k-mer frequency analysis tool into my application, I would need to look into efficiently counting k-mers too.

1.3 Analysis

After understanding the project topic and problem a little better, I decided upon a number of requirements of the application to begin with. Some of these were definite goals, some stretch goals and some future development tasks if I were to finish all else or were to continue the application after the project deadline. I broke the problem down into its two core components steps, the analysis and the report. I felt that the resulting list that can be seen below was enough to work with based on the knowledge I had gained from my background reading and what I thought would be appropriate for the time allotted for the project.

1.3.1 Quality assessment

The quality assessment had to use a number of methods suitable enough to produce some data or statistics that could indicate to the user a measure of quality of their assembly data. For this end, I decided upon a number of objectives:

1.3.1.1 Contig length

FR1 'Give the user control over the minimum length a contig should be to be considered.'

This measure would be good in displaying where an assembler could not find any reads to match with a single read and so didn't do anything more with it than output it as a contig of an individual read length. This would most likely indicate that it is of no use to a user, as a contig the size of a read length is unlikely to contain any useful genome data. Allowing a user to set the minimum length threshold lets them set their own size, be it the known read length size of their data, or a size they think would be appropriate to start seeing some usable data from contigs with length over a particular amount/number of read lengths.

1.3.1.2 Number of unknown characters

FR2 'The application should count the number of unknown (N) characters within a contig.'

When an assembler cannot understand what to do with a character, or a sequence of characters, it may insert an 'N' character. Indicating to a user how many of these exist in a contig, and what percentage of the whole they make up is a helpful indication of whether their data is of good quality or not, with the less or no unknowns the better the quality.

1.3.1.3 GC Content

FR3 'Conduct a GC Content percentage analysis in sliding windows of a size set by the user.'

The GC content is the percentage of 'G' (guanine) and 'C' (cytosine) characters within a sequence [22]. When the application takes a contig, it should break the contig up into windows of a size set by the user. If a contig was of length 30,000, they might break it into 100 windows of 300, for example. These window sizes should then have their GC content percentage calculated, and the mean of the entire contigs GC content worked out. Using these values it then becomes possible to detect potential anomalies in the percentages of individual windows that have a percentage value drastically different than the mean.

1.3.1.4 Open Reading Frame Locations

FR4 'ORF locations should be highlighted to indicate whether a GC window is outside of the mean because it aligns with an ORF location rather than being a chimeric change.'

This addition to the requirements came later in the projects development as I came to understand GC content and how a window that looks like an area of a chimeric change may actually not be. We can look at a contiguous sequence and find Open Reading Frame [24] locations, where the GC content percentage is often naturally different than outside of these regions. [27] This is due to ORFs being subsequences of DNA that may encode proteins, where the GC content is often higher.

An ORF Location is found through looking for where a location starts with ATG and where it ends with TGA, TAA or TAG. There are 6 frames to be found in sequences, where three are 'forward' and 3 are 'backwards' with the opposite characters of the base pairs to the forward frames. For both forward and backwards, the first frame is the original sequence, the second

frame begins one character into the sequences and the third frame begins two characters into the sequence. A very small example just for basic understanding is provided below:

```
Sequence:
TTGATGGCGCATAG

Frame \#1 (fwd):
TTGATGGCGCATAG
Frame \#2 (fwd):
TGATGGCGCATAG
Frame \#3 (fwd):
GATGGCGCATAG
Frame \#4 (bck):
CTATGCGCCATCAA
Frame \#5 (bck):
TATGCGCCATCAA
Frame \#6 (bck):
ATGCGCCATCAA
```

By finding the ORF Locations, we can use these in the quality report to match up to windows that may at first seem like anomalies in the contig where it is instead actually a natural occurrence. There is a preexisting tool for this ORF finding functionality by NCBI [23]. While it might not be the case that an ORF Location matches with an out of threshold GC content window, or that they do match but it is still in fact an anomaly, it was a requirement of the application I wished to include in order to give the user another tool to use for inspecting their data.

1.3.1.5 K-mer frequency analysis

FR5 'The application should conduct a k-mer frequency analysis, or use output of k-mer frequency analysis tools.'

Conducting k-mer frequency analysis in window sizes has a similar output as GC content, except we look for the frequency of particular 'k-mer' (where 'k' is a number of how many characters to be considered, e.g. 3mer 'ATG', 4mer 'ATGA'). Through measuring the frequency in windows we could see if there was an even distribution of frequencies across a contig, and if any windows had large changes in the frequencies that could indicate a potentially bad quality contig. [11]

This was set as a stretch requirement, as the process of efficiently conducting k-mer frequency analysis and the research into the possibility of writing my own software to do it or consuming output of another application was expected to take over the time for the project, after my background reading and understanding was completed and the other features previously mentioned implemented.

1.3.1.6 NCBI reference data

FR6 'The application should compare the contigs to known reference data in the NCBI database with BLAST.'

Using the known reference data, much like with MEGAN, it would be possible to see if any of the contigs in the user data match to known reference data. If a full contig matches a species with BLAST, perhaps it is of good quality and demonstrates a genome. However if a contig matches one species from BLAST in one half of the data and the other half matches another species from BLAST, then we could conclude that this contig shows evidence of being a chimeric contig. This was a stretch requirement though, with the idea more in mind for if the application were to be continued to be developed after the project deadline.

1.3.2 Quality report and data input

1.3.2.1 Read user input in FASTA format

FR7 'A user should be able to paste or upload their data in FASTA format.'

The FASTA format is a commonly used format for sequence data. It is 'a text-based format for representing either nucleotide sequences or peptide sequences' [28]. It is formatted to have a header of single line description, containing information about the way the data was created, and then lines of sequence data afterwards. An example of this can be seen in the figure below. I set myself the task for allowing the upload and paste of data, beginning with pasting the data into a form, as uploading data could add a larger realm of security and data size issues and so I wanted to have the simplest thing possible first.

```
>NODE 1 length 180 cov 7.511111
CTCCCTCTTTTTTCGGATATGCTGGTGATTTGGCGACGAACAGTATCTGGGCAGCTCTAT
GGTTTGCCAGTTTCCTATCTGGTCGGATAAACTAGTGATTTGGCGACAGGCGGTATCGGG
TAAGCTTGTTTTATCCCGACGATCAGTGCCTGAAATACTGACAGATACTG
>NODE 3 length 74 cov 9.391891
ATTAAAAAGGGTGTAAATTGAATATTATAATATGTATAGGAGTCGGATAGTCGTGACCTA
\tt GTTTGTGTCGTCTAACTGTTGACGAGAAGCGTGTAAAGAGATCGCGTGCTACACAA
>NODE_9_length_70_cov_41.885715
TAATATTGTAATTATTTTGAAATGTTTAATTTGGGTTTAAATTTGTCAATAAAATATTGT
>NODE 16 length 154 cov 11.662337
TGCCTATTACAGGTCTTTCTTTAGGCCTGAAATTTTATCTAAATTATTAAAGATACATTA
ATTAACATATCCAGACCATTAATGAACTCTCGATGTCTACACTAGCCTTTGATATTGACT
GATAATTAAATCAAAAAATGGATAATCGAAACTATGCTTGTTATTATACTGACTTTATTA
ATTTCTATTCCTTAATTCTTGGTt
>NODE_23_length_138_cov_4.391304
GTACCATCCAAAGTACTAGACATTGTTTCTTGTCAGACCCAATTTTACATCACTGATATG
GGGAAAGCAGATTAGAAAATTGGTGAGGACCTTTTCTCTGCCCATAGAGGAAAGGCAACC
GGGACCTTCTGGTAACTGTGCACCCCACCCTCCTCTTTACTTATTAaacaggcatctac
taagttga
>NODE_32_length_78_cov_4.871795
TCCGAAATTCGAACTCATCCTTCCTTAGGGTGTAACGGTGATATAATCAAACGATTTTTT
AAATCGGa
```

Figure 1.2: A small example of the contents of a FASTA file, with very short contigs. The example is just to show the format of how multiple contigs are stored in the same FASTA file.

For the FASTA format my application accepted, I planned for it to begin only taking into account ATGC & N as characters to deal with, where N was 'unknown/unclear' and the other characters were valid. Uppercase and lowercase characters would be preserved in any output display of the application, but capitalization would not taken into account for any of the processes as they did not make a difference to the quality assessment techniques.

1.3.2.2 Display a list of the contigs

FR8 'A user should be able to see the contiguous sequences listed in their FASTA data.'

I decided it would be a good idea to display to the user the list of contigs that their file contained, with small bits of relevant information, such as the size of the contig, the header of it and the previously mentioned 'N' count and percentage. From here a user would be able to select a contig they wish to further inspect for quality issues.

1.3.2.3 User control over GC Content, ORF Location and k-mer frequency analysis parameters

FR9 'When conducting a quality inspection, a user should be able to adjust the parameters of the methods based upon their expectations, data input and choice.'

Since the data uploaded by a user would have been assembled from reads of a particular length, the appropriate size of GC content and k-mer frequency windows, minimum size of ORF Location lengths and minimum length of contigs should be set by the user. It could have been a possibility to guess what a good size would be for the user (i.e. find the shortened contig in the data and consider that that might be the size of a single read, and suggest length parameters based on that).

1.3.2.4 Visual reports of the quality inspection methods

FR10 'A user should be able to see a visual representation of a contigs GC content and any anomalies.'

In order for a user to be shown whether their data is of good quality or not, visualizations would be required that highlight any problem areas, or if areas are not highlighted, at least give them a broad sense of the layout of the results of the quality inspection to allow them to analyze the results from themselves. This should include displays for GC content in windows, ORF Locations broken into individual frames, k-mer frequency analysis in windows and a comparison shown between the GC content and ORF Locations.

I did consider how I could potentially provided a quality confidence factor to a user, or how I might demonstrate to them in a statistical way that their data was of good or bad quality. However, with the amount of methods that could be used, the unclarity of the data itself and the users expectations, I felt that it would be better to instead provide the user with visuals to analyze the results of my application themselves, and I would instead try to highlight problem areas without necessarily saying clearly 'This is/is not a problem area'.

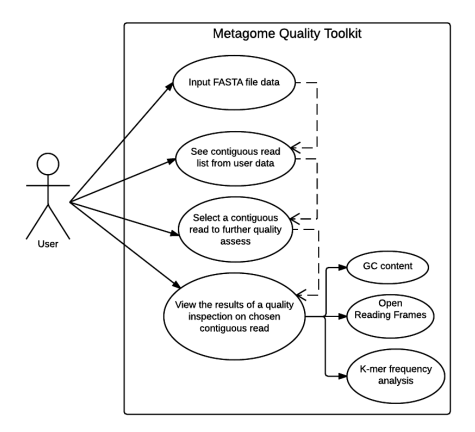


Figure 1.3: A use case diagram demonstrating the expected functionality for a user based on the functional requirements laid out in this section.

1.3.3 Implementation - 3rd Party vs Self-Created

Based on the requirements I had selected, I decided that I wanted to write my own software for each functional component. While solutions for each component exists individually, I felt there would be more benefit to a user if I could produce my own software to support the functionality, over requiring them to use third part software in order to use my application too.

Additionally to this, I wanted the technical challenge of writing my own software, and enabling the application to be maintainable and expanded upon in the future rather than relying on third party support applications in order to process a users input. The process of developing the application from scratch would also give me the opportunity to learn more about the domain and technologies for development than just consuming output of other existing applications, even if achieving the end result would take more time.

While I knew the project would come to a close on development at the end of the semester, I also wanted to envision this as a project that could be taken forward in the future to be further developed and maintained. For this reason it also made sense to develop my own application without consuming output from third party software, excluding the FASTA files from an assembler. If one of these third party applications was no longer available or their output changed, it would render my application unusable or require more future modification to adapt to changes by outside sources.

1.4 Process

An agile methodology seemed most appropriate for this project development, due to the unclear requirements (based on my lack of understanding and the open tasks at the start of the project) and would allow me to break down each requirements into stories, and then into tasks and get a thin slice of the application done quickly, and build outwards from there, always ensuring I had some finished product at each functional completion. While my project could well follow a plan driven approach, such as waterfall or the spiral model, I believed that an agile approach would be more suitable for developing it.

I chose to use Scrum for my framework, and used aspect of Extreme Programming for my daily development cycle, adapted to a one person project. Considering the nature of my projects requirements, I considered if I should instead use Feature Driven Development, but decided upon Scrum due to my experience with it through my industrial year and in that I could see how I could break the requirements down in to user stories and tasks.

1.4.1 Scrum

Scrum is an agile framework for developing complex products [21]. In brief, it works in iterations (Sprints) where work is broken into user stories and tasks for those stories, there are key members of the team who have designated roles (Engineer, QA, Scrum Master, Product Owner, etc) and where each Sprint has meetings involved in it to help the flow of work: Sprint Planning & Sprint Retrospectives, Sprint Review and Daily Stand-ups. I decided that adopting this framework would help me structure my weekly work and aid me in planning and development for the project.

