

## ABERYSTWYTH UNIVERSITY

Computer Science and Statistics (GG34) CS396: MINOR PROJECT

# **Application of Machine Learning Techniques to Next Generation Sequencing Quality Control**

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Draft April 30, 2014

### **Declaration**

I certify that except where indicated, all material in this thesis is the result of my own investigation and references used in preparation of the text have been cited. The work has not previously been submitted as part of any other assessed module, or submitted for any other degree or diploma.

Sam Nicholls 2014

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# **Chapter 1**

# Introduction

Over the past few years advances in genetic sequencing hardware have introduced the concept of massively parallel DNA sequencing, allowing potentially billions of chemical reactions to occur simultaneously, reducing both time and cost required to perform genetic analysis[1]. However, these "next-generation" processes are complex and open to error[2], thus quality control is an essential step to assure confidence in any downstream analyses performed.

### 1.1 Project Aims

The project consists of two sub-projects:

- Analysis of a current quality control system in place
- Identification of quantifiable sample properties that affect downstream analysis

### 1.1.1 Analysis of Current System

With the support of the Wellcome Trust Sanger Institute in Cambridge, this project works with the Human Genetics Informatics team to investigate **auto\_qc**[3], the institute's current automated quality control tool.

During genetic sequencing a large number of metrics are generated to determine the quality of the data read from the sequencing hardware itself. As part of the institute's vertebrate sequencing pipeline[4], **auto\_qc** is responsible for applying quality control to samples within the pipeline by comparing a modest subset of these metrics to simple hard-coded thresholds; determining whether a particular sample has reached a level that requires a warning, or has exceeded the threshold and failed entirely. Whilst this catches most of the very poor quality outputs, a large number of samples are flagged for manual inspection at the warning level; a time consuming task which invites both inefficiency and error[5].

In practice most of these manual decisions are based on inspecting a range of diagnostic plots, which suggest that a machine learning classifier could potentially be trained on the combinations of quality control statistics available to make these conclusions without the need for much human intervention.

The first part of the project aims to apply machine learning techniques to replicate the current **auto\_qc** rule set by training a decision tree classifier on a large set of these quality metrics. The idea is to investigate whether these simple threshold based rules can be recovered from such data, or whether a new classifier would produce different rules entirely. During this analysis it is hoped the classifier may be able to identify currently unused quality metrics that improve labelling accuracy. An investigation on the possibility of aggregating or otherwise reducing the dimensions of some of the more detailed quality statistics to create new parameters will also be conducted.

The goal is to improve efficiency of quality control classification, whether by improving accuracy of pass and fail predictions over the current system or merely being able to provide additional information to a lab technician inspecting samples labelled with a warning to reduce arbitrary decisions.

### 1.1.2 Identification of Properties that affect Downstream Analysis

The other half of this project is motivated by the question "What is good and bad in terms of quality?"

To be able to classify samples as a pass or a fail with understanding, we need an idea of what actually constitutes a good quality sample and must look at the effects quality has on analysis performed downstream from sequencing. An example of such is **variant calling** – the process of identifying differences between a DNA sample (such as your own) and a known reference sequence.

Given two high quality data sources where DNA sequences from individuals were identified in two different ways (one of which being next-generation sequencing) it would be possible to measure the difference between each corresponding pair. Using this, we could investigate the effect of leaving out part of the next-generation sample during the variant calling process. If we were to leave a part of a sample out of the variant calling pipeline would the variants found be more (or less) accurate than if it had been included? Would they agree more (or less) with the variants called after using the non next-generation sequencing method?

Having identified such sub-samples, can quality control metrics from the previous part be found in common? If so, such parameters would identify "good" or "bad" samples straight out of the machine. Samples that exhibit these quality variables will go on to improve or detriment analysis.

### 1.2 Project Method

### 1.2.1 Methodology

Clearly some team-based practices invited by agile methodologies – pair programming immediately comes to mind – are not applicable in a solo project. It is also unreasonable to expect an "on-site" customer for this particular project. In *The Case Against Extreme Programming*, Matt Stephens describes a "self

referential safety net" where the perceived traps in each practice are supported and "made safe" by other extreme programming (XP) practices. This would rule out XP as a viable methodology for a solo project as cutting out some of the processes that allow this form of evolutionary design to work (and flatten that cost-of-change curve) can introduce serious flaws to the management of a project and potentially result in failure. In the same breath it is important to remember that not all agile processes need be discarded just because XP seems incompatible. Indeed, some processes are common sense, for example: frequent refactoring, simple design, continuous integration and version control. Test driven development could also prove a useful process to consider as part of a methodology for this project as setting up a framework that allows for quick and frequent testing (before coding) and ensuring that any refactoring has a positive (or at least non-negative) effect on the system could be a worthwhile contribution to efficiency.

Could a more plan driven approach or form of agile-plan hybrid be considered appropriate here? In *Balancing Agility and Discipline*, Boehm and Turner introduce the idea of "homegrounds" for both agile and plan driven approaches; noting here that for projects that require high reliability and feature a non-collocated "CRACK customer" in fact align with some of these homegrounds for plan driven development. Combined with the thought that the project requirements will also be relatively stable it would seem that there may be no reason to switch to a more agile methodology as its primary feature is the welcoming of change that is not even needed? Perhaps this is the naivety of an optimist.

Personally I think I would approach this with a form of agile-plan hybrid; I like the idea of quick iterations and getting feedback as opposed to leaving acceptance testing until the end of the project, but I also want a somewhat detailed feedback process. In Neil Taylor's *Agile Methodologies* course it was suggested that it is dangerous to pick and choose processes (don't anger the Ring of Snakes) and also merely paying "lip service" to agile must be avoided (otherwise what's the point?), I feel that on this occasion it can be justified by the size of the project itself.

This project will consist of many research steps, each requiring some form of computational process to prepare the data for the next step. Whilst the implementations of the algorithms themselves pose computational complexity, there appears to be little challenge from a planning perspective and in fact a looser overall plan should be considered as we must account for unforeseen and unexpected outcomes from each research step.

The most important part of ensuring this project stays on track will be the development of a sensible testing methodology to ensure we are not only moving in the right direction in terms of which algorithm and parameters to use but also in terms of reliably measuring performance over time in a way that allows justification of such design choices.

Despite this trail of thought, given the research grounding this project entails it might be required to look beyond traditional and even modern software development methodologies and investigate a more scientific approach. A simple scientific method would involve establishing a null hypothesis that can be proven false by testing (e.g. "auto\_qc classifier is more accurate than the new classifier") and executing experiments that attempt to prove this null hypothesis false in favour of an alternative hypothesis (typically the opposite, e.g. "The new classifier is more accurate than the old classifier"). This form of hypothesis testing could essentially become the project's acceptance tests (providing we have an empirical definition of what "more accurate" means in terms of this system) and any modification can be classed as an experiment ("Do these parameters allow us to reject the null hypothesis?"). Although care must be taken not to let this descend into unstructured

cycles of mere hack-and-test, code-and-fix style programming.

Overall it is rather difficult to select a methodology for a project such as this as the research element makes it almost impossible to draw on previous personal experience for ideas of what development processes would be effective.

### 1.2.2 Task Management

It is useful to be able to keep track of current tasks preferably via a medium that would allow some method of sorting and filtering. For various personal projects and the second year group project I have used **Redmine**, a Ruby-based web application designed for bug tracking. However over time I have come to find keeping the task information stored in Redmine a task in itself. Attempts to extend the platform to implement additional functionality have been fruitless.