As a one-person project, a number of the components of Scrum either had to be dropped or adapted, for example, the roles of team members were all adopted by myself rather than having actual team members. I worked on a weekly Sprint cycle, where at the beginning of every Tuesday a new Sprint would begin, and I would review the work done and have a Retrospective with myself and plan out the next weeks tasks, often through talking with my supervisor. As for Daily Standups, I was able to carry these out through holding them with my peers, Alex Jollands and Sion Griffiths. Between the three of us we would meet in-person or using Internet calls on weekday mornings at 10am and discuss our work from the day before, the planned work for the current day and anything we wanted to soundboard off one another.

The Scrum framework had me taking my initial planned out requirements and turn them into Stories with their own tasks. Stories are structured in the way of:

```
'As a <type of user>, I want <some goal>so that <some reason>.' [17]
```

Once a user story has been created, I would then consider it and see if I could break it down into a 'narrow slice'; the simplest, thinnest possible thing I could do for a feature implementation that would produce results. There is an example of this in Appendix 3 'Examples and Extras' under 'Example User Story Breakdown'. Breaking down the requirements into user stories and then tasks helped me focus on what was important, and gauge what needed to be done and how much effort vs time it would take.

1.4.2 Extreme Programming

For the day to day development outside of the Scrum framework, I decided to adopt a modified Extreme Programming (XP) approach [16]. While I couldn't do all of the XP techniques (pair programming, for example), I strongly took the elements of Test Driven Development (TDD), refactoring, simple design (coding only what needs to be done based on the TDD and refactoring) and continuous integration. I felt that using a combination of Scrum and XP techniques was a suitable process for my application, and would aid me in my design, implementation, testing, meeting the requirements and focusing my motivation.

The motivation for taking the aspects of TDD and Refactoring were that I could build and design my application iteratively, as I understood and refined the requirements and functionality of the application as I went, while still having a usable application from an early stage that could be developed upon in each Sprint. This helped me in having an evolving design for the application, while not having a design document but still laying out the structure of the application to a good standard.

TDD ensured that the application would be developed in the simplest way possible, and always meet the requirements currently tested for by doing the unit test development up front. Refactoring the work as I went and where needed meant that I would keep my code maintainable while developing in this way. The end result through selecting this methodology was to aim for a robust, maintainable, well-design application with test coverage for all functionality and programmed efficiently through proper time management.

Using continuous integration meant that I could be sure that whenever I completed a new bit of code, it would be run against all existing tests to check that nothing had broken and was impacting previously working functionality.

1.4.3 Pomodoro Technique

On the matter of time management during development time, I found it useful to break my work into small cycles using the Pomodoro technique [3] to increase my focus and productivity. The aim of the Pomodoro technique is to spend 25 minutes with full focus on the work, and then spend 5 minutes to stop and have a break. They are also useful for keeping track of work done and visually seeing how much time has been invested into the project for the day. To record my pomodoros I used http://www.tomato.es/.

Due to the short length of each pomodoro, it was also seen as an effective method for squeezing in work where there was time. By quantifying time into small slots, it made it possible to consider how much time I might have between other events in the day and work out where it could be possible to put in a single pomodoros worth of work. This is instead of perhaps not working because I felt I did not have the time, or working and eating into other daily activities if I didn't block out the time and be aware of when to stop.

1.4.4 Project blog

While developing my application I kept a blog about my project, any milestones I reached or interesting sections. While I did not designate a particular time to blog, I believed it would be a useful tool in reflecting on the work done when writing this report, something for my supervisor

to be able to check upon my progress between meetings and give me the time to reflect on the work done and any upcoming tasks to help me mentally process where the project was in its life cycle against the planned objects. The blog may be seen at http://users.aber.ac.uk/jee22/wordpress/.

Chapter 2

Design

2.1 Overall Architecture

The design of the application evolved over time due to being developed using agile methodologies. This meant that each step had an initial design and was built upon every time a new feature was added, being refactored along the way. There were a number of prototypes made during the course of the project's development, and this chapter will discuss the design of the end product, and go into detail where the design may have been previously different.

The resulting application evolved into a web service, with a Model View Controller (MVC) framework and resources structured for maintainability. Following XPs guidelines, at each step of the way the applications design was made as such to be the simplest yet most maintainable it could be through refactoring, cutting down on duplicate code, structured logically and choosing smart data structures and Objects to represent aspects of the application and quality results.

2.1.1 Choice of technologies

At the beginning of the application, I felt that the application should be programmed in a language that would be able to be ported across multiple platforms for used by anybody who wished to use it. Additionally to this, I wanted to have ways of representing the components of my application as Objects and would need some way of presenting a UI to the user, even if my initial application would only output command line results, in order to follow the XP value of Simplicity (YAGNI - "You Ain't Gonna Need It", until you do).

For this reason, I selected to develop my application using Java to begin with as it filled the criteria of being Object Oriented and was portable through using the JVM on different operating systems. I resolved that I would select a UI package to present the report of my results once some of the core functionality had been implemented. I considered instead using Ruby or C++, but due to my familiarity with Java I believed it would be a better choice to stick with what I knew. At this point, I did not consider having the application as a web service, and so using Ruby with Rails was not something I had thought of.

As the application developed and I started reading in files and outputting results from the GC Content process, I found that I was having a hard time finding a quickly usable GUI I could work with for Java to display the results in a way that I wanted. It was at this point I started considering

a technology change to Ruby on Rails until Sion Griffiths, a peer of mine, suggested I could keep my application in Java and turn it into a web service using Spring Framework [19], and using Spring Boot to easily set up the application with Tomcat for me [18]. This seemed like the perfect solution to my problem, allowing me to generate my charts and results using HTML5, JavaScript and HTML5 Canvas, along with JavaScript libraries such as Plotly.js [9], a charting library for creating explorable, interactive charts for the web.

Spring is an open source application framework for Java that is highly useful for building web applications. Using Spring Boot makes generating a Spring application quick and simple, taking most of the configuration code for running things like Tomcat servers and helping configure them up for you. I used Maven as my build manager and controller of dependencies. This tied in with using Spring Boot as it managed the pom.xml file used to configure Spring through Spring Boot, helped me install and update any dependencies necessary for building and running my application and produced the executable Jar file for distributing the application. Tomcat is a deployable web server that I use to host my application as a web serice with.

The conversion to Java Spring was not a painful one, and only involved setting up a new Maven project and declaring it to run as a Spring application in the pom.xml file, then copying over my previous code and structure into the new project. From there I could deploy the application as a web server using Tomcat and I was back to my previous position of working out a User Interface solutions. Thankfully, there is a package called Thymeleaf [7] that works with Spring to allow access to Objects from within the Model of the Java code (placed there by the Controller) using the View dynamically as the View is generated through HTML templates and fragments.

```
model.addAttribute("contiglist", contigList);
model.addAttribute("discardedcontigcount", contigResult.getDiscardedContigCount());
model.addAttribute("contiguousread", new ContiguousRead());
return "list";
```

Figure 2.1: Adding an object to the Model via the Controller to be accessed in the View. The View here is 'list', which is a Thymeleaf template that will dynamically build the page using the data from the Model added here.

On top of using Thymeleaf for accessing data put into the Model, it has the additional benefit of using fragments that can be imported into different HTML templates. This design choice made it so I was able to cut down on writing duplicates of code. For example, I wrote the header and footer of the UI design in a Thymeleaf HTML fragment, and then on each page just needed to call one line in order to import it into that page, rather than built the entire thing again. An extra benefit to this is that it allowed me to keep the HTML templates clean and easier to maintain by reducing their size and separating out different aspects of the UI. Having the header, footer and a number of forms in Thymeleaf HTML templates meant that these sections could be edited without impacting any page they are imported into.

Figure 2.2: Including a thymeleaf fragment in a page. You can see how the call is made from one 'th:replace' with the name of the html fragment file and then the fragment to be included from that file, in this case 'inspectbox'.

Figure 2.3: Including a thymeleaf fragment in a page. This is the declaration of the fragment being included in the previous figure, declared as such through 'th:fragment'.

For the GUI itself, I elected to use Plotly.js [9] for representing my GC content charts for the level of detail and control it gives a user over inspecting charts, large and small, that worked no matter what GC window size the user selected. For the ORF Locations frames charts I wrote my own code using JavaScript and displayed it with HTML5 Canvas, as it allowed me control over what should be shown depending what a user clicks upon and I could work with the data from the Model in ways I saw fit.

2.1.2 MVC Framework

As I was building an application with a GUI, the application was built using a Model View Controller framework design, separating out the components into their different types. This helped the separation of code and responsibilities. The class diagrams presented in this chapter are based on the final product design and based on a page per page view from the UI and what classes are used on the HTML request of that page.

Builds and returns view on user request/interaction Data supplied from Model by Controller View Model

Figure 2.4: The MVC framework the application is designed upon. The data flows between the Model and the View via the Controller, based on the user requests and interaction.

2.1.2.1 Model

The Model was designed to contain the data and methods for processing a users input. This contains the data structures and objects for handling a users data as they traverse from page to page of the view. I designed the Model in a way that certain objects would be Bean objects (classes with getters and setters for their properties) that could be accessed by the View in the way that Thymeleaf required.

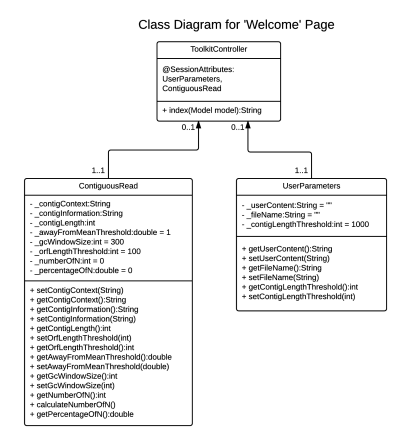


Figure 2.5: Class diagram for the 'Welcome Page', and what classes are used upon a Request for the page.

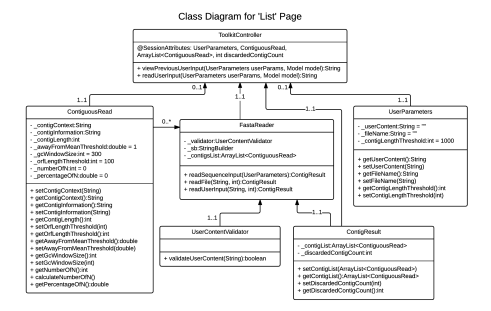


Figure 2.6: Class diagram for the 'List Page', and what classes are used upon a Request for the page.



Figure 2.7: Class diagram for the 'Results Page', and what classes are used upon a Request for the page.

The structure evolved over time due to the nature of XP and using an agile methodology. Even so, the result was that there was a logical sense to the structure of classes, methods belong in the class they live in, some utility classes exist for carrying out operations that don't necessarily need to be within a class that needs that utility and this also left the code easier to maintain and separated. By using utility classes and methods, they can be called by other classes in the future, and the class that uses them doesn't need to know about the implementation internals of the methods they use, only the results.

2.1.2.2 View

The View of the application is where the GUI is presented to the user to be seen in their browser. Designing the application to use the browser was chosen to allow anyone to be able to use the application as long as they are using a modern browser. It is using HTML, HTML5 Canvas and JavaScript, dynamically built using Thymeleaf templates. If any of the template does not process or there is an issue with the users data they are presented with an error page and told to return to the menu and try their task again.

The data for the View comes from the Model, passed by the Controller. In line with keeping responsibilities separated, the Model does not care about how the View displays the data, and the View does not care about the content of the Model data it is passed as long as the data types it expects are present in the Model from the Controller. For example, the ORF Location data is sent to the View as an OpenReadingFrameResult Object, that has a list of OpenReadingFrameLocations and other data about the process. It is then the responsibility of the View to use this data in displaying it in HTML5 Canvas with JavaScript. No additional processing or modifications are carried out on the data at this point, the View just picks up the data and places it on the Canvas element where it expects it should go, based on the content of the data, e.g. an OpenReadingFrameLocation with the 'frameIndicator' set as 0 would indicate to the View that it should be placed on Frame 1.

This type of responsibility separation is consistent throughout the application, with one exception. There is a GcResultDataView, that is the GcResult data processed into an easy to handle class. This is to make the process for the View far faster than having to get the View to carry out some processing to determine the GcWindows that are out of the mean threshold and additional results to display on the bar chart for the GcResult. While it would be possible to alter the application in order to leave this up to the View, I felt that it would be better to instead serve the View with the window data it needed to put on the chart and the colour codes for each of those window data bars. If the View for this result were to change in the future, no additional processing would be required within the Model to match this, as the results are still present in their raw form for the View to use as it sees fit, and so I felt this was an acceptable design choice.

2.1.2.3 Controller

The Controller serves as the master for how data flows through the application, making calls to methods from the Model to process the users data, receiving requests from the user via the View and HTML Requests and handling what data is used in the View. The user may send HTML Requests to the different pages of the application and will receive a response based on the '@RequestMappings' of the Controller.

For example, a GET Request on the 'list' page will return a page expecting the user to have

already submitted some data that has been turned into a list of contigs. A POST Request on the 'list' page however will be expecting that this is a new set of data and process the data submitted into a list of contigs and put them into the users session attributes.

Through every page, a user carries a Session with them and particular session attributes are expected at certain points in @SessionAttributes. For example, a user must have UserParameters and a ContiguousRead in their session parameters in order to be able to view the List and Results page. If they have not visited the Welcome page and submitted data, however, it is not possible for them to have these attributes, and so they are locked out of those pages. This is good behavior as the user should not be able to try and view results for data they have not submitted.

2.1.3 Naming Conventions

I followed a naming convention of attempting to name classes, methods, fields and variables in such a way that a reader could understand what the function of that particular thing was just from the name alone. Private field names all begin with an underscore, e.g. '_privateField' and everything is written in camel case, starting with a lowercase character, e.g. 'thisWouldBeAMethod-Name()'.

2.1.4 User Input

Based on the background reading, the decided file format to accept was FASTA format. The design allows for user uploading and user pasting of files in the Model methods. However, in the end I decided to keep the design just to handling pasting of data. This was based on a time constraint and issues with file upload limits, security and the need to test. While the functionality has been left in the code, it has been left as a 'if I were to continue' functionality to be expanded upon if I had more time to work on file uploading as a priority over different techniques for the quality assessment.

User data is also not kept by the application, it is stored in a user session that expires once they leave the page. This is handled by Java Spring, and set as one of the @SessionAttributes. If the user does not have their session variables, they are presented with the Error page, so they cannot try and access areas of the application/web service where they currently do not have access or the data to do so. This means that the data they have is their own, and not retained by the application or possible for other users to access.

2.1.5 Directory Structure

The directory of the applications structure was designed such that it would reflect the MVC aspect of the application, and make finding resources and particular Classes easier for a reader. The design makes the application easier to maintain in the future and is another way of helping to enforce separation of responsibilities between classes and resource types. In the figure below you can see the way that the directory structure is designed to back this up.

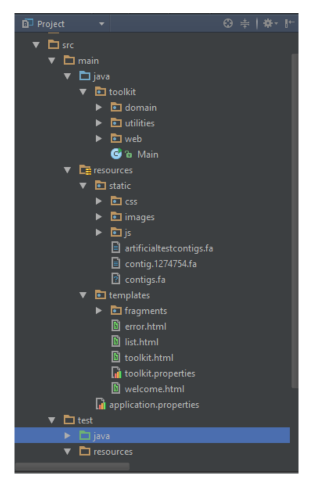


Figure 2.8: The layout of the directory structure for the application.

Within the 'src/main/java' folder, there are the sub folders 'domain', 'utilities' and 'web'. Separating the classes out this way helps with the previously mentioned separation of responsibilities. Domain contains the classes for the domain objects themselves and the data and processing (the Model, so GcResult, OpenReadingFrameLocation, etc), Utilities contains utility classes for use by any of the other classes and to allow their functionality to be independent of any domain class and Web contains the Controller classes (ToolkitController, ErrorController).

2.1.6 QualitySummary and Results

In the application, the quality assessment results are returned in a QualitySummary object. This QualitySummary contains references to GcResult and OpenReadingFrameResult. There is the option to have these two implement an interface that might be called QualityResults, and have just an array of QualityResults in the QualitySummary rather than defined result types. This would allow us to add any type of result without needing to know what is in there. However, I chose not to do this as considering the way in which we serve data to the View it seemed okay to be able to have the View called results from the model directly, and through XP developing this any further would go against YAGNI.

I am aware that in the future if a user was allowed to select what type of processing they wish to

use and as more techniques for quality assessment are added, this technique would be very useful to implement. It would be possible to do this and in the View have it check for the existence of the expected results, or check if they were empty/null, and then only dynamically include fragments of results that are present.

2.2 User Interface

Below is the finalized designs of the user interface, shown through screen shots, along with an explanation for the design choices in how the pages are laid out for the user to navigate through. Each page has a header, containing the title of the application and the logo I made.

The logo is based on the mythical creature 'chimera', a creature made from multiple parts of other creatures (often a lion, snake and a ram). The inspiration for the name and design comes from the application hoping to highlight potential Chimeric regions of contigs to a user, hence 'Chimera - Metagenome - Chimeta'. The 'K' in Chimera is simply because 'Chimeta' sounds too much like it would be pronounced 'chai-meta' not 'kai-meta' as in 'kai-meera' for Chimera. It isn't an important aspect of the design.

2.2.1 Welcome

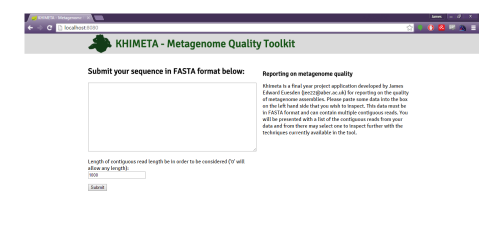


Figure 2.9: The 'welcome' page that greets the user upon requesting the home page of the web service.

The welcome page is structured so that there is an explanation for the user of what the tool is, and an area for them to paste their FASTA data in. There wasn't much else needed for this page, but were the project to be continued this is where there would also be the option to upload user files instead of pasting in data. The user also specifies the minimum length a contig must be in order to be considered here.

The original design for this page also included the parameters for modifying the quality assessment, as this tied in with when the application dealt with only one contig at a time. As the application evolved, the design changed and was refactored to shift the parameters for quality

inspection to the next page where they could be tied to contig and not to the assembly data submission as a whole.

2.2.2 List

The below images show the 'List' page, where a user can see all of the contigs within their uploaded data file that are above the minimum length threshold they specified. The page also displays the number of discarded contigs due to the length, as I felt this would be useful information to give to a user.

For each contig, I also believed it would be useful for the user to see the contigs header (name), the length (how many characters) and the number of unknown 'N' characters, along with the percentage of those that made up the contig. This could be used for a user to consider whether a contig has too many of these characters to the point that they feel it is a bad quality contig.

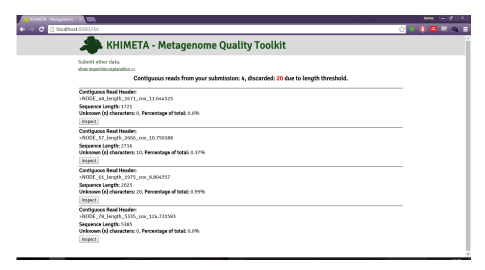


Figure 2.10: The 'list' page that is displayed after a user submits their (valid) assembly data in FASTA format.

If a user wishes to further inspect a contig, I designed it such that when they click to 'Inspect' on a contig, a box appears underneath the contig information for them to modify the parameters for the quality assessment process. I had previously considered a design where these parameters were on the right side of the list, and would stay static there as the user scrolled down the page. However, if a contigs name was very long it would run 'underneath' the box and be hidden. I also found it just did not look very appealing, and so I opted for the design you can see below instead. When the user clicks to run the process the 'Inspect' button changes to 'Loading...', to inform the user that the process is working.

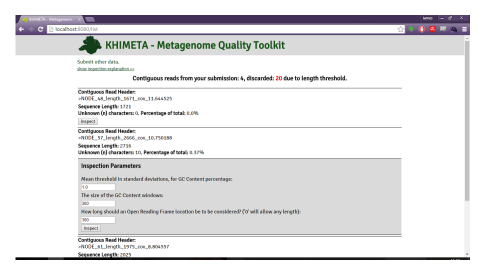


Figure 2.11: When a user clicks to inspect a contig, this menu appears giving them options to alter the parameters of the quality assessment inspection.

In order for a user to understand what the inspection process is, I included a box that details the process, which appears when they click the words for 'Show inspection explanation'. This box can be hidden by clicking the 'Hide inspection explanation'. Currently, the link to display the explanation is at the top of the page, and a user must be at the top of the page to click and view it. It could instead be possible to have the list of contigs scroll in their own box, such that the inspection explanation is always available to be seen and viewed when clicked. However, it did not fall into scope of my Sprints while working on the functionality.

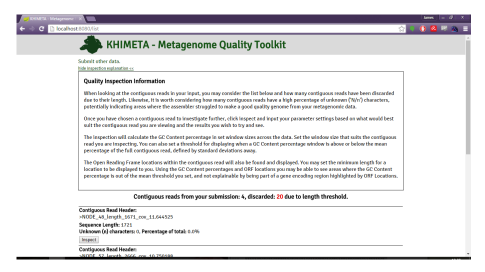


Figure 2.12: If the user clicks to see an explanation of the process, they can click the text that says to 'Show explanation', and this text appears.

2.2.2.1 Parameter choices

The choice of parameters that the user can modify were selected based upon what could be altered about the inspection processes. For example, allowing a user to change the minimum length of

an Open Reading Frame location gives them control over what size they think is important, based upon what their assembly data is. It could be that they care about any sized potential protein coding region, or that they only wish to see ones of size 1000 or above, as they know that anything less is unlikely to be an actual protein coding region. Giving them this parameter choice allows them to decide this for themselves, as the application won't know what is the best size for the user.

Similarly, the GC content window size should be set by the user, again based on what they think is a reasonable size. If a user were to think that they might want windows of size 100, that probably wouldn't return them great results on a large contig, as the GC content could vary so much between windows where the data is just too small to be useful. Likewise if the window size is too big, this data may also be useless, as they cannot properly compare their GC window sizes to the expected size of what they want to see from protein coding regions.

A reasonable window and minimum ORF Location size can only be selected by the user, and so the choice was made to allow the user to select their own choices, even allowing them to select values to high or too low, it is their decision to make based upon the data they have provided. In future versions of the application, it would be interested to infer a decent window size for the user and suggest it to them, but for the purpose of this project it was out of scope for the time allotted with the other functionality to be implemented.

The final parameter is how many standard deviations away from the mean of all GC content windows the user would like to consider is enough to warrant highlighting a window as being a potential issue. I selected standard deviation as it is a standard way of determining a value away from the mean that could be useful to inspect. Whether the user cares how much or how little this value actually is is irrelevant. The user may select what number of these they feel is useful to consider just for highlighting bars in the results chart.

2.2.3 Results

For the display of the inspection results, the user is shown a set of labeled tabs, that correspond to the techniques used. It is designed this way so that more tabs can be added in the future as more techniques are set in the inspection process, and to keep the page from being too crowded with results that would require a large amount of scrolling. The code for making the tabs themselves work is credited to Catalin Rosu [2] and their original code can be seen in the appendices.

2.2.3.1 GC Content

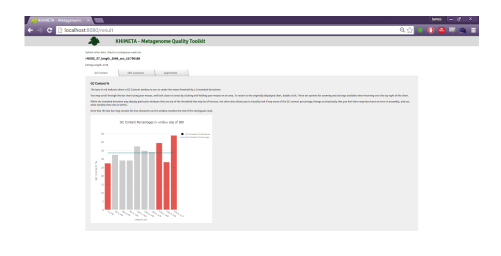


Figure 2.13: The GC result tab shown to the user on inspection of a contig. The browser has been zoomed out in order to display the page in full.

The GC Chart result is displayed to the user in order to give them a visual reference of how the GC content in their contig is. Through this they may detect single windows that may have chimeric properties through a difference in GC content above or below the mean, which can be highlighted in red and are set by being a number of standard deviations away from the mean. It also allows them to look and see if there is a naturally large difference at points between the GC content windows that are not necessarily picked up by the threshold.

The chart is built using Plotly.js [9], as it allowed me to give the user more control over how they view the GC content results in a chart than if I spent the time implementing my own solution to give them the same amount of fine-grained control. A number of the options offered to a user by the Plotly.js chart are probably far more than they need to be and may be confusing to a user. If the project were to be continued it may be worth developing a chart system for displaying the GC content without using Plotly.js, which could also help with developing a way of showing when there are drastic changes in the GC content not caught by the threshold.

Above the chart is a small explanation of what the chart is displaying, and what the user might look for in order to understand the chart and whether there could be potential issues in their contig displayed by these results.

2.2.3.2 Open Reading Frames

The tab for the Open Reading Frame displays a graphic made using JavaScript and HTML5 canvas, that gives the user a visual representation of 6 frames of the contig and any protein coding regions within those frames. The design was inspired by NCBI's orffinder [23].

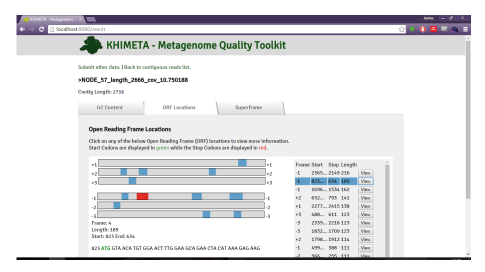


Figure 2.14: The Open Reading Frame results view tab.

I designed it in a way that allows the user to view the full list of ORF Locations (protein coding region) on one side, including their character length, start and stop point and which frame they are in, organized by largest to smallest. This is reflected in the frame chart where a user can see where the frames physically lie within the contig. If a user clicks an ORF Location, either on the chart or in the list, it highlights that ORF in the list in blue, and red in the chart. This also displays more information about the ORF Location underneath the chart.

The choice to display the ORF Location information was out of interest to the user if they wish to use the tool to look at the protein coding regions. The information displayed about an ORF Location includes the length, start and stop point and the characters within the ORF Location. While these might not be useful to the user for determining quality, the information is used in the comparison for GC content to find if the GC content is explained by an ORF Location, and so I believed the information might be useful to display to a user either way.

Each frame has its own canvas element, and they are aligned using a HTML table. I tried a design where each of the 6 frames was made up in a single canvas, and while this worked, it restricted the amount of formatting that could be done around each frame and had much more code for detecting whether a user has clicked within a frame, delaying the response a user got when clicking.