Out of frustration with thus and other alternatives – including **TaskWarrior** whose simple but effective command line interface was overshadowed by its occasional storage corruptions, I wrote my own open source web based task management application; **Triage**[6]. This will be useful in keeping track of current objectives and allow prioritisation in a quick and simple manner.

The list used to organise my project is also publicly available[7] for transparent progress tracking by my supervisor and those interested at the Sanger Institute.

### **1.2.3** Time Considerations

It must be remembered that this project needs to meet the requirements for a *minor* project and is to be completed alongside the study of several other modules which each have their own assignments and obligations. It would be easy to become overly ambitious and thus aims and goals will need to be revised as both obstacles and breakthroughs are encountered over the lifetime of the project.

# Part I Analysis of Current System

# Chapter 2

# **Introduction and Background**

### 2.1 Introduction

This part of the project can be outlined as follows:

- Collect data sets on which a machine learning classifier is to be trained
- Construct a program capable of processing and storing such data sets such that required subsets of the data can be quickly and easily returned for further analysis
- Select a suitable machine learning framework to handle the training and validation of a classifier
- Ensure a robust validation methodology exists for assuring quality of our own results
- Set up an environment capable of allowing results from such a classifier to be stored and compared
- Training a suitable classifier on the collected data sets
- Perform experiments by selecting subsets of the variables and observations and measure whether classification accuracy is improved

### 2.2 Concepts and Terminology

### 2.2.1 DNA and Bases

...DNA is made up of four bases; G, C, T and A...

### 2.2.2 Next Generation Sequencing

...the process of recovering or extracting the sequence of nucleotide bases...

### 2.2.3 Single Nucleotide Polymorphisms

...SNPs...

### 2.2.4 Samples, Lanes and Lanelets

A **sample** is a distinct DNA specimen extracted from a particular person. For the purpose of sequencing, samples are pipetted in to a *flowcell* such as the one in Figure 2.1 - a glass slide containing a series of very thin tubules known as **lanes**. It is throughout these lanes that the chemical reactions involved in sequencing take place.

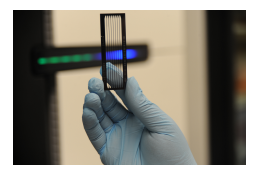


Fig. 2.1 An Illumina HiSeq Flowcell[8]

Once inserted, samples are amplified *in situ*, in the flowcell itself. A process in which the genetic material of each sample is caused to multiply in magnitude to form a dense cluster of the sample around the original. Millions of clusters will be created throughout each lane of the flowcell.

Note that a lane can contain more than one sample and a sample can appear in more than one lane; this is known as *sample multiplexing* and helps to ensure that the failure of a particular lane does not hinder analysis of a sample (as it will still be sequenced as part of another lane).

The more abstract of the definitions, a **lanelet** is the aggregate read of all clusters of a particular sample in a single lane. Figure 2.2 attempts to highlight examples of a this (circled in blue – not all lanelets are highlighted). For example Lane 5 shows the four clusters (in reality there would be millions) of Sample A combine to represent a lanelet. A lane will have as many lanelets as it does samples.



Fig. 2.2 Example of flowcell with some lanelets highlighted

# Chapter 3

# **Materials and Methods**

### 3.1 Input Data and Format

### 3.1.1 "BAMcheckR'd" Data

As part of the project I have been granted access to significant data sets at the Sanger Institute, unlocking quality control data for two of the largest studies currently undergoing analysis. A wide array of quality metrics are available for each and every lanelet that forms part of either of the two studies, totalling 13,455 files.

The files are created by **samtools stats** – part of a collection of widely used open-source utilities for post-processing and manipulation of large alignments such as those produced by next-generation sequencers that are released under the umbrella name of "SAMtools"[9] (Sequence Alignment and Map Tools). **samtools stats** collects statistics from sequence data files and produces key-value summary numbers as well as more complex tab delimited dataframes tabulating several metrics over time.

The output of **samtools stats** is then parsed by an in-house tool called **bamcheckr**<sup>1</sup> which supplements the summary numbers section of the **samtools stats** output with additional metrics that are later used by **auto\_qc** for classification. This process appends additional key-value pairs in the summary numbers section. A truncated example of a "bamcheckr'd" file can be found in Appendix A.1.

It is these summary numbers that will be the main focus of our learning task.

### 3.1.2 auto\_qc Decision Data

To use these "bamcheckr'd" files for training and testing a machine learning classifier, it is necessary to map each file to a classification result from **auto\_qc**. The one-to-one mapping between each input file and its label

<sup>&</sup>lt;sup>1</sup>Named such as **samtools stats** now incorporates **bamcheck** and the tool is written in R

are provided by the Sanger Institute in a separate file hereafter referred to as the AQC Decision Matrix or AQC (Decision) File.

A truncated example of such a file can be found in Appendix A.2. Only the first few columns are included – indeed we are only interested in the *lanelet* and *aqc* fields which provide an identifier that maps the row to a given input file and its classification by **auto\_qc** respectively. Latter columns pertain to a breakdown of decisions made by **auto\_qc** which are not included in the example for confidentiality (and brevity).

### 3.2 Development Environment

### 3.2.1 Language

Python was selected for the language of the program designed to handle this vast array of input data, more out of personal taste rather than a detailed analysis of required performance and features. From previous experience I was happy with the performance of Python when processing large datasets in terms of both file handling operations and storing the data in memory for later use. Python's generous choice of both built-in and third-party libraries have proven useful. Due to its concise and flexible nature it is possible to rapidly develop applications and its readability eases ongoing maintenance; useful given the short time-span allocated for this project and the possibility of others wishing to contribute to the project codebase after completion.

Whilst the choice was made primarily on preference, this is not to say other options were not considered: a highly popular Java-based collection of data mining tools, **WEKA**[10] would certainly have provided a framework for building decision tree classifiers but did not appear to offer any significant features that were unavailable elsewhere, whilst Java itself has the added constraint of requiring a virtual machine to be installed which could be undesirable from a performance or security standpoint when the application is deployed to servers at the Sanger Institute.

Difficulty was also encountered finding example implementations for **WEKA** with most documentation and tutorials providing information for performing analysis via the graphical "Explorer" interface instead, which would not be appropriate for quickly setting up and repeating experiments automatically.

Given the quality data we'll be using to train a machine learning classifier is output from the previously mentioned R script, **bamcheckr**, it was worth briefly investigating the options available for R itself as the potential of integrating the learning and predicting functions right in to the same process that outputs the data seemed convenient.

Whilst the **tree**[11] and **rpart**[12] packages are available for constructing decision trees in R (and actually **RWeka** provides an R interface to **WEKA**) neither appeared to be as robust as other more well-known frameworks. Also putting it politely, the programming paradigm of R[13] is rather different to other languages and can significantly increase development time (and frustration[14]) if one is not well versed in the patterns and grammar of the language and it seemed best to stick to one's comfort zone given the brief timescale for the project.

Had performance been a critical decision factor, lower level languages such as C, C++ or even Fortran could be

used. Briefly looking at frameworks available for C++ in particular, two popular solutions include: **dlib**[15], which did not support tree-based classifiers but did offer implementations of other potentially useful algorithms such as multi-class support vector machines; and **Shark**[16], which supported both decision tree and random forest classifiers. Both packages also provided a series of utility functions for performing mathematical or statistical operations on vectors of data.

### 3.2.2 Framework

Having studied the *Machine and Intelligent Learning* module in my final year, the prospect of getting stuck in to the deep of a machine learning algorithm was exciting. However the reality is a lot of cumulative time and effort has gone in to the creation and optimisation of a framework, which is unlikely to be surpassed successfully by a short-term one-person project. Thus utilisation of a third party machine learning library seems a wise investment for the project's codebase.

There are clearly numerous machine learning frameworks available in many languages, some of which were touched upon in the previous section and formed part of the development environment decisions. Whilst it is obviously unnecessary to select a framework which uses the same language as the project, it seemed counter-intuitive to select otherwise, for the establishing of additional arbitrary output and input steps to move data between the two environments could impede prompt experiment repeatability and introduce error.

A mixed bag of machine learning frameworks exist in Python[17], ranging from several highly active general purpose libraries to a multitude of smaller projects that focus on one specific learning task.

I investigate two of the larger libraries; **scikit-learn**[18] and **Orange**[19], primarily as their general purpose nature will allow exploration of various classifier solutions (given time) without the need to integrate many packages together, but also for their built-in functionality that aids the measuring of classifier performance and accuracy. These two packages were also selected for investigation on their recommendation from the project supervisor.

**scikit-learn** is somewhat the "new kid on the block" in terms of machine learning packages for Python, originally starting as a Google Summer of Code project in 2007 under the name of **scikits.learn**<sup>2</sup> the package was not officially published until 2010 where it quickly gained popularity and built an international team of contributors[20].

Arguably one of the most useful features of **scikit-learn** is its seamless integration with the "big names" of scientific computing in Python: **NumPy**[21] and **SciPy**[22], making heavy use of the optimised data structures and mathematical algorithms these packages offer, not only providing improved performance against packages that implement their own structures but also easing interaction by not requiring users to populate abstruse structures with their data or implement such algorithms themselves.

**scikit-learn** offers support for a wide range of algorithms covering many different machine learning tasks, including decision trees and random forests. Also of interest are the various utility subpackages, taking particular note of one for measuring the performance of the various classifiers; including functions for executing

<sup>&</sup>lt;sup>2</sup>Named as a **SciPy** Toolkit for Learning Tasks

cross-validation and generating confusion matrices; and another which assists with the selection of parameters (features) to use in a machine learning model.

The library boasts a growing community of users and a dedicated team of developers<sup>3</sup> who submit hundreds of commits to the project repository each month according to Github Pulse.

**Orange**, like **scikit-learn** is designed to be a general purpose machine learning framework and provide a wide array of tools to work with many different types of problems. Setting it apart from many of the other packages is the inclusion of a graphical user interface (GUI) that presents drag-and-drop-eqsue access to widgets from which users assemble a workflow to apply to their data. The GUI also allows visualisation of elements such as decision trees or confusion matrix plots without any effort from the end user.

Despite the repository commit history stretching back to mid 90s, only recent improvements including a major redesign of the GUI, significant changes to its object hierarchy and its 2013 academic publication have brought more attention to the framework.

At first glance **Orange** appears to offer a larger collection of functions to work specifically with classification tasks when compared to **scikit-learn**. Indeed it would later be discovered that it ships with features useful to the project that have not been implemented in **scikit-learn** including tree pruning and printing.

However it does not feature the same integration with Python's scientific computing packages; **NumPy** and **SciPy** and also requires that end users input their data in a structure defined by the framework rather than a generic data structure. Thus use of **Orange** would introduce more data wrangling and munging steps to ensure that data is represented in the correct format for a classifier to perform its learning and prediction role and also to interpret returned objects.

Although **Orange** affords a suitable (and in some cases better) range of functions to use for decision trees, I was concerned at the push to use its GUI as opposed to the package API, potentially impeding experiment repeatability by having to reset the graphical environment each time.

Despite the **scikit-learn** package not yet reaching a major version milestone, its feature set, interaction with **NumPy** and **SciPy** and useful built-in subpackages led me to choose **scikit-learn** as the machine learning framework for this part of the project.

### 3.2.3 Additional External Libraries

As discussed, the project will make use of the open source<sup>4</sup> **SciPy Stack**, consisting of several core packages including **NumPy**[21]: a package for efficient numerical computation which defines generic N-dimensional array and matrix data containers; and **SciPy**[22], which provides a multitude of optimised numerical routines.

Whilst the stack also includes **Matplotlib**, a highly popular graphing package, I typically prefer to use the R package **ggplot2**[25] for creating graphs. However it is useful to note that **Matplotlib** integrates easily with the data structures of **NumPy** (and thus **scikit-learn** too).

<sup>&</sup>lt;sup>3</sup>Some full time developer positions are funded by *INRIA*, the French Institute for Research in Computer Science and Automation

<sup>&</sup>lt;sup>4</sup>Components of the SciPy Stack are distributed under the 3-clause Modified BSD License[23][24]

**argparse** is a third-party Python library<sup>5</sup> used for specifying the arguments and options of a command line interface via a simple API. **argparse** spares the developer from having to check the presence and validity of a user's arguments to a Python program themselves whilst also automatically generating a friendly interface to the program for the end user based on the definitions provided by the developer.

### **3.2.4** Testing

As discussed in Chapter 1.2.1, testing forms a critical part of the project given the need to monitor the impact of changes to classification accuracy as well as to ensure the program is working correctly. Ideally, execution of a test suite should be simple and easily repeatable. Results that pertain to accuracy should also be stored for future reference to monitor ongoing performance of the classifier.

Such requirements could be fulfilled by a continuous integration platform – a server dedicated to the building and testing of the code contained in a centralised repository typically to which an entire team will have write access[27]. Whilst in this scenario there will be much less "risk" from integration issues due to the single person team size, the themes of automated building and self-testing code can still be taken on board.

**Jenkins**[28] is a highly popular[29] example of such a platform with which I am familiar. Although an out-of-the-box **Jenkins** instance is suitable for variety of software engineering projects, it would be necessary to invest some time to install and tweak plugins to perform actions on test results (such as failing a build that causes accuracy to decrease). Previous experience found that more specific tasks will often require a plugin to be authored to overcome limitations in the feature set of a more generic plugin, which given the intricacies of the **Jenkins** package layout could easily turn in to a project of its own. Unfortunately, other features that would be useful to the project including the indexing and searching of logs are somewhat lacking in **Jenkins**.

Online solutions such as **Travis**[30] and **Wercker**[31] are free for open source projects and could potentially offer a quicker set up, as both services merely require a small configuration file in the root of the repository and a hook to be registered (allowing builds to be triggered on a code push for example) before being able to deploy a build in the cloud.

However these online services would not have been able to handle the return of build artifacts (such as logs, graphs or dot files containing a decision tree) without some convoluted solution of uploading them to another service or committing them to a private code repository during execution of the job – the non-persistent nature of these nodes would cause any artifacts to be destroyed when the node is terminated at the end of the build.

Also, as these virtual nodes are isolated from any sort of persisting file system it would be necessary to upload large quantities of training data any time a build and test job is to be run. This sounds rather inefficient in terms of both bandwidth and time and would more than likely constitute unreasonable use in view of the platform's "fair use" terms and conditions too.

It would seem that none of the widely available solutions would provide an environment suitable for the testing of this project. I wanted to provide my own platform for this problem but given the time constraints of the

<sup>&</sup>lt;sup>5</sup>Although since Python 2.7, **argparse** has been included as part of the standard library[26]

project this was simply not practical and in the end we settled for well formatted log files which could be searched and processed with command line tools such as **awk**.