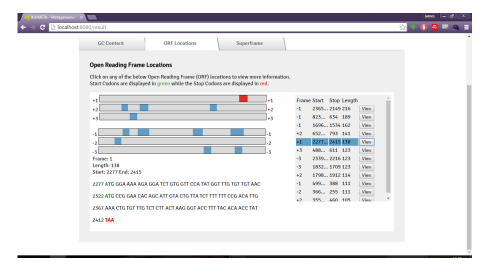


Figure 2.15: The Open Reading Frame results view tab, displaying that when a user clicks on a different ORF Location, the highlight on the frame chart and list changes to reflect this.

2.2.3.3 Superframe

Superframe is where the GC content and ORF Location charts are compared in order to help a user see if there are any areas of GC content difference that might be explained by the ORF Location. It is simply an overlaying of the 6 frames into one chart (all in the same sequence direction) and overlaying the GC content chart results for each window.

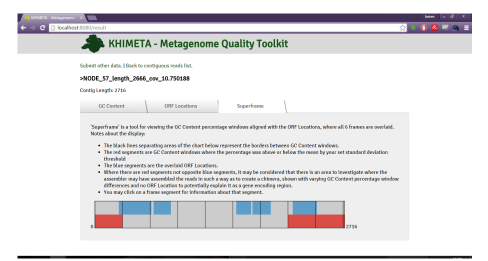


Figure 2.16: The 'Superframe', a comparison between the ORF Locations within the contig and the GC content windows that are above the threshold..

If the user clicks on a particular window, they can see information about what window that is (where it starts and ends) and the GC content percentage. I felt this tool could be useful for a user to be able to determine themselves if they felt there were enough individual windows of GC content that were out of the mean threshold and not explainable by ORF Locations and so felt the quality of the contig was or was not good. I attempt to explain this with the description above the

chart.

It is worth noting, the GC content areas highlighted for the Superframe are only those that are above the threshold of the mean, not below it. This is because it is those areas that are above could be explained by an ORF Location, as the GC content tends to be higher in protein coding regions, and so there is no use in displaying the GC content windows out of threshold by being below, which cannot be explained by the ORF Locations.

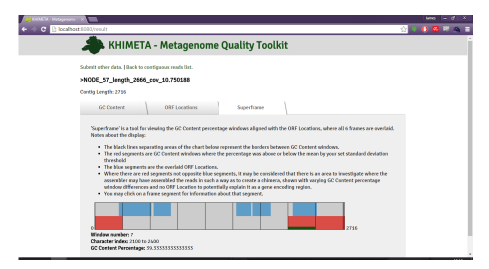


Figure 2.17: The result of when a user clicks to view a window data, they can see the particular percentage of the GC content of that window, and where that window starts and finishes, if they wished to inspect the contig themselves using those numbers.

2.2.4 Error

If a user encounters an error on the website, such as attempting to access pages without first submitting data) the page in the figure below will be shown.



Figure 2.18: If an error occurs as the user is browsing the web service, i.e. they try to access a page when they do not have the required data submitted, they are presented with this page informing them of an issue.

While not very descriptive, it serves to return the user back to data submission, which is the best place to start again if something bad went wrong or they somehow tried to access a page when they hadn't previously submitted the required data. If there were a future revision of this page, it could be more descriptive of the exact reason why the error occurred, but was out of scope for the current project to make a fully detailed error reasoning. There is an ErrorController class in place however that can deal with this functionality in the future if the application were to be continued. Currently any errors are run through the Error controller but it only returns this page to the user.

2.3 Support Tools

For writing the code, I used JetBrains IntelliJ Community Edition IDE. This included writing all the code be it Java, HTML, CSS, Javascript, ThymeLeaf and any additional properties for setting up Spring. This was very useful for text highlighting, debugging the code, and tied into my version control, using Git and hosted on GitHub.

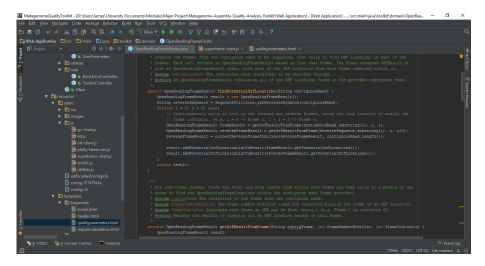


Figure 2.19: The Jet Brains IntelliJ Integrated Development Environment (IDE) I used for developing my application.

I also used http://www.tomato.es/ for counting my pomodoros. The web site provides a timer and a count for how many pomodoros have been completed during the time the browser has been opened.



Tomatoes is a pomodoro tracker, a Pomodoro technique $\ensuremath{\mathfrak{B}}$ driven time tracker.

Use the Pomodoro technique ${\bf @}$ to track your working time in 25-minutes slot called $\it pomodoro.$

Figure 2.20: A countdown of the pomodoro tomato timer on http://www.tomato.es/, used for breaking up development time and keeping track of work done over time.

2.3.1 Version Control and Continuous Integration

I used version control to keep a repository of my code in case of losing data in event of hardware crash or software corruption. For this I selected to use GitHub as I already had a repository on there and Git is supported in the IntelliJ IDE I was using. This can be found at https://github.com/coderghast.



Figure 2.21: A few of the commits made to my major project repository.

In addition, every time I checked in, I used continuous integration with CodeShip [4], allowing me to receive e-mails any time I made a commit to my repository that didn not pass the tests written. This was very helpful in discovering issues with failing builds when checking in a change and forgetting to update tests or breaking previously written code as the design changed.

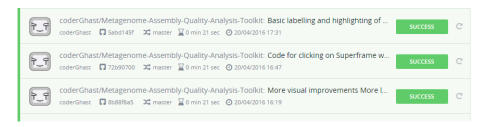


Figure 2.22: Examples of the continuous integration on CodeShip running my tests every time I committed to my repository.

Chapter 3

Implementation

This chapter serves to discuss the implementation specifics of the design of my application, including the development environment and any issues I encountered in implementing the features set out in my project objectives.

3.1 Development Environment

The application was developed primarily on a Fujitsu Lifebook A series with an Intel i5-3230M CPU @ 2.60GHz processor, 4GB of RAM and running Windows 10. The code was developed using Jet Brains IntelliJ IDE and Google Chrome developer console.

3.2 Features

3.2.1 Reading User Input

3.2.1.1 Implementation

The user input was one of the first parts of the application to be implemented, and consisted of reading in a FASTA file and outputting the content of the file to the console. Using TDD, this was tested by matching the output of the file read from a specific test file to the known file contents. The process was developed further to extract out the header that describes each contig, checking that it began with the '>' character and then every other line after this header was included as part of that contigs sequence data.

Originally it dealt with only one contig at a time, but was then expanded to be able to read in a full list of contigs and separate them based on where the header line starts. The code was eventually modified so that the user could paste data, that would be broken into components and processed in the same way. This was for the pasting of the user data rather than file upload, as I decided I would rather have pasted data than uploads for the current state of the application.

Figure 3.1: Part of the code for reading in a users data when they have pasted it into the text area of the web service. Deals with creating new ContiguousRead objects and adding them to a ContigResult every time it finds a new header for a contig (or reaches the end of the input).

3.2.1.2 Issues

It took some time to deal with the formatting of the file and knowing when one contig starts and one ends. The current system effectively finds every contig within the user content and extracts them out into objects, and so this is no longer an issue. It was solved through detecting when a new header contig header appears, put the contig sequence data and previous header into a new ContiguousRead object (if there is any data yet), and then start the reading process again under this new header. The final contig is constructed by reaching the end of the file and packing up whatever data is left over into a new ContiguousRead object.

While not an issue as such, the original version of the application would carry out the quality assessment of each contig as and when they were read in, to avoid holding too much data in memory. As the design evolved this was no longer a possibility, as it didn't allow the user to see their contigs before they were processed, or do any additional inspection on the contigs without holding the entire user data submitted in memory and going through it a second time upon an inspection request to find the contig they wanted. This option is still a possibility, but is an optimization aspect that wasn't in scope for the project. Additionally, the separation of responsibilities is better in this version of the code after being refactored from the time when the read class both read and assessed the contigs.

It did take longer to implement this than I had anticipated due to me changing from file reading to user pasting, and trying to find contig start and ends to read in the data versus read and assess at the same time.

3.2.2 Counting GC Content & Percentage

3.2.2.1 Implementation

Counting the GC content of the application involved breaking up the contig characters into sizes based on the set window length by the user, then working out the percentage of G and C characters in each of those windows. Each part of the contig that is split is put into a 'GcWindow' object, that has methods for counting the number of G and C characters within it.

```
private static ArrayList<GcWindow> splitIntoWindows(String contiguousRead, int windowSize){
    ArrayList<GcWindow> windowedAssembly = new ArrayList<>();
    for(int i=0; i < (contiguousRead.length()); i += windowSize){
        StringBuilder stringBuilder = new StringBuilder();
        if(i + windowSize < contiguousRead.length()) {
            stringBuilder.append(contiguousRead.substring(i, i + windowSize));
        } else {
            stringBuilder.append(contiguousRead.substring(i));
        }
        windowedAssembly.add(new GcWindow(stringBuilder.toString()));
}
return windowedAssembly;
}</pre>
```

Figure 3.2: The code for splitting a contig into windows available for calculating the GC content window percentages.

Once split, the GcWindows are passed to have their percentages calculated and added to an ArrayList of Doubles This is used for calculation with the mean, standard deviation and for returning results to the user for use in the View.

```
private static ArrayList<Double> calculateGcContentPercentages(ArrayList<GcWindow> windows) {
    ArrayList<Double> percentages = new ArrayList<>();

    for (GcWindow gcWindow: windows) {
        int gcContent = gcWindow.getCCount() + gcWindow.getGCount();
        int contentWindowSize = gcWindow.getWindowContentSize();
        if(gcContent > 0) {
            percentages.add(((new Double(gcContent) / new Double(contentWindowSize)) * 100));
        } else {
            percentages.add(new Double(0));
        }
        return percentages;
}
```

Figure 3.3: The code for calculating the GC content percentages for each window passed into the method.

3.2.2.2 Issues

In principle this is, and was, an easy task, yet took me longer to complete because my domain knowledge was still somewhat lacking and I got caught up on the little details that kept me from progressing, even though I didn't need to know of them or use them in the process of working out

the GC content windows. Overall there were no issues with the implementation once I got my head around why it was worth doing this and how it should be presented as a quality measure to the user.

3.2.3 Displaying GC Content percentage

3.2.3.1 Implementation

I began to attempt to implement GC content viewing with Plotly.js right from the beginning, and so I had a prototype up and running fairly quickly where I manually took the results from the console output of the Java application and pasted them into the final containing my Plotly.js prototype to be used as data.

Next I worked this in with Thymeleaf to get the results directly from my Java application, then turned them into JavaScript and used the data from that for the chart. The x axis labeling for the chart is manually created in order to represent where the GC window for each bar starts and finishes. The earlier design had window numbers along the x-axis, populated by Plotly.js. These were of no use to a user though, as if they wanted to find out what GC window they were looking at they would have to manually work out where the window started and ended from their window size multiplied by the window number of interest.

```
/*Script th:inline="javascript">
/* Setup data needed */
/*<![CDATA[*/
    var windowdata = [[${gcResult.gcPercentages}]];
    var windownums = [[${gcResult.gcWindows}]];
    var gccontentmean = [[${gcResult.gcAverage}]];
    var gcColours = [[${gcResult.gcBarColours}]];
    var gcMeanForAllWindows = [[${gcResult.gcMeanForAllWindows}]
    var windowSize = [[${gcResult.gcMeanForAllWindows}]];
    var contigLength= [[${contiguousread.gcWindowSize}]];
    var contigLength= [[${contiguousread.contigLength}]];
    var contigContent = [[${contiguousread.contigContext}]];
    var orfData = [[${orfResult}]];
    function init(){
        drawGcChart();
        setupOrfChart();
        setupSuperframeChart();
    }
    init();
/*]]>*/
```

Figure 3.4: Extracting the data from the Model, provided by the Controller to the View, using Thymeleaf's inline tag to be able to use CDATA to convert the Thymeleaf extracts into JavaScript.

The red bars of the GC chart displaying where the percentage of a window is over or under the threshold of the mean of all GC window percentages was implemented as a useful aid to the user, and the colours are set in the Model rather than made in the View. The reasoning for this was to implement a system that would provided the View with what it needed and not needing the View to calculate anything, only consume the values (in this case the GC window data and RGB data) to display results.

```
public class GcResultViewData {
    public ArrayList<Double> gcPercentages;
    public int[] gcWindows;
    public double gcAverage;
    public ArrayList<String> gcBarColours;
    public double[] gcMeanForAllWindows;
}
```

Figure 3.5: 'GcContentViewData', the object provided to the View in order to display the GC content data to the user.

The final thing to add was the Mean line. This is just displaying to the user where the mean of all the GC windows lie, and helps with the visualization of if any windows of their contig look like they might have issues.