### **3.2.5** Tools

Version control is critical, git

# Chapter 4

# **Pre-Implementation**

### 4.1 Classification Correlation

An important consideration for statistical analysis is the relation between observations. The "bamcheckr'd" input data described in Chapter 3.1.1 is available per lanelet, however as shown in Chapter 2.2.4 a lane may contain more than one lanelet. Herein lies the trouble: if during a sequencing run the flowcell is somehow subjected to abnormal conditions (*e.g.* a temperature increase due to an air conditioning failure) or the device is depleted of reagents then every lane (and thus all lanelets within) will be of considerably poor quality.



Fig. 4.1 **Heatmap of lanelet QC status by lane**: Lanes are vertical bars with each lanelet cell coloured red to represent a failure, yellow for a warning and grey for a pass.

In such a case there would appear to exist a relationship between the respective qualities of each lanelet in

a lane as well as each lane in a sequencing run. To examine this further, an R script utilising **ggplot2** was authored to visually inspect whether correlation existed and if so to what degree that data is affected.

Figure 4.1 displays a plot of **auto\_qc** classification for each lanelet in a lane. The plot itself is a dense heatmap where each lane stands as a vertical bar, broken in to horizontal cells, each of which represents a lanelet that was sequenced in that particular lane. These lanelets are colour coded using; red for failures, yellow for warnings and grey for passes (to allow the other two classes to be more easily seen).

Therefore an unbroken vertical red line indicates that all lanelets that comprise of that line failed to pass some aspect of the current **auto\_qc** thresholds. In reality there are few conditions under which a lanelet would fail irrespective of the status of the rest of the lanelets in the same lane which typically involve an error during the preparation of the sample (an easy to spot result as it will cause poor quality across all lanelets using that sample).

Overall there are a series of instances appearing to support correlation for whole-lane failures and warnings but despite this there do appear to be occasions where a lanelet has failed where the remainder of the lane has not. Having discussed this with the project supervisor and contacts at the Sanger Institute we decided to continue to the implementation stage, agreeing that whilst some evidence of correlation between lanelets in the same lane has emerged, we will still be able to recover parameters that will be useful to quality control and statistical testing may be required following this analysis to describe how powerful such parameters are taking this possible correlation in to account.

It should be noted that the proportion of failures and warnings is considerably smaller than passes and so care will need to be taken to find a balance; for example it would not be feasible to merely discard lanelets from lanes that have failed entirely as there'd hardly be any data on which to train a classifier. Indeed other solutions may be possible, perhaps weighting observations which exhibit similar behaviour to other lanelets in their lane so as to give their parameter values less priority during the construction of the classifier itself.

It is worth noting that although the plot does not make a particular distinction between lanes in the same flow cell, lanes are sequentially identified so the red bars of thicker-width arguably display some failures across entire flow cells.

As a final note it should be stressed that this plot should be regarded as a diagnostic rather than an experiment with a direct conclusion. Given more time it would be useful to investigate the nature of these possible correlations, given a lane that has failed across all lanelets: do those lanelets actually express similar quality metrics?

### 4.2 Recovering Ratios

An initial scan of the summary numbers available in the "bamcheckr'd" files introduced in Chapter 3.1.1 revealed 82 different parameters. However, when comparing this parameter set to the threshold rules of **auto\_qc**, it appeared that some parameters were "missing".

In the pursuit of replicating the decisions of the existing system, it is ideal to have all the parameters used

in the making of those decisions at hand. In particular, the missing parameters were of a normalised nature, typically in the form of a ratio or percentage which should make them more valuable than a parameter that just represents a raw count of some property.

It was found that these missing parameters are calculated in a step preceeding **auto\_qc** as part of the **vr-pipe** pipeline<sup>1</sup>.

Whilst I could have attempted to set-up my own instance of **vr-pipe** to recover this data, speaking with contacts at the Sanger Institute, it was decided that this would prove troublesome work; requiring an involved deployment to a cloud based facility such as Amazon's Elastic Compute Cloud and installation of various Perl dependencies as well as requiring access to many controlled databases within the institute.

Most of the functions that calculated these parameters turned out to be straightforward and could easily be ported to another application. At first it was intended to add these functions to the program authored for this part of the project, but the Sanger Institute suggested it would be more useful to implement such functionality in **bamcheckr** directly, removing some of **auto\_qc**'s dependence on its position in **vr-pipe**.

```
# Install and include the 'devtools' package
install.packages("devtools")
library(devtools)

# Install package directly from Github repository
install_github("samstudio8/seq_autoqc", subdir="bamcheckr")

# Install package from local directory
install("/home/sam/Projects/seq_autoqc/bamcheckr")
```

Listing 1: Installing an in-development R package with **devtools** 

With the help of **devtools**[32] (see Listing 1) it was simple to develop and test contributions to **bamcheckr** locally without needing to re-publish the package after changes. An additional script was added to **bamcheckr**'s NAMESPACE... to recover the missing ratio and percentage based parameters...

...unfortunately, the performance of the script was poor, taking 5.5s on average and increasing to 16.1s when enabling a complex function containing many vector operations (potentially inefficient due to my lack of R experience)... for overlapping\_base\_duplicate\_percentage

...utilising the program designed and implemented in the following chapter... loaded required information and used the API to calculate the missing parameters...

```
...Morandat[33]
```

...with such an intriguing difference in performance with significantly more time it would be appealing to explore the idiosyncrasies of each implementation ...although such an investigation would quite likely sufficiently

 $<sup>^{1}</sup> https://github.com/wtsi-hgi/vr-pipe/blob/hgi-release/modules/VRPipe/Steps/vrtrack\_auto\_qc\_hgi\_3.pm$ 

generate an entire project of its own.

### 4.3 Contributions to bamcheckr

...R CMD BATCH issue ...Fixed a graph plotting failure.

# Chapter 5

# **Frontier**

This chapter introduces **Frontier**: the main programming effort for this part of the project. Frontier is a Python package that serves as a data manager, providing both interfaces to read inputs into structures in memory and to retrieve them in formats acceptable to a machine learning framework.

### 5.1 Design

### 5.1.1 Purpose

Frontier's purpose is to supplement analysis with **scikit-learn** by allowing a user to read in and parameterise data in a format that can then be used for analysis by the **scikit-learn** library. Frontier was designed to simplify the process of setting up machine learning tasks and enable experiment repeatability by removing the need for users to spend time constructing classes and functions to parse their input data and to just get on with analysis using **scikit-learn**.

Initially Frontier was to act as a wrapper around **scikit-learn**, essentially removing the end user's interaction with the library and merely providing an interface for data to be passed in and some sort of classifier to be returned. However this quite clearly limited Frontier's audience by tying it to a particular framework and would quickly become unmanageable in the task of providing wrappers for all aspects of an external library.

Instead it was decided that Frontier would be used alongside a user's chosen machine learning framework, providing a useful API to parse and extract observations and variables from input data and arrange them in structures suitable for processing with **scikit-learn**.

If possible, Frontier could also handle any objects returned, displaying or logging any textual or graphical information pertaining to a returned classifier's accuracy to assist a user in the ongoing performance monitoring of changes to used data subsets or parameters.

### **5.1.2** Format

Frontier is designed as a Python package, allowing a user to import its functionality in to other programs. The result of this project's technical output could almost be considered as two separate entities: Frontier itself, the package designed to ease user interaction with scikit-learn and **Front**, a Python script which implements Frontier's functionality in order to interact with scikit-learn to conduct analysis on the **auto\_qc** data.

### **5.1.3** Method

.. the result of abstracting code and tools created during the use of **scikit-learn** for analysis of the current **auto\_qc** system .. initially hard-coded to suit the specific needs of the project but followed an evolutionary design process...

designed specifically for the given learning problem, with hard coded classes and encodings...

### 5.2 Concepts

In this section we introduce some terminology and ideas that will assist understanding of the application's purpose and terminology used in the following implementation section.

### **5.2.1** Observations and Parameters

An **observation** refers to a distinct object or "thing" from which we will attempt to understand the relationship between its known properties and classification to be able to label future observations with unknown class based on those properties. In the context of this project, this would be a lanelet. Other documentation in the field may use the term *sample* but to avoid confusion with the definition of sample introduced in Chapter 2.2.4 we will use the term observation.

The aforementioned *properties* or *attributes* of an observation will be referred to as the **parameters** of that observation. Traditionally these may be described as *features* but it was felt that this wording may have connotations with discrete data. Early versions of Frontier referred to parameters as *regressors*, using terminology from statistical modelling. This wording was dropped to remove any confusion between regression and classification machine learning problems.

### **5.2.2** Data and Targets

**Data** will be used somewhat generically to refer to a matrix in which observations act as rows and their parameters act as columns. It is expected that all observations will have the same set of parameters.

For problems of a classification nature such as ours, an observation's **target** is the known classification of that particular observation. Depending on the context of the learning problem targets may be discovered manually

(e.g. counting leaves from images of plants, one will need to manually count all the leaves in the image before being able to use it for learning) or as the output from another system such as **auto\_qc**.

In the context of this project where the objective is to begin learning the rules of **auto\_qc** the targets refer to whether a particular lanelet observation was labelled as a pass, fail or warn by the current system.

### **5.2.3** Class Labels and Codes

An observation's target may also be known as its **label**. A labelled observation is said to be a member of that label's **class**. For example a lanelet that has failed **auto\_qc** is said to be labelled as a fail and is a member of the class of failures.

String based class labels can pose difficulties when handling data later on, not only taking more memory to store as opposed to a type such as an integer but also introducing accidental subsetting or discarding of data whose labels differ unexpectedly. For example are "fail", ,"fial", "FAIL" and "Failed" supposed to be members of the same class?

Often such labels are **encoded** in to simpler types such as integers.

### 5.2.4 Training and Testing

...

### **5.2.5** Python Data Structures

...Python dictionaries offer (constant) lookup access and more importantly querying the structure for whether it contains a particular key can also be performed in constant O(1) as opposed to searching a list for membership[34]...

a structure in which hashed keys are mapped to an arbitrary value or object.

### **5.3** Implementation

This section investigates the implementation of Frontier's major components:

- Informing the package of the problem domain...
- Interfaces provided for reading in data
- Storage of read data in memory
- Retrieval of stored data
- · Interaction with scikit-learn

### **5.3.1** Class Definitions

Frontier was designed to support classification machine learning problems, to adequately support this task, the package must be aware of each of the possible classes in the problem space. Early versions of Frontier were specifically designed for training and testing data from **auto\_qc** and could only support encoding and decoding of the pass, fail and warn classes.

However this implementation was clearly esoteric and held little to no further use outside the domain of the project's learning task. What would happen if a class label were to be added or removed in future? Most likely many lines of code would need to be re-written to handle such cases; the package was inflexible.

To counter this, Frontier was refactored to remove hard coded label definitions enabling its use as a more general purpose tool where users can specify the domain's classes and their associated labels and encodings. Listing 2 shows the definitions used by Front to work with the **auto\_qc** data.

```
CLASSES = {
    "pass": {
        "names": ["pass", "passed"],
        "code": 1,
    },
    "fail": {
        "names": ["fail", "failed"],
        "code": -1,
    },
    "warn": {
        "names": ["warn", "warning"],
        "code": 0,
    },
}
```

Listing 2: Class definitions for auto\_qc as passed to Frontier

As per Listing 2, to define classes a user must provide a Python dict containing the following for each label:

· class String

The dictionary key is used as the canonical name of the class label

• names [String]

A list of labels that denote membership of this class

• code Integer

The encoded representation of this class (See Chapter 5.2.3)

• \_recode Boolean, Private

A flag to indicate whether an API action has changed the code of this class, typically used when treating

all members of one class as a member of another – A user should never set this manually

### • \_count Integer, Private

The number of observations with this label, typically used when calculating weightings based on the proportions of class sizes and outputting logging information – A user should not set this manually

This simple structure allows Frontier to be compatible with almost any classification learning task with minimum input from the end user. Additional utilities provided by the Frontier utils subpackage use this structure to automatically classify and encode labels without user intervention with the following functions:

### classify\_label

Attempt to classify a label by comparing a given string to each set of *names*, locating an exact match will return the relevant canonical class label

### · encode class

Given a canonical class label, return its code

### decode class

Given a *code*, return the canonical class label unless *recode* is True

### · count class

Increment the *\_count* for a particular class given its canonical label

These functions are put to use throughout Frontier and are essential for reader classes (detailed in the next chapter) to parse, classify and then encode observation targets automatically from relevant files.

### 5.3.2 Input Handling

Frontier's modular nature allows users to write their own Python classes to read data from any form of input file or stream. Two examples of which are the classes used to read from the "bamcheckr'd files" documented in Appendix A.1 and the **auto\_qc** decisions matrix briefly demonstrated in Appendix A.2.

These classes are described as **Readers** and implement a common base class, **AbstractReader** which takes care of setting up the file handler, including functions to both close and iterate over the file's contents. It will also automatically call its own **process\_file** function that skips over any header lines before passing each line in the file stripped of any newline characters to **process\_line**.

It is expected that derived classes will at least provide their own implementations for **process\_line** and **get\_data**. Failing to do so will cause Frontier to throw a **NotImplementedError** when attempting to use the class to read data.

**process\_line** defines the line handling operations that extract and store desired data found in a given line. This responsibility includes returning None for lines that contain comments and irrelevant data.

**get\_data** must return any read in data in a suitable structure for storage by Frontier. Typically this will be a Python dictionary using some unique identifier for each observation as a key, mapping to an arbitrary value or

object containing that observation's parameters. Further discussion on Frontier's storage of data and targets is to follow.

The **AbstractReader** class is designed to simplify the process of reading in observations and their targets for end users, however it is still up to the author of the dervied class to set up any structures to store data (which cannot be done automatically without likely enforcing potentially unhelpful constraints) before initialisation of the inherited base class (via the call to **super**) as shown in Listing 3.

```
class BamcheckReader(AbstractReader):
    [...]

def __init__(self, filepath, CLASSES, auto_close=True):
    self.summary = SummaryNumbers()
    self.indel = IndelDistribution()
    super(BamcheckReader, self).__init__(filepath, CLASSES, auto_close, 0)

[...]
```

Listing 3: Extract from **BamcheckReader** class documenting initialisation of necessary data structures and calling for initialisation of its inherited base class

As shown in Listing 3, the initialisation of the **AbstractReader** allows four arguments:

### · filepath String

A relative or absolute path to a file from which to extract data or targets

### • CLASSES Dictionary

A dictionary of user specified class labels defined as described in the previous chapter

### • auto\_close Boolean, Optional

Whether or not to close the file handle immediately after executing **process\_file**, this is True by default to prevent either memory leaking when users are reading in a large number of files and are perhaps unaware that they require closing or the throwing of an IOError caused by having too many file handles open at once

### • header Integer, Optional

The number of lines to ignore before the reader should begin passing stripped lines to **process\_line**, defaults to 0

Currently it is also the responsibility of the author of a derived class to perform relevant sanity checking of any extracted data. For example the **BamcheckReader** class checks for the presence of multiple entries of a particular metric which will print a notice if found, unless the entries have differing values, upon which an exception is thrown and the process is halted.

Once a reader has been defined for a particular file format, a user need only provide a directory of files to be parsed and the name of the class designed to complete the parsing. Frontier will then take care of executing

the parsing process on all files in the directory. After a derived reader has completed file handling, Frontier will call its **get\_data** function to "move" the extracted data to its own storage.

At this point Frontier will also check the integrity of the data, primarily that all parameters have a non-zero variance. Users will be warned when this requirement is violated; parameters with no variance cannot provide much information for successful classification as their values are equal for all class labels!

In future it would be useful to investigate whether it would be feasible to perform such sanity checking in a generic manner to ensure it could be applied to a wide enough range of scenarios to make it worth including functionality in the **AbstractReader** directly.

With more time, future improvements could overhaul the reader interfaces to allow users to simply specify the format of a file in a string that can be parsed by Frontier's IO subpackage, rather than having to write their own derived class. Classes could also list file extensions they are capable of processing which could potentially be used to automatically determine which readers to use without requiring the user to explicitly specify.

### 5.3.3 Storage

Frontier specifies a class called the **Statplexer**<sup>2</sup> which provides users with a single point of access to all read in data. The reader interfaces described in the previous chapter implement their own loading functions to populate the **\_data** and **\_targets** class members of the **Statplexer** object, both of which are Python dictionaries.

During the parsing of observation data with the relevant reader class, the reader is expected to locate an appropriate unique ID for each observation. In the case of processing of **auto\_qc** data, this would be the lanelet's barcode which is collected from the filename of that particular lanelet's "bamcheckr'd" file. This identifier is then used as a key in both the **\_data** and **\_targets** dictionaries to map to a structure (typically another dictionary or an arbitrary class) that stores that observation's parameters and its known target (encoded class label), respectively.

Although these attributes can be manipulated directly (and indeed they are for testing purposes) the leading underscore follows a popular convention defined in Python's style guideline, PEP8[35], where class members with leading underscores should be treated as non-public. Python doesn't have private variables such as those that may be found in other languages like Java, indeed the Python style guide points out that "no attribute is really private in Python"[35]. In an interesting StackOverflow answer on the subject, a user describes that this is "cultural"[36] and that Python programmers are trusted not to circumvent convention and "mess around with those private members". Users are therefore expected to use the functionality Frontier provides for controlled getting and setting of data stored in these pseudo-private \_data and \_targets members.

**load\_data**, for example, should be used exclusively when populating the two dictionaries and is automatically called on construction of the **Statplexer** if the construction arguments are valid. **load\_data** will automatically call other necessary (pseudo-private) functions of the class including **\_test\_variance** which checks that parameter variances are non-zero and also warns users if new observations are overwriting old ones, or if an observation does not appear to have a corresponding target.

<sup>&</sup>lt;sup>1</sup>Rather, a pointer to the address of the extracted data's dictionary hashmap will be copied to memory inside a Frontier class

<sup>&</sup>lt;sup>2</sup>A somewhat contrived contraction of 'Statistics Multiplexer'

The following section details the retrieval of data and targets from the **Statplexer** which are returned to the user in **NumPy** arrays. Why does **Frontier** not just store the read in data in such a structure to begin with instead?

This is primarily due to the underlying layout of a list structure, which must be represented in memory as a contiguous block. Great expense is therefore incurred if the list grows beyond its bounds and must be relocated elsewhere in memory to resize to a larger contiguous block. Given the nature of input data, the number of observations and parameters are unknown before the input data is actually read and so memory cannot be reserved to prevent these operations. The potential number of input files renders reading through the data once to reserve the right amount of memory for a second read-through impractical in terms of time.

It is arguable that despite this, the data could be unloaded from the \_data and \_targets dictionaries in to some form of array at the end of the call to load\_data. However as the Statplexer allows loading of additional data at any time, which would not only risk increasingly expensive resizing operations as the list continues to expand but the sanity checking that takes place in load\_data involves checking for membership of a given ID in \_data and \_targets which is constant for dictionaries and significantly slower for large lists (see Section 5.2.5).

It is possible that an implementation could use both a dictionary and an array; appending the observations to the array and entering a mapping between the observation ID and its index in the array. Though this would still not avoid the issue of relocating the array once it grows beyond its bound in memory.

Worse still, as results requested from the **Statplexer** API are returned to the user in a sorted matrix<sup>3</sup> (*i.e.* rows and columns are ordered alphanumerically by the observation ID and parameter name, respectively), if observations are not processed in a sorted order (which is not a requirement) then returning results will involve accessing the proposed **\_data** and **\_targets** arrays in a non-linear fashion, losing any efficiency that could be gained from using an array in the first place.

Whilst not entirely optimal<sup>4</sup>, in terms of a compromise between memory complexity and avoiding expensive operations on data input and retrieval, dictionaries offer the best balance.

It should be noted that the **Statplexer** stores a copy of the user defined CLASSES dictionary (as the class member **\_classes**), which is an argument to its construction, allowing the **Statplexer** to share this information with any readers or utility functions who require it.

### 5.3.4 Retrieval

The Statplexer is designed to provide functions to an end user for extraction of desired data in a format suitable for an external framework or library, in our case; scikit-learn...

...leverages numpy...

...Frontier can be used to inspect the parameter set extracted from the observations... some of the methods provided for this purpose include:

<sup>&</sup>lt;sup>3</sup>A two-dimensional **NumPy** array

<sup>&</sup>lt;sup>4</sup>Dictionaries still require use of **sorted** when creating a result matrix via the API

### · list\_parameters

Return a sorted list of all parameters

### find parameters

Given a list of input strings, return a list of parameters which contain any of those strings as a substring

### • exclude\_parameters

Given a list of input strings, return a list of parameters which do not contain any of the input strings as a substring, or if needed an exact match

...data is returned sorted by key, ...with parameters sorted alphanumerically... ...targets map 1:1 with the data array...

### • get\_data\_by\_parameters

Return data for each observation, but only include columns for each parameter in the given list

### get\_data\_by\_target

Return data for each observation that have been classified in one of the targets specified and additionally only return columns for the parameters in the given list

### 5.3.5 Interaction with scikit-learn

### **Cross Validation**

...method in which to measure classification accuracy... ...potentially use a weighting to penalise mistakes in smaller classes...

...K fold cross validation ...using stratified K fold cross validation...

### **Confusion Matrices**

"Normal" confusion matrix and "Warnings" confusion matrix...