3.2.3.2 Issues

The main issue I had with displaying the GC content percentages with Plotly.js was getting data to actually display as I was unfamiliar with Plotly.js and Thymeleaf once I started using the Spring Framework. Thymeleaf makes getting data from the Objects in the Model very easy and simple, but since I was trying to integrate two technologies together that I had never used before, it took some time to find the right way of accessing the data from my Object through Thymeleaf, into JavaScript and then the correct format for use by Plotly.js.

The next issue was trying to get the line for the Mean to show. The way I got this to work was using an additional data input for the same chart as displaying the GC content window bars and having the same number of data points as there are GC windows, where each data point is the mean number, in order to get the chart to display a line across the entire chart to match the GC windows.

3.2.4 Finding Open Reading Frames

3.2.4.1 Implementation

The implementation of Open Reading Frames happened in a few steps, with the design and code evolving as I understood more what they were and how best to present the results. At first, I was only finding a single, longest ORF Location, as I did not fully comprehend what an ORF was. This was just looking at a contig and finding the first start codon of ATG and the last stop codon of TAG/TAA/TGA. Once I understood them though, the process became about how to find whatever ORF Locations I could. Using Test Driven Development was extremely useful for this step, as I wrote tests for finding specific ORF Locations in small sequences I created myself and ran the code against them.

After a number of attempts at trying to find the ORF Locations within a contig frame, I settled on a method of going through the entire frame, breaking it up into characters of 3 and only keeping the start and stop codons in lists of each. Each codon was stored with data about where it starts and ends, in order to be able to later reconstruct the contig between a start and stop codon based

on the character positions.

```
private OpenReadingFrameResult getOrfResultFromFrame(String contigFrame, int frameNumberModifier, int frameIndicator)
    OpenReadingFrameResult result;

ArrayList<Codon> startCodons = new ArrayList<>();
ArrayList<Codon> stopCodons = new ArrayList<>();

for (int i = 0; i < contigFrame.length(); i += 3) {
    if ((i + 3) <= contigFrame.length()) {
        if (contigFrame.substring(i, i + 3).equalsIgnoreCase("ATG")) {
            startCodons.add(new Codon("ATG", i + frameNumberModifier));
        }
    }
}

for (int i = 0; i < contigFrame.length(); i += 3) {
    if ((i + 3) <= contigFrame.length()) {
        String nextCodonInFrame = contigFrame.substring(i, i + 3);
        if (stopCodonDetected(nextCodonInFrame)) {
            stopCodons.add(new Codon(nextCodonInFrame, i + frameNumberModifier));
        }
    }
}

result = constructOrfsFromCodons(startCodons, stopCodons, contigFrame, frameNumberModifier, frameIndicator);
return result;
}</pre>
```

Figure 3.6: Finding all of the start and stop codons from within the passed frame from the contig.

Each frame is separated by creating a substring of the original contig, removing 0, 1 and then 2 characters for the first three frames, and the same for the reverse frames but with the contig sequence reversed with the opposing base pair of each character used.

```
public OpenReadingFrameResult findPotentialOrfLocations(String contiguousRead) {
   OpenReadingFrameResult result = new OpenReadingFrameResult();
   String reverseSequence = SequenceUtilities.getReverseSequence(contiguousRead);
   for(int i = 0; i < 3; i++){
        // Simultaneously calls to look in the forward and reverse frames, using the loop iterator to modify the
        // frame indicator. (e.g. i = 0 -> Frame 1, i + 3 = 3 -> Frame 4)
        OpenReadingFrameResult frameResult = getOrfResultFromFrame(contiguousRead.substring(i), i, i);
        OpenReadingFrameResult reverseFrameResult = getOrfResultFromFrame(reverseSequence.substring(i), i, i+3);
        reverseFrameResult = correctReverseFrameIndicies(reverseFrameResult, contiguousRead.length());

        result.addPotentialOrfLocationListToResult(frameResult.getPotentialOrfLocations());
        result.addPotentialOrfLocationListToResult(reverseFrameResult.getPotentialOrfLocations());
    }
    return result;
}
```

Figure 3.7: Extracting each frame and calling to run the ORF Finding process.

For each ORF Location found in a reverse frame, the start and stop indexes are swapped, representing that what is the 'Start' of an ORF Location is actually closer to the end of the contig. This is so that it can be displayed appropriately in the View, and match up with the proper direction with the forward frames.

Figure 3.8: Getting the base pair characters of a reverse frame is as simple as a switch statement and building the reversed contig from back to front.

Through this process, it made it possible to find which codons came before and after which within the contig in order to find which would construct the longest ORF Locations, which start codons were redundant (as they were between an earlier start codon and the next stop codon), and in my first completed version of the algorithm, where the last stop codon before the next start codon was.

At this time, I still had a misunderstanding about protein coding regions, in that I believed the end of an ORF Location was at the last stop codon before the next start codon, when in fact it should have been the first stop codon it reaches is the end of the current codon. This was a simple fix though as it just involved removing a section of my code and fixing my tests.

Figure 3.9: 'Zipping' together ORF Locations from start and stop codons within the frame.

Once each frame has been processed to find the ORF Locations within them, the results of each frame are combined together in a single OpenReadingFrameResult. Each ORF Location is aware of what frame it belongs to, and so combining them all into one result doesn't make a difference when determining what ORF Location belongs where.

3.2.4.2 Issues

The two main issues with implementing the Open Reading Frame finding algorithms were my lack of understanding of what they were when I began, leading to the second issue of implementing more than was necessary. It took me longer than I had anticipated to implement the code for finding ORF Locations because I believed I had the additional requirement task of finding the last stop codon before the next start codon, where every stop codon between the initial start codon and that last stop codon was to be included in the ORF Location but ignored as an actual start and stop.

Thinking up how to do this algorithmically took some planning and time, and eventually I came to the conclusion of finding all start and stop codons and then running a while loop, checking that there was at least 1 of each left in each list to make an ORF Location, where the start came after the stop (for reverse frames, this logic still applies as the indexes are only reverse after the ORF Locations have been constructed) and then carrying out a number of conditional checks to remove any start and stop codons that came between the longest ORF that could possibly be made.

Once I had realized this was erroneous and the first stop codon encountered is the actual end of an ORF Location and not to continue, it was too late to recover any time and I just had to remove the code that caused the issue and rewrite my tests to match the actual expected outcome. Aside this, once I designed the code based around the 'zipping' of start and stop codons together, the implementation was relatively simple, using unit tests to ensure I was correctly implementing the algorithm for expected results.

3.2.5 Displaying ORF Locations

3.2.5.1 Implementation

The first part of the ORF Location displaying for the View takes place in the Controller, where it strips away ORF Locations under minimum length threshold set by the user and then sorts all ORF Locations by length using a comparison method. The display of the ORF Locations in their particular frames is done using HTML5 Canvas, one for each frame. As each ORF Location holds an integer value representing what frame it belongs to, when painting the canvases it is a simple matter finding the correct context for the particular frame from an array of canvas contexts

```
var currentContext = contextList[orfData[i].frameIndicator];
```

Reverse and forward frame ORF Locations are painted in the same way, except the start and stop indexes are swapped for reverse frames. Likewise, for the highlighted ORF Location to be painted, the only difference is that the current highlighted frame is stored in a global variable (updated any time an ORF Location is clicked in the list or on a frame) and when re-painting the frame canvases and iterating through the list of ORF Locations, when the loop reaches that ORF Location the fill colour is set to be different than every other ORF Location.

Finding a click within a canvas frame is as simple as checking if the click is horizontally within a frame location by calculating where the start and stop points of all the ORF Locations (with that frame in particular) are and if the click falls within one of the the ORF Locations.

Figure 3.10: Finding if a click if within an ORF Location is as simple as going through the list of ORF Locations, only checking against those within the same frame as the click, then looking at whether the click is within the start and stop points of an ORF Location within the canvas, based on the size and location of where it was painted (for reverse frames, the start and stop points are swapped, as reverse frames are displayed in the same direction as forward frames, with the indexes in the right order for a reverse frame).

When an ORF Location is clicked to be viewed in more detail, the characters within that sequence needed to be formatted to be able to be reasonably displayed. This process took some time to build, and through refactoring was re-written twice over until I found a result that was reasonably fast at formatting and worked the way I wanted it to. This is that it highlighted every start and stop codon within a sequence, split the characters into groups of 3, every 15 groups of

3, a new line is to be created and where the start of the line has the index within the contig where that line of the ORF Location begins.

Figure 3.11: Formatting the characters from within an ORF Location into HTML tags to display the data in the way I wanted it to be designed, then inserting it into the div for displaying ORF Location information within the page.

3.2.5.2 Issues

When formatting the ORF Location characters into the list, I found I kept having results where the newline break would be incorrect, or not all of the start codons were highlighted. This issue was because the way I built the formatted text was by adding text into an array, and then breaking up the text to add newlines whenever the array had a remainder of zero when divided by 15. This meant that it was including the indicator of what index the line of the ORF Location was on as part of the calculation. The result was that the line would be split incorrectly. Once I changed how the formatting was done and included the character index as part of the first set of characters of each line, the calculation worked correctly for splitting the lines where I wanted them to be split.

Although not an issue as such, my first version of drawing the canvas frames involved a single canvas. While the displayed result was the same, it meant that the detecting of user clicks within ORF Location algorithm took longer to process as it had to determine which frame the user was clicking on and then find if it was within an ORF Location. Additionally, having it all in one canvas made the formatting of the canvas results on the page a little more challenging as I couldn't place values (like the frame indicator) exactly next to each frame without painting them within the canvas.

By just splitting the single canvas into 6 different canvases, one for each frame, it allowed

me to easily surround the canvas elements with text for frame indicators and cut down on the code for detecting where and what the user was clicking on within each frame, as shown in the implementation of the click detection above in the section above.

3.2.6 Superframe Comparisson

3.2.6.1 Implementation

Implementing the Superframe was a relatively simple procedure as all of the data required, and part of the code, had already been done within the GC content and ORF charts. Since the Superframe just displays both of these together, it was simply overlaying the 6 frames together and painting on the GC content windows that were out of the threshold (above it, not below) and painting lines to represent the breaks between windows. The algorithm for detecting and highlighting user clicks within windows is also much the same as the code user for the ORF Location clicks and highlighting, just modified to detect within windows rather than within the frames.

The canvas is set up so that it could also be used with other future techniques from within windows, such as k-mer frequency analysis results, and be a tab page where any comparison or overall results could be displayed.

3.2.6.2 Issues

There were no issues implementing this content as the code had mostly been done previously. The main challenge was trying to think of a good way of displaying to the user where there might be potential issues, and giving them a report on the quality. What the application does is just displays them this overlay of the techniques used right now and leaves them to come up with their own decision.

It would be nice in future to be able to expand on this and perhaps be able to truly give the user a report that tells them where there are good or bad areas of their data, instead of making them infer it for themselves through this chart, but without additional techniques this is a challenging prospect.

3.2.7 Implementation Review

During implementation, I found that GC Content percentage and ORF Location finding took a lot longer than expected, especially considering that I implemented an additional part of ORF Location finding that wasn't needed, and refactored it 3 times until I reached the point where it correctly found them.

The user interface design and coding to format it also took a lot longer than I had originally anticipated. I wanted to present the information in a clean and useful way, without cluttering the page, and a number of redesigned happened due to the underlying code evolving over time to meet the requirements. Getting used to Thymeleaf and how it took the data from the Objects took a while to get used to, too, as I was unfamiliar with Spring and Thymeleaf and wasn't sure how I was adding the data to the Model and then not clear on how it accessed it, or how it was used in form submissions.

The Superframe also needs additional consideration. A lot of the quality assessment is still on the user to look at the produced reports and their content to consider whether they themselves feel their assembly file is good or not through factors highlighted by my application.

Overall, these issues and lack of understandings delayed me to the point where I didn't have time to implement k-mer frequency analysis or any of the 'nice to have' features, and there are certainly areas of the current implementation where I feel bits are lacking, such as file upload, verbose logging and validation and the aforementioned Superframe. What I have got working though, I feel is of decent quality, my process of iterative development and design evolution with prototyping worked quite well and the requirements for those sections are fulfilled.

Chapter 4

Testing

4.1 Overall Approach to Testing

My approach to testing was using Test Driven Development and refactoring, taking note from XP practices, ensuring that I only developed as much as was required to make tests pass based upon my requriements for eachi part of the applications intended functionality. The aim was to have as many tests automated as possible, as this would allow me to test frequently and reliably and tie into testing everytime I checked into my Git repository with CodeShip's continuous integration of my code. Where automation could not be carried out, I created test files and used a test table to determine whether my application met the requirements I could test against.

4.2 Automated Testing

4.2.1 Unit Tests

Writing unit tests before writing any functional code to match against was very useful, allowing me to develop code againsts the tests, written one by one, so that I would not develop any more than necessary. This made is so that what was developed fit the requirements, as the tests written before the code were based upon the exact specifications of the requirements. Refactoring the code after each test kept my design evolving and improving maintainability and simplicity to remove duplicate and unnecessary code.

Figure 4.1: An example of a unit test used in my application.

Each unit test was made to only carry out one function check at a time, and the naming of each test reflected what the functionality of the system part under test should do. It is for this reason that the item under test is called 'sut', system under test, and each test is named with a convention

that has a 'should' clause in the name, to demonstrated that the function called should produce particular expected results.

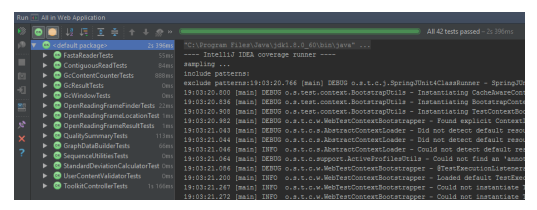


Figure 4.2: Results of running the set of unit tests developed for the application.

IntelliJ gave me the tools to run my unit tests with coverage, allowing me to see how much of my code was actually tested against. In the figure below it can be seen that most of my methods were covered. Those methods that were not covered I chose not to test against, as they are mostly setters and getters with no further functionality. In practice, it would perhaps be best to test against these too, in case for some reason the way a value is set or retrieved had to take into account some processing.

Coverage All in Web Application							
+	80% classes, 84% lines covered in package 'toolkit'						
⊢Ba	Element	Class, %	Method, %	Line, %			
L ₀	domain	85% (12/14)	83% (60/72)	87% (275/313)			
₹.	utilities utilities	100% (4/4)	100% (5/5)	92% (47/51)			
Ţ	web web	50% (1/2)	16% (1/6)	19% (4/21)			
	☑ Main	0% (0/1)	0% (0/1)	0% (0/3)			
₽.							
×							
?							

Figure 4.3: The test coverage of my unit tests over the developed application code.

Coverage All in Web Application							
↑ 85% classes, 87% lines covered in package 'domain'							
⊢ B a	Element	Class, %	Method, %	Line, %			
10	C Codon	100% (1/1)	66% (2/3)	83% (5/6)			
₹.	© ContigResult	100% (1/1)	100% (5/5)	100% (10/10)			
Ţ	ContiguousRe	100% (1/1)	42% (6/14)	68% (24/35)			
	G FastaReader	100% (1/1)	100% (4/4)	90% (55/61)			
1	G GcContentCo	100% (1/1)	100% (3/3)	90% (20/22)			
×	G GcResult	100% (1/1)	100% (5/5)	94% (16/17)			
3	G GcResultView	0% (0/1)	0% (0/0)	100% (1/1)			
?	G GcWindow	100% (1/1)	100% (5/5)	100% (13/13)			
	© OpenReading	100% (1/1)	100% (6/6)	100% (65/65)			
	© OpenReading	100% (1/1)	100% (8/8)	83% (26/31)			
	© OpenReading	100% (1/1)	100% (5/5)	100% (19/19)			
	© QualitySumm	100% (1/1)	100% (7/7)	100% (11/11)			
	© QualityToolkit	0% (0/1)	0% (0/1)	0% (0/9)			
	© UserParameters	100% (1/1)	66% (4/6)	76% (10/13)			

Figure 4.4: The test coverage of my unit tests over the developed application code, broken down for each class.

4.2.2 User Interface Testing

For testing the user interface, I used Chrome's Developer Tools for checking the correct running of the JavaScript, and debugging the code as it ran on loading of the page. I also confirmed that each of my pages loaded and functioned correctly in Google Chrome, Firefox and Microsoft Edge.

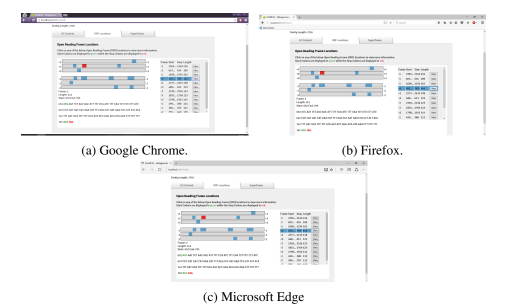


Figure 4.5: The same page and results from the application shown in different modern browsers.

For testing the speed performance of the web pages of the applications loading time, I used YSlow [13] which runs a number of checks on the page setup and loading speed to return a report about whether a page loads quickly or slowly, and how it could be improved.

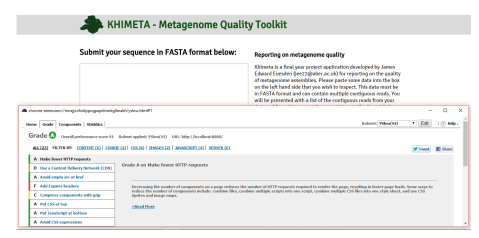


Figure 4.6: YSlow report after running on the Welcome page.

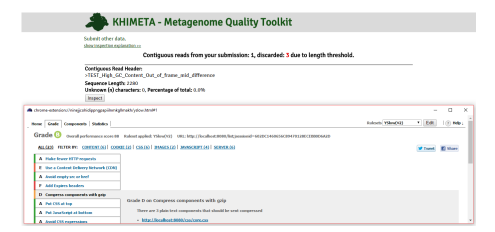


Figure 4.7: YSlow report after running on the List page.

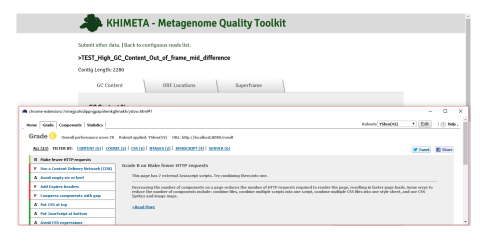


Figure 4.8: YSlow report after running on the Results page.

My results showed me that for the most part there was not much I could do to improve my application outside adding expiry headers to some session variables and implementing a Content Delivery Network (CDN), which was far outside the scope of the project.

4.2.3 Manual Testing

A number of artificial files were created in order to be used for testing. Some of these were used in automated tests, such as checking for a files existence and possibility of being read, or throwing exceptions when a test file did not exist while other files were for running manually, either being entirely artificial and checking for individual expected behavours, or composed of actual data from multiple species that I manually split and combined together to view the results of. The completely artificial file data is included in the appendices. While the results of the mixed real species data is not included, the results of one very obvious combining of species can be seen in the figure below.

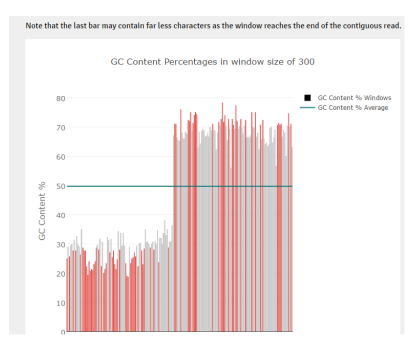


Figure 4.9: Combining two species contigs together at 50% of each of the file (first half one species, second half another species), we see a very obvious split in the GC chart.

Running this test showed me that it is possible to see where there is a huge change in GC content, but that the threshold won't pick it up because with a case like this, while extremely unlikely to happen where there is a mix of only two species, the threshold only shows those outside of the mean, not drastic changes. It highlights that there is room for improvement with detecting these changes and reporting on them to the user, rather than having them infer it themselves.

4.2.3.1 Real Data

I also tested the application using real metagenomic data. This data was provided by Sam Nicholls, and was taken from the gut of a limpet. There is a large file of thousands of contigs, with the shortest being under a hundred bases long and the longest being above twenty five thousand bases.

Running this real data and looking at both the short and long contigs allowed me to see that my application did produce results for displaying areas where there could be chimeric regions in contigs, displaying ORFs and highlighting areas where an ORF may align with a highlighted GC content window.

4.2.3.2 Test Table

For confirming functional requirements were met and testing things that couldn't be done with automation, I produced a test table with tests matching requirements, expected results and avoiding unwanted results. The tests from the table were carried out by fulfilling the action in the table, and then viewing the results and marking whether the expectations were met or not. You can see the System Test Table included in the Appendices.

When it came to running large files, or files with many contigs, I ran a number of my laptop and for the most part it handled them quite well with a few hundred contigs of moderate length, although with contigs in the sizes of a hundred thousand characters and up, the process of reading in the contigs and then processing one starts taking noticeably longer. I believe this is a restriction with my own machine, however, as should the JVM be allotted enough memory while the application is hosted on a larger, faster machine it is likely it could perform far better.

Chapter 5

Evaluation

This chapter attempts to evaluate my project as a whole, including how well I felt I carried out my analysis of the problem, designed and implemented the application, whether I feel it would be useful to anybody, whether the resulting application met the project objectives and what I could have done differently, or could be improved upon.

5.1 Requirements Analysis

When looking back on the original problem stated, the requirements I set for the application and the resulting application now that the project has finished, I both believe that I made some good choices and bad choices in what I selected to do, and whether it was worthwhile to produce for an actual user.

On the one hand, by selecting to create a web service that allows users to put in FASTA files and inspect the contigs within that file and carry out a number of quality inspections on the data is a great idea. Having a single place to do this with a page that allows the user to view a number of statistics about their data could be highly useful. However, when I think about my choice of working with GC content and Open Reading Frames, and not understanding the domain enough to implement them faster than I did, I feel like perhaps it would have been a better idea to start with k-mer frequency analysis, as this may have been a better measure of quality than what the current application does.

While this may be the case, the way I did develop the application probably helped me understand the domain a little better than if I did begin with k-mer frequency analysis, even if I just consumed output from 3rd party software, and so while the application may lack in some usefulness, what I gained from following the route that I did may have been better in the long run, if I were to continue developing the application and add the functionality of k-mer frequency analysis.

It is for this reason I also feel justified in designing my application as a web service, where adding additional functionality to serve to a user from somewhere it is hosted on allows multiple features to be added without the demand of processing power or memory on a users machine if the application were a regular software application for them to run locally. It does come with the downside that in order to be able to process very large user requests it needs to be hosted on a powerful machine, but as the cost of computing power and memory lowers year by year, it is possible to do this if the demand were there.

5.2 Technical Achievement

The resulting application is a piece of software that carries out a number of tasks, and fulfills each of the functional requirements I set out to do, except for those previously mentioned, has test coverage for the functionality and could have a potentially useful role with those who deal with sequence assembly files. The technical challenge of implementing such an application began with understanding the domain, selecting suitable technologies and then carrying out the development within my development lifecycle plan with XP and Scrum.

The application does give a user access to information about their assembly file that they wouldn't have unless they used multiple different tools elsewhere and combined those results themselves. It is built upon Spring Framework, using the MVC framework design pattern and presented as a web service with a lot of room to grow over time with new techniques for quality assessment. I believe that the technical achievement I made from developing this project is pretty good. I got to use technologies I was unfamiliar with (Spring Framework, Thymeleaf, Plotly.js), develop using an agile methodology to appropriately evolve the application design(XP, Scrum), use known software development techniques and practices (MVC, RESTful web service, etc) and came out with an application that produces results that can be tested against requirements.

I am pleased with my choice of technologies, and know that while it would have been possibly to develop the application in a different language (Ruby, C++, etc) or even develop it as a standalone application and not as a web service, I stand by my decisions that they were what I felt were the most suitable choices for this application for my implementation, understanding and knowledges. There are a number of technical improvements that could be done, and these will be discussed in the next section.

5.2.1 Future Work

5.2.1.1 Improvements

Within the current application, there are a number of improvements I would like to carry out. Understandably, there is always room for more refactoring and improvement of the code. As my experience as a software developer grows, so will my skill in refactoring and code design. I cannot claim anything as the best it can possibly be, and so I will always say there is room for improvement in the code and application design.

On the subject of design, the front end design of the user interface also has much room for improvement. This can come in the form of better HTML and CSS layouts, colour choices, better explanations of the applications use, and most importantly, better design of the results. I am pleased with how some of the results are displayed, in particular the display of the list of contigs and the Open Reading Frame results tab are both areas I am pleased with. The GC content chart I feel could be better in demonstrating areas of drastic differences rather than just differences in individual windows over and under a threshold with the mean. This could be done with sliding windows with multiple passes using different window sizes (of reasonable length) and then display better results to a user.

Likewise, the 'Superframe' could also be improved. While its function is useful to display to a user where there are matches between protein coding regions and areas of high GC content, outside of the threshold from the mean, I know that if I had more time to work on the application

I could find more and better ways of demonstrating this to a user, and better ways on reporting on which GC windows could be 'natural' occurrences of high GC content and which cannot be explained (through a lack of an ORF Location or being lower than the threshold of the mean).

There is also room for improvement in the validation of parameters, and giving the user a reason about why their input of parameters or data failed. Right now the application will return a generic error page any time an exception is thrown in the application, or if they input bad data for parameters the application will ignore it and put them back on the page to try again, but won't indicate why this is the case. I attempted many times to get this functionality to work with Thymeleaf and Spring but couldn't get it to function. It is another feature that I feel if I had more time to put into the project I could find the solution.

5.2.1.2 Additional Functionality

Along with the improvements to the existing code and functionality, there are a number of features I would like to implement too. First and foremost, once the improvements had been carried out, I would like to allow a user to upload their own file rather than just pasting in data. This would include decent validation of the users file and implementing restrictions on file size too (this could be altered, based on where the application was hosted and how much memory would be allowed to be alloted to the application).

I would definitely want to implement k-mer frequency analysis, too. Beginning with a prototype, even if it was not very efficient, I would attempt to develop my own algorithm for the application, and also consume the results from the applications that already do k-mer frequency analysis and compare my own results against it, and then decide from there the best way to implement it. I do feel that self-implementing it would be better than 3rd party software, however, as I still stand by that having to rely on 3rd party systems can sometimes lead to issues in the future with maintainability should they be removed, modified or no longer supported, unless there is no other choice but to use them.