### **5.4** Usage Example

```
from Frontier import frontier
from Frontier.IO import DataReader, TargetReader
data_dir = "/home/sam/Projects/owl_classifier/data/"
target_path = "/home/sam/Projects/owl_classifier/targets.txt"
CLASSES = {
        "hoot": {
            "names": ["owl", "owls"],
            "code": 1,
        },
        "unhoot": {
            "names": ["cat", "dog", "pancake"],
            "code": 0,
        },
}
statplexer = frontier.Statplexer(data_dir,
                                  target_path,
                                  CLASSES,
                                  DataReader,
                                  TargetReader)
```

Listing 4 : Example usage of Frontier

Constructing the Statplexer requires the following arguments:

### • data\_dir String

Root data directory under which all files will be parsed for observation data

### • target\_path String

Path to file to be parsed for observation targets

### • **CLASSES** Dictionary

A dictionary of user specified class labels defined as described in Chapter 5.3.1

### • DataReader Module

Module containing the class (of the same name) to be used to parse each file in the *data\_dir* tree for observation data

### • TargetReader Module

Module containing the class (of the same name) to be used to parse the target\_path file for target data

# 5.5 Testing

...

# Chapter 6

# **Results**

- 6.1 Introduction
- **6.1.1** Why Decision Trees?
- **6.1.2** CART

### **6.2** Parameter Selection

...important to find the "best" parameters ...what is best? scikit-learn uses total gini information ...frontier uses two methods:

- Backward elimination; pruning parameters with the lowest total gini
- Call scikit-learn's SelectKBest

<sup>\*</sup> incorrect degrees of freedom \* warnings: /usr/lib64/python2.7/site-packages/sklearn/feature\_selection/univariate\_selection.py: RuntimeWarning: invalid value encountered in divide, causing NaN \* Replaced univariate\_selection with version from master \* needed use force np.float64 \* ...actually data was 0... gg

- 6.3 Trees
- **6.3.1** Initial Trees
- **6.3.2** Parameter Sets
- **6.3.3** Ignoring Warnings

# Part II

# Identification of Qualitative Sample Properties

# **Introduction and Motivation**

## 7.1 Introduction

The second part of the project can be outlined as follows:

- Collation of variant locations...
- Construction of script to select a representative genomic region...
- Extraction of such regions...
- Merging of regions to one file...
- Establishment of pipeline...:
  - Select each lanelet in turn...
  - **–** ...
  - Call variants on k-1 lanelets...
  - Compare concordance...

## 7.2 Concepts and Terminology

### 7.2.1 Variants and SNPs

...for the purpose of this study we are only interested in single nucleotide polymorphisms (SNPs) "snips" ...

...their *position* refers to the index of the base of the chromosome in which the polymorphism exists... ...commonly these are 1-indexed (although this is not always the case) and thus care must be taken to ensure this is accounted for at any step where positions are considered...

## 7.2.2 The 'Goldilocks' Region

Having recovered from the miscommunication that led me to attempt to find reverse strands in VCF files with already known errors, I've been making performance improvements to the script that finds candidate genomic regions.

The task poses an interesting problem in terms of complexity and memory, as the human genome is over three billion bases long in total which can easily lead to data handling impracticalities.

For the next step of the project we are looking to document what effects quality has on analysis that occurs downstream from sequencing, for example: variant calling - the process of identifying bases in a sample that differ from the reference genome (this is a little simple as you also need to discern as whether the difference is actually a polymorphism or not).

To investigate this I'll be performing leave-one-out analysis on the whole-genome data we have, consisting of leaving a lanelet out, performing variant calling and then comparing the result of the left-one-out variant call and the corresponding variant call on our "SNP chip" data.

The basic idea is to answer the question of what constitutes "good" or "bad" lanelet quality? Does leaving out a lanelet from the full-sequence data lead to variant calls that better match the SNP chip data, or cause the correspondence between the two sets to decrease? In which case, having identified such lanelets, can we look back to the quality parameters we've been analysing so far and find whether they have something in common in those cases?

If we can, these parameters can be used to identify "good" or "bad" lanelets straight out of the machine. We know that lanelets that exhibit these quality variables will go on to improve or detriment analysis.

However, variant calling is both computationally and time intensive. Whilst the Sanger Institute have significant computing resources available, my dissertation has an end date and with the time I have we must focus the leave-one-out analysis on a subsection of the whole genome.

It is for this reason we're looking for what Josh at Sanger described as a "representative autosomal region". The candidate must not have too many variants, or too few: a "Goldilocks genome".

### **7.2.3** Groups

ichip vs gwas... ...will become clear later ...interested keeping the variants from the two data sets we have seperate...

### 7.2.4 Length and Stride

...length will refer to the number of bases included in the region ...stride will refer to the number of bases to be used as an offset between the starting base of one region and the beginning of the next...

...a length of 1000 and a stride of 500 for example ...the first region for example will consist of 1:1000, then 501:1500 inclusive

...ensure sensible choices of these parameters!

## 7.2.5 Candidates

...a **candidate** is a region of *length*... ...a candidate will contain a number of variants, counted for each *group*... ...simple object consisting of start and end base...

# **Materials and Methods**

## 8.1 Input Data and Format

## 8.1.1 Variant Call Format

```
# Install
git clone htslib
git clone bcftools
make #(requires htslib to be above samtools/bcftools dir)
sudo cp bcftools usr/local/bin
# Tabix
# Download from sourceforge
# http://sourceforge.net/projects/samtools/files/tabix/
sudo cp tabix /usr/local/bin
# Generate an index (vcfidx) (not necessary)
vcftools --gzvcf cd.ichip.vcf.gz
# Query VCF (can be done with awk, cut etc.)
vcf-query cd.ichip.vcf.gz -f '%CHROM:%POS\t%REF\t%ALT\n'
# ! A faster alternative to using vcftools exists in bcftools ! #
bcftools query -f '%CHROM: %POS\t%REF\t%ALT\n' cd-seq.vcf.gz > cd-seq.vcf.gz.q
\# Complains about lack of tabix index but still makes the file...
```

```
# Index with tabix
# "The input data file must be position sorted and compressed by bgzip
# which has a gzip(1) like interface"
tabix -p vcf file.vcf.gz
diff cd-seq.vcf.gz.tbi diff cd-seq.vcf.gz.tbi.sanger
# Shows no difference :)
# http://samtools.sourceforge.net/tabix.shtml
# http://ucftools.sourceforge.net/perl_module.html
# http://www.biostars.org/p/51076/
# http://ucftools.sourceforge.net/htslib.html#query
```

## 8.1.2 Variant Query File

...named such for the command that creates it, the Variant Query File is the proposed input format for **Goldilocks**... ...listing z shows the commands used to extract this information in the required format... ...line-delimited, tab-separated entries with the first field a colon-separated pair of a chromosome and position... (*e.g.* 1:5000 is the base at position 5000 on the first chromosome).

## 8.1.3 Paths File

...esoteric file format to handle grouping of Variant Query files... ...very simple, lines beginning with a '#' delimit the desired groups, any line following a group line is considered to be a member of that group... ...entries are tab-delimited, with two fields; a unique slug for simple identification of the file and its actual path (relative or absolute) respectively...

## 8.2 Development Environment

## 8.2.1 Language

Once again, Python was the language of choice for this script... ...for many of the reasons previously discussed in Chapter 3.2.1 ...we needed this quickly, Python is useful for fast prototyping... ...expected reasonable performance without the need for a lower level language like C...

## **8.2.2** Testing

...a little more difficult ...discussed more at length in the following chapter...

## **8.2.3** Tools

As before... (Chapter 3.2.5)

## **Goldilocks**

This chapter introduces **Goldilocks**; the first programming hurdle for this part of the project. **Goldilocks** is a Python script designed specifically to read in and store locations of user-selected variants on the human genome and to then find regions of a given size on the genome where variants are of a desired density.