The potential to tie the application in with sending requests to NCBI's BLAST for checking if contigs, or parts of them, have reference sequences would also be a great feature to provide to the user as a way of determining if the have a useful and quality contig too. This ties in with thinking of other and better ways of reporting on the quality of the users assembly data. If given more time, and with the domain knowledge now, I could think of other ways that could demonstrate quality to a user, on their own file, on individual contigs, on portions of data, through visual techniques, confidence factors and more. There is so much room to expand the application that it could become a fully fledged software application taken forward over a year or more, if someone wished to do so, to present the ultimate toolkit for reporting on metagenome assembly quality.

If I were to begin the project again from scratch, with my current knowledge of the application and the domain, I feel like I could implement what I have done already a lot faster, finish the improvements, work through k-mer frequency analysis and try to find additional ways of reporting on the quality of the data, perhaps seeking out groups of potential users and really drilling into the project problem and what would be the most useful to them outside of my own ideas.

5.3 Project Management

Now that the project has been completed, I look back on my decision to use and adapt Scrum and Extreme Programming agile approaches for the project lifecycle and think that they were very useful in helping me design, plan, test and work through the tasks that need to be done, but that perhaps it would have been a better choice to use Feature Driven Development (FDD). I cannot say that what would have been produced if I used FDD would have been produced any faster or better than using the Scrum/XP hybrid, but that it could be a possibility to use if I started the project again.

For the most part, I found that using Scrum and XP helped, especially the practices of XP. There are a number of XP practices that I could not implement, however, such as pair programming, which is key to the success of XP in that it ensures that the Test Driven Development and refactoring carried out on the application are reasonable and logical, and not a single developer in a silo believing themselves to be doing the right thing but instead making errors or being inefficient.

With Scrum, while breaking down functionality into User Stories was helpful, sometimes it became trivial to break things down into much smaller slices and create tasks for simple processes, such as 'Add button for inspection'. On the other hand, iterative development through Sprints, being able to reflect on the application with Sprint Retrospectives and daily stand-ups with my peers was extremely useful for motivation and discussions of development to help my thought processes.

Using the Pomodoro technique was also useful, as it helped me keep track of how I was doing for work each day, stay focused, find time to work when I might normally put off doing something and made sure I didn't overwork too. It'd definitely a technique that I will put to good use in the future of both software engineering and other creative projects.

5.4 Final Conclusion

Overall, I look back on the project and see an application that met some of the original requirements, where the requirements were slightly hazy and unknown to begin with but produces a potentially useful tool to those who want to look for quality issues in their assembly data. There is large room for improvement and advanced functionality with the tool to be even more useful to users, and if given more time this could be done. The project process was mostly successful, although motivation during the beginning of the project was low due to not fully understanding the domain or requirements of the finished application.

Through developing the application to find a solution to the project topic, I learned a number of things about the domain of metagenomics and technologies that exist to find quality in sequence data, new technologies and frameworks and about myself, what motivates me and what would be good practices to use for future self-driven projects. There are multiple things I would do differently if I were to start again, such as those mentioned in the previous section, but for the most part I feel I have accomplished a body of work that set out to do a task and completed enough of it that a user could feasibly get some use out of the tool, with the code design being good enough to be maintained into the future.

Appendices

Appendix A

Third-Party Code and Libraries

jQuery - The jQuery library has been used to display tabs for the results page and help detecting user clicks on HTML5 Canvas elements. The library is open source and it is available from the jQuery Foundation. [25]. The library is released using the MIT license. [5]. This library was used without modification.

Plotly.js - A library for displaying graphs and charts, used in the application for creating and displaying the GC content results. The library is open source and is available from Plotly. [9]. The library is released under the MIT license. [5]. This library was used without modification.

Spring Framework - The project is built using Java and Spring. The Spring Framework supports the application for being hosted as a web service. The library is open source and it is available from Pivotal Software. [19]. The library is released under the Apache License 2.0. [1]. This library was used without modification.

Thmyeleaf - The project used Thymeleaf for accessing data from the Java code with the Spring Framework Model and display it in the View through JavaScript. The library is open source and it is available from The Thymeleaf Team. [7]. The library is released under the Apache License 2.0. [1]. This library was used without modification.

1.1 Tabs for results

The code for displaying the results in tabs using CSS3 and Jquery [2]

1.1.1 HTML

```
<div id="tab2">...</div>
    <div id="tab3">...</div>
    <div id="tab4">...</div>
</div>
1.1.2 CSS
#tabs {
 overflow: hidden;
 width: 100%;
 margin: 0;
 padding: 0;
 list-style: none;
#tabs li {
 float: left;
 margin: 0 .5em 0 0;
}
#tabs a {
 position: relative;
 background: #ddd;
 background-image: linear-gradient(to bottom, #fff, #ddd);
 padding: .7em 3.5em;
  float: left;
  text-decoration: none;
  color: #444;
 text-shadow: 0 1px 0 rgba(255,255,255,.8);
 border-radius: 5px 0 0 0;
 box-shadow: 0 2px 2px rgba(0,0,0,.4);
#tabs a:hover,
#tabs a:hover::after,
#tabs a:focus,
#tabs a:focus::after {
 background: #fff;
#tabs a:focus {
 outline: 0;
#tabs a::after {
 content:'';
 position:absolute;
```

```
z-index: 1;
  top: 0;
  right: -.5em;
 bottom: 0;
 width: 1em;
 background: #ddd;
 background-image: linear-gradient(to bottom, #fff, #ddd);
 box-shadow: 2px 2px 2px rgba(0,0,0,.4);
  transform: skew(10deg);
 border-radius: 0 5px 0 0;
}
#tabs #current a,
#tabs #current a::after {
 background: #fff;
  z-index: 3;
}
#content {
 background: #fff;
 padding: 2em;
 height: 220px;
 position: relative;
  z-index: 2;
 border-radius: 0 5px 5px 5px;
 box-shadow: 0 -2px 3px -2px rgba(0, 0, 0, .5);
}
1.1.3 Jquery
<script src="http://code.jquery.com/jquery-1.7.2.min.js"></script>
<script>
$ (document).ready(function() {
```

```
script scri
```

```
}
});
});
</script>
```

Appendix B Ethics Submission

Appendix B

Ethics Submission

Ethics Application Number: 4295 Ethics Submission document overleaf.

AU Status

Undergraduate or PG Taught

Your aber.ac.uk email address

jee22@aber.ac.uk

Full Name

James Edward Euesden

Please enter the name of the person responsible for reviewing your assessment.

Reyer Zwiggelaar

Please enter the aber.ac.uk email address of the person responsible for reviewing your application rrz@aber.ac.uk

Supervisor or Institute Director of Research Department

CS

Module code (Only enter if you have been asked to do so)

CS39440

Proposed Study Title

MMP A Toolkit For Reporting On Metagenome Assembly Quality

Proposed Start Date

25th Jan 2016

Proposed Completion Date

4th May 2016

Are you conducting a quantitative or qualitative research project?

Quantitative

Does your research require external ethical approval under the Health Research Authority?

No

Does your research involve animals?

No

Are you completing this form for your own research?

Yes

Does your research involve human participants?

No

Institute

IMPACS

Please provide a brief summary of your project (150 word max)

Creating a web service tool that a user may submit some metagenomic sequences in FASTA format (ATGC characters) that returns them an estimated report of how good the 'quality' is from the created assembly of the sequence based on techniques such as GC Content counting and finding ORF Locations. The web service stores no data locally and processes the submitted data per user session. The data files accepted are just string characters that could be any sequence. The report produced is a visual report using charts to look for areas where there may be chimeric data

Where appropriate, do you have consent for the publication, reproduction or use of any unpublished material?

Not applicable

Will appropriate measures be put in place for the secure and confidential storage of data?

Does the research pose more than minimal and predictable risk to the researcher?Not applicable

Will you be travelling, as a foreign national, in to any areas that the UK Foreign and Commonwealth Office advise against travel to?

No

Please include any further relevant information for this section here:

If you are to be working alone with vulnerable people or children, you may need a DBS (CRB) check. Tick to confirm that you will ensure you comply with this requirement should you identify that you require one.

Yes

Declaration: Please tick to confirm that you have completed this form to the best of your knowledge and that you will inform your department should the proposal significantly change.

Yes

Please include any further relevant information for this section here:

Appendix C

Examples and Extras

3.1 Example User Story Breakdown

NOTE: This break down is from my blog entry at http://users.aber.ac.uk/jee22/wordpress/?p=147.

As a researcher

I want to get a report on the quality of my metagenome So that I know whether it is of good or bad quality

Okay, super high level. This can be broken down into:

As a researcher

I want to get a report on the GC content of my metagenome So that I can see where there might be inconsistencies

So, that could be explained better (i.e. what are inconsistencies? Areas where there might be a split/chimera, or just gene encoding regions and completely natural).

As a researcher

I want descriptions of the GC content of ${\tt my}$ metagenome So that I can pinpoint areas of interest

Perhaps a better way of making a story for GC content in this instance. What about the report?

As a researcher

I want a textual and graphical description of my metagenome quality So that I can see and understand where there might be quality issues

Again, quite high level, but not too bad. This could be broken down further.

As a researcher

I want a graph plotted to show me the GC content in my metagenome So I can visualise the distribution of GC content to better understand my metagenome

From some of these, further tasks can be broken down, so, lets take one and do that with the last story I defined. I suppose, before we can do that though, since we dont have an application developed, we might need some initial setup stories.

```
As a researcher I want an application to read in my metagenome assembly So I can see it outside of the FASTA file
```

Maybe thats pushing it a little. Theres not really much to be gained from this in business value, but, as far as development goes it can give us some nice little tasks:

```
Read in FASTA file
Output display of metagenome visually for researcher to understand
```

Thats just two simple tasks. Read in a file type, and with the contents, display it. It might not be much, but its a start where we can say to a hypothetical researcher Okay, weve taken your file, and we can show you that your metagenome looks like this. Theres no processing done to it, but you can see how with this visualisation, theres the room for labelling and noting the interesting points later. What do you think?

This story with others could then help us propose a Sprint Goal, as below:

Sprint Goal: To display a metagenome in an application after reading in a FASTA file, with the look at implementing GC Content counting should time allow.

3.2 Artificial Test File

>TEST_High_GC_Content_Out_of_frame_mid_difference CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG >TEST_Low_GC_Content_In_Frame_Throughout CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG >TEST_Sinlge_Frame_Many_Start_Stop_Codons CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG ATGGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCATGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGATGCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCTAGGCCCGGGCCCGGG CCCGGGCCCGGGCCCGGGCCCGGGTAAGGGCCCGGGCCCGGGCCCGGGTGAGGG CCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGGCCCGGG >TEST_50_percent_N_characters

 Appendix D System Test Table

Appendix D

System Test Table

System Test Table is included overleaf

System Test Table

Khimeta System

ID	Requirement	Description	Inputs	Expected outputs	Pass/ Fail	Comments
M1.1	FR1	On Welcome page: User can request the minimum size of a contig	Valid contig over 1000 Minimum length of 1000	The contig appears on the List page	P	
M1.2	FR1	On Welcome page: User can request the minimum size of contig	Valid contig under 1000 Minimum length of 1000	The contig does not appear on the List page	P	
V1.1	FR1	On Welcome page: User cannot input incorrect parameters	Invalid contig Valid minimum length	User is returned to Welcome page to reinput parameters	P	Could do with improvement by giving reason why the parameter(s) were bad
V1.2	FR1	On Welcome page: User cannot input incorrect parameters	Valid contig Invalid minimum length	User is returned to Welcome page to reinput parameters	P	Could do with improvement by giving reason why the parameter(s) were bad
M2.1	FR2	On List page: User should see number of N characters of a contig	Contig with no N characters	N character count is 0, percentage of N is 0.0%	P	
M2.2	FR2	On List page: User should see number of N characters of a contig	Contig with 240 N characters of contig length 480	N character count is 240, percentage of N is 50.0%	P	
M3.1	FR3	A contig should have its GC content calculated and GC content results should be available in the Model	An artifical contig with known GC values in selected areas	GC content windows high in GC where expected from the test contig	P	
M4.1	FR4	The application should find the ORF locations of a contig in each frame	An artifical contig with known ORF Locations	The ORF Location results from the contig are found	P	

ID	Requirement	Description	Inputs	Expected outputs	Pass/ Fail	Comments
M4.2	FR4	The application should not find the ORF locations of a contig in each frame where there are non	An artifical contig with no ORF Locations	The ORF Location results from the contig should have no ORF locations	P	
M5.1	FR5	The application should carry out k-mer frequency analysis	A contig with known k-mer frequencies	The expected k-mer frequency analysis results	N/a	Not implemented
M6.1	FR6	When a contig aligns with known reference sequence using BLAST it should be found and compared	A contig that will align to some sequence data in an NCBI database	A match and comparison result	N/a	Not implemented
V2.1	FR7	The user can paste assembly sequence data into the application to be read	A number of contigs to be pasted in	The contigs from the pasted data should be found and read	P	
V2.2	FR7	The user can upload a file containing assembly sequence data into the application to be read	A file containing a number of contigs	The contigs from the file data should be found and read	N/a	File reading is implemented but no way for a user to manually upload. Needs front end implementatio n
V3.1	FR8	On the list page: A user can see their list of contiguous reads	There has been some sequence data put into the users @SessionAttributes	A list of contiguous reads	P	
V3.2	FR8	On the list page: A user cannot view any contigs in the contig list without submitting contig data on the welcome page first	No input, fresh session	User is taken to Welcome page	P	Could do with improvement by giving reason that the data was missing