## 9.1 Design

## 9.1.1 Purpose

Expanding on the introduction, **Goldilocks** is responsible for parsing input files that contain chromosome and base position entries.. Input files can be grouped, pooling the set of unique listed variants between the files and treating them as one search space.

For each file group, a list of positions for each chromosome is stored in memory. Once all variants have been loaded **Goldilocks** is designed to iterate over each chromosome<sup>1</sup>, load all of the variants from the lists in to an array and essentially count the number of variants present (in each group) between a given start and end position, determined by the arguments specified by the user as described in Section 7.2.4.

This combination of a start and end position and the counts of variants present inbetween forms the structure of a **candidate**. Having performed this search over all the autosomes, candidates can then be ranked by user-specified criteria regarding the density of the counted variants.

#### **9.1.2** Format

**Goldilocks** is a modestly sized but elegantly complex Python script which can also serve as a stand-alone module to be imported in to other programs. Whilst the use case is certainly more esoteric than **Frontier**, I

<sup>&</sup>lt;sup>1</sup>Strictly speaking we are only handling the 22 autosomes and ignoring the remaining 2 sex chromsomes (allosomes)

made the choice to modularise the functionality, in the hope that if an end user were to find some function of the script particularly useful they may call it as part of their own program.

With more time it would be interesting to research other potential uses for such a script...

### **9.1.3** Method

Development was akin to a more traditional methodology; requirements were simple and available up front, testing could wait until all the components had been completed to properly measure whether outcomes were as expected.

Goldilocks had to be constructed quickly but carefully. Although avoiding a quick and dirty solution, the initial codebase was arguably cluttered with blocks of repeated code to perform actions for different hard coded groups. Testing was not automated and utilised eyeballing of resulting data and occasional checking of input files to ensure results were as expected.

The first completed version consisted of a somewhat monolithic script, prime for breaking up in to a more organised set of functions...

#### 9.1.4 Pitfalls

...chromosome six must be ignored... ...chromosomes are 1-indexed!

## 9.2 Implementation

...we investigate the following implementation stories:

- Handling user input of grouped Variant Query files
- Loading variants from identified Variant Query files
- Storage of read data in memory (and troubles thereof)...
- · Striding and window size and data structures thereof
- Final output and selection, counters and buckets (avoid chr6)
- Testing, generate output files, call functions akin to API

## 9.2.1 Loading Variant Query File List

**Goldilocks** requires the user provide their own valid "pathsfile", meeting the specification demonstrated in Section 8.1.3 (an actual example of such a file can be found in Appendix A.3).

merely lists locations of input files and groups them together as desired ...all files in a particular group will be considered as a whole (all variants from all files in the group will be added to the set of chromosomes for that group)

**load\_variant\_files** is a simple function to read the pathsfile and initialise the relevant data structures for each group... ...each group will have a three data structures created...

## 9.2.2 Loading Variants from Input Files

**load\_variants\_from\_file** is responsible for loading the file referenced by a particular path and adding each valid variant record to the correct file group...

...this function also keeps track of the "furthest along" variant for each chromosome over all groups, this is used in the search step later to ensure each group has the same number of regions...

...no duplicate checking to avoid needing to check for list membership ...this is not an issue later (setting a flag to 1 multiple times will cause no effect)... seemed faster to repeat those actions than to check for membership...

...however this process of appending variants to a list would suffer from expensive memory moving operations and is ripe for alteration... ...could use a dictionary... ...but perhaps a set would be more appropriate... ...investigating the difference in performance would be of interest...

...although the input for this part of the study handles <x> variants in <y> seconds and thus time was better spent elsewhere...

### 9.2.3 Storage

As I discussed previously, on the hunt for my Goldilocks genomic region, it is important to consider both time and memory as it is simple to deliver a poor performing solution.

Searching for candidates regions over the entire genome at once is probably unwise. Luckily, since our candidate must not span chromosomes (telomeres – the ends of chromosomes are not very good for reads) then we can easily yield a great improvement from processing chromosomes individually.

The process is to extract the locations of all the variants across the genome from the SNP chip VCF files (these files list the alleles for each detected variant for each sample), load them in to some sort of structure, "stride" over each chromosome (with some stride offset) and finally list the variants present between the stride start and stride start plus the desired length of the candidate. These are our regions.

Due to the use of striding, one does not simply walk the chromosome. A quick and dirty solution would be to just look up variants in a list:

```
for chromosome in autosomes:
   for start in range(1, len(chromosome), STRIDE):
        for position in range(start, start+LENGTH):
```

```
if position in variant_position_list:
    # Do something...
```

This is of course rather dumb. Looking up a list has  $\mathcal{O}(n)$  performance where n is the number of variants (and there are a lot of variants). Combining this with the sheer number of lookups performed, with the default parameters the list would be queried half a billion times for the first chromosome alone.

You could improve this somewhat by dividing the variant\_position\_list in to a list of variants for each chromosome, so at least n is smaller. It is pretty doubtful that this would make any useful impact given the number of lookups. I'm happy to say I bypassed this option but thought it would be fun to consider its performance.

A far more sensible solution is to replace the list with a dictionary whose lookup is amortized to a constant time. The number of lookups is still incredibly substantial but a constant lookup is starting to make this sound viable. Using the variant positions as keys of a dictionary (in fact let us use the previous minor suggestion and have a dictionary for each chromosome) we have:

```
for chromosome in autosomes:
    for start in range(1, len(chromosome), STRIDE):
        for position in range(start, start+LENGTH):
        if position in variant_position_dict[chromosome]:
        # Do something...
```

This still takes half an hour to run on my laptop though, surely there is a better way. I wondered if whether dropping the lookup would improve things. Instead, how about an area of memory is allocated to house the current chromosome and act as a variant "mask" – essentially an array where each element is a base of the current chromosome and the value of 1 represents a variant at that position and a 0 represents no variant.

We of course have to initially load the variants in to the chromosome array, but this need only be done once (per chromosome) and is nothing compared to the billions of lookups of the previous implementation.

Rather to my surprise this was slow (maybe if I have time I should check how it compared to looking up with a list). Was it the allocation of memory? Perhaps I had run out of RAM and most of the work was going in to

fetching and storing data in swap?

I switched out the comparison (== 1) step for the previous dictionary based lookup and the performance improved considerably. What was going on? There must be more to looking at a given element in the numpy chromosome array, but what?

After a brief spell of playing with Python profilers and crashing my laptop by allocating considerably more memory than I had with numpy.zeroes, I read the manual (!) and discovered that numpy.zeroes returns an ndarray which uses "advanced indexing" even for single element access, effectively copying the element to a Python scalar for comparison.

That's a lot of memory action.

It then occurred to me that we're interested in just how many variants are inside each region and our chromosome is handily encapsulated in a numpy array. Why don't we just sum together the elements in each region? Remember variant positive base positions are 1 and 0 otherwise, useful. So the work boils down to some clever vector mathematics calculated by numpy.

We don't even lose any detail because the actual list of variants inside a region can be recovered in a small amount of time just given the start and end position of the region.

```
for chromosome in autosomes:
    chro = np.zeros(len(chromosome), np.int8)
    for variant_loc in snps_by_chromosome[chromosome]:
        chro[variant_loc] = 1

for start in range(1, len(chromosome), STRIDE):
    num_variants = np.sum(chro[start:start+LENGTH])
    # Do something...
```

With conservative testing this runs at least 60 times faster than before, the