ID	Requirement	Description	Inputs	Expected outputs	Pass/ Fail	Comments
V4.1	FR9	When inspecting a contig, a user should be able to modify the parameters of the inspection	With known test sequence data already in place: Valid Threshold highlighting Valid GC window size Valid minimum ORF location length	The results page, including the expected GC content chart, expected ORF Location chart and Superframe chart results	P	
V4.2	FR9	When inspecting a contig, a user should not be allowed to input invalid parameters for inspection	With known test sequence data already in place: Invalid Threshold highlighting Valid GC window size Valid minimum ORF location length	The user is kept at the List page to re-input the inspection parameters	P	Could do with improvement by giving reason that the Threshold highlighting parameter was invalid
V4.3	FR9	When inspecting a contig, a user should not be allowed to input invalid parameters for inspection	With known test sequence data already in place: Valid Threshold highlighting Invalid GC window size Valid minimum ORF location length	The user is kept at the List page to re-input the inspection parameters	P	Could do with improvement by giving reason that the GC window parameter was invalid
V4.4	FR9	When inspecting a contig, a user should not be allowed to input invalid parameters for inspection	With known test sequence data already in place: Valid Threshold highlighting Valid GC window size	The user is kept at the List page to re-input the inspection parameters	P	Could do with improvement by giving reason that the minimum ORF location length parameter was invalid

ID	Requirement	Description	Inputs	Expected outputs	Pass/ Fail	Comments
			Invalid minimum ORF location length			
V4.5	N/A	Results page: User cannot view without submitting contig data first	No input, fresh session	User is shown the error page	P	Could do with improvement by giving reason that the data was missing

Appendix E

Definitions of Technical Terms

DNA sequence: A sequence of letters that indicate the order of nucleotides within DNA (ATGC). It has the potential to represent information which has direct impact on the functions of living things.

Read: A short subsequence of DNA after a DNA sample has been run through a Sequencer. Often between 50-100 characters of DNA base pairs long.

Sequencer Breaks down and reads a DNA sample to produce many small reads in text format. Attempts to determine the order of the four bases, ATGC. The text character 'N' may also be included, where a sequencer encounters an error, due to sequencing technology limitations and/or difficulties reading DNA structures.

Bases G (guanine), C (cytosine), A (adenine) and T (thymine). Combinations of these bases create the DNA sequences.

Genome: A sequence that fully represents the genes or genetic material within a cell or organism. **Assembler:** An assembler is used to try and align and merge reads to try and create contiguous reads that represent the original DNA sequence that the reads have been created from by a DNA sequencer.

Contig: A contiguous sequence that represents a region of DNA, made up of overlapping reads from a DNA sequencer. Produced by assembly software.

Codon A series of three DNA nucleotides that represent an amino acid or stop signal, e.g. GAC, CTA, or stop signals such as TGA, TAG, TAA.

Annotated Bibliography

[1] Apache Software Foundation, "Apache License, Version 2.0," [Online] Available: http://www.apache.org/licenses/LICENSE-2.0, 2004, [Accessed on: 1 May 2016].

The Apache License 2.0.

[2] Catalin Rosu, "CSS3 & jQuery folder tabs," [Online] Available: http://red-team-design.com/css3-jquery-folder-tabs/, 2012, [Accessed on: 29 April 2016].

The tutorial and code for creating tabs using CSS3 and Jquery that I used in order to implement tabs for the inspection results page.

[3] F. Cirillo, The pomodoro technique. Lulu, 2009, 1445219948.

The pomodoro technique, used during the development of my application for efficient time management.

[4] Codeship Team, "Continuous Delivery with Codeship: Fast, secure and fully customizable." [Online] Available: https://codeship.com/, 2016, [Accessed on: 27 April 2016].

CodeShip, the web service I used to help provide me with contiurous integration for my project, connected to my GitHub repository to be run every time I checked in to the repository. This was integral to developing my application as it highlighted any time that I broke my build through forgetting to modify tests when content was updated, or when I broke parts of the application and so old tests would fail.

[5] —, "The MIT License (MIT) — Open Source Initiative," [Online] Available: https://opensource.org/licenses/MIT, 2016, [Accessed on: 1 May 2016].

The MIT license, for jQuery.

[6] N. C. B. I. R. Coordinators, "Database resources of the National Center for Biotechnology Information," *Nucleic Acids Research*, vol. 41, no. D1, pp. D8–D20, Jan. 2013.

The NCBI database and resources was extremely useful to me while developing my application. It gave me access to papers and research articles, data sets for downloading sequences to play with and explore and tools for techniques I wanted to implement in my own application. It's safe to say that without the existence of the NCBI resources, I would have understood a lot less about my project while beginning my development.

[7] Daniel Fernndez, "thymeleaf/thymeleaf," [Online] Available: https://github.com/thymeleaf/thymeleaf, 2016, [Accessed on: 27 April 2016].

The GitHub repository of Thymeleaf, for use with Java Spring Boot. Invaluable for me when accessing objects from the Model via the View of my web service.

[8] D. Huson and S. Mitra, "Introduction to the Analysis of Environmental Sequences: Metagenomics with MEGAN," in *Evolutionary Genomics*, ser. Methods in Molecular Biology, M. Anisimova, Ed. Humana Press, 2012, vol. 856, pp. 415–429.

A useful chapter in reading about a tool that has its own attempt at analysing metagenomic data. While the tool does not look at the quality, it looks at what the user provides and indicates to them if anything they have is already existing in reference data, and how their data compares to similar sequences already known. This could be considered as a quality measure, as if a sequence can be found to match a reference sequence, we know it is most likely not chimeric.

- [9] P. T. Inc. (2015) Collaborative data science. Montral, QC. [Online]. Available: https://plot.ly Plotly.js, the JavaScript library I used for creating the GC content window chart.
- [10] V. Kunin, A. Copeland, A. Lapidus, K. Mavromatis, and P. Hugenholtz, "A Bioinformatician's Guide to Metagenomics," *Microbiology and Molecular Biology Reviews*, vol. 72, no. 4, pp. 557–578, Dec. 2008.

This paper was invaluable to me at the beginning of the project, understanding what metagenomics is, and how the quality of an assembly can be full of errors, and why. While much of the paper was not necessarily relevant to my project as it deals with the full process of working with a metagenomic sample, for my background reading it was very useful.

[11] B. Liu, Y. Shi, J. Yuan, X. Hu, H. Zhang, N. Li, Z. Li, Y. Chen, D. Mu, and W. Fan, "Estimation of genomic characteristics by analyzing k-mer frequency in de novo genome projects," Aug. 2013. [Online]. Available: http://arxiv.org/abs/1308.2012

A paper highlighting the usefulness of k-mer frequency analysis and distribution in determining characteristics of genomes. Similar to my projects objectives where I wished to have my project use k-mer frequency analysis for quality assement.

[12] N. J. Loman, C. Constantinidou, M. Christner, H. Rohde, J. Z. M. Chan, J. Quick, J. C. Weir, C. Quince, G. P. Smith, J. R. Betley, M. Aepfelbacher, and M. J. Pallen, "A Culture-Independent Sequence-Based Metagenomics Approach to the Investigation of an Outbreak of Shiga-Toxigenic Escherichia coli O104:H4," *JAMA*, vol. 309, no. 14, pp. 1502+, Apr. 2013.

This paper was useful in understanding what the study of metagenomics can be used for, and so helped me better understand what metagenomics is and how it could be useful to have a tool for quality control.

[13] M. Duran, "YSlow - Official Open Source Project Website," [Online] Available: http://yslow.org/, 2016, [Accessed on: 30 April 2016].

YSlow, a tool for performance testing the loading of html pages, which I used for testing each of the pages of the application.

[14] G. Marçais and C. Kingsford, "A fast, lock-free approach for efficient parallel counting of occurrences of k-mers," *Bioinformatics*, vol. 27, no. 6, pp. 764–770, Mar. 2011.

Jellyfish, a tool for k-mer frequency anaylsis. A tool I looked into briefly at the beginning of my project when considering what techniques to use for my application.

[15] P. Melsted and J. K. Pritchard, "Efficient counting of k-mers in DNA sequences using a bloom filter," *BMC Bioinformatics*, vol. 12, no. 1, pp. 333+, 2011.

BFCounter, for k-mer frequency analysis. A tool I considered when thinking about the requirements and techniques involved in the development of my application.

[16] Mike Cohn, "Extreme Programming: A Gentle Introduction." [Online] Available: http://www.extremeprogramming.org/, 2016, [Accessed on: 26 April 2016].

The description of Extreme Programming (XP) that helped me understand the XP methodology and adapt it for my own use in this one person project.

[17] —, "User Stories and User Story Examples by Mike Cohn," [Online] Available: https://www.mountaingoatsoftware.com/agile/user-stories, 2016, [Accessed on: 26 April 2016].

An explanation of how a User Story should look, when utilizing them to plan tasks for use with implementing functions of my application.

[18] Phillip Webb, Dave Syer, Josh Long, Stphane Nicoll, Rob Winch, Andy Wilkinson, Marcel Overdijk, Christian Dupuis, Sbastien Deleuze, "spring-projects/spring-boot," [Online] Available: https://github.com/spring-projects/spring-boot, 2016, [Accessed on: 27 April 2016].

The GitHub repository of Spring Boot and documentation for using Spring Boot. Used by me in order to build my application as a web service with Java using Spring.

[19] Pivotal, "spring.io," [Online] Available: https://spring.io/, 2016, [Accessed on: 1 May 2016].

The Spring Framework, allowing me to create a web application service using Java for my project.

[20] R. Schmieder and R. Edwards, "Quality control and preprocessing of metagenomic datasets," *Bioinformatics*, vol. 27, no. 6, pp. btr026–864, Jan. 2011.

A tool that does what my project topic describes, with more understanding than I had at the start of my project. An interesting look at what other software is out there to do what my application also wanted to do and was a baseline for me analysising what sort of techniques I may wish to use to give the user of my own application a report on the quality of their assembly.

[21] K. Schwaber and J. Sutherland, "The Scrum guide," 2001.

The Scrum Guide, useful for knowing how to use the Scrum framework as part of my projects lifecycle model.

[22] J. C. Segen, The dictionary of modern medicine. Parthenon Pub. Group, 1992, 1850703213.

For the reference of what GC Content is. The most referenced link seen else where online (e.g. from Wikipedia) is to a web link that no longer exists. This reference backs up the description of GC Content, and is how I understand what GC Content is outside of Wikipedia's reference to the missing web link.

[23] Tatiana Tatusov and Roman Tatusov, "ORF Finder," [Online] Available: http://www.ncbi.nlm.nih.gov/projects/gorf/, 2016, [Accessed on: 25 April 2016].

A tool from NCBI for finding Open Reading Frames within a pasted sequences. I liked using this to compare my own ORF results against and get an idea of how I might want to display my own results in the quality report too. It was also nice to use for helping me understand what an Open Reading Frame actually is.

[24] The Board of Regents of the University of Wisconsin System, "Translation and Open Reading Frames," [Online] Available: http://bioweb.uwlax.edu/GenWeb/Molecular/Seq_Anal/Translation/translation.html, 2008, [Accessed on: 25 April 2016].

I started to understand what an Open Reading Frame was from reading this web page. It helped me get a grasp of what ORFs were, how they were constructed and what I should do to write code for my own application in how to find ORF Locations.

[25] The jQuery Foundation, "jQuery," [Online] Available: https://jquery.com/, 2016, [Accessed on: 1 May 2016].

jQuery, used for the code for tabs in CSS3, and for some click detection.

[26] T. Thomas, J. Gilbert, and F. Meyer, "Metagenomics - a guide from sampling to data analysis." *Microbial informatics and experimentation*, vol. 2, no. 1, pp. 3+, 2012.

A useful article in understanding what metagenomic study was. It was helpful to read as I tried to understand metagenomics at the beginning of the project.

[27] J. D. WUITSCHICK and K. M. KARRER, "Analysis of genomic g + c content, codon usage, initiator codon context and translation termination sites in tetrahymena thermophila," *Journal of Eukaryotic Microbiology*, vol. 46, no. 3, pp. 239–247, 1999. [Online]. Available: http://dx.doi.org/10.1111/j.1550-7408.1999.tb05120.x

An article that includes a discussion of the increase in GC count within coding regions. Useful for me to understand why using Open Reading Frames could be good for a user to see GC regions that were higher with potentially explainable reasons within their contiguous reads.

[28] Zhang Lab, "FASTA format," [Online] Available: http://zhanglab.ccmb.med.umich.edu/FASTA/, 2016, [Accessed on: 25 April 2016].

A description and example of what a FASTA file is and what they do. Gives a nice example of a fake FASTA file and helped me know what to expect with the file format when developing my application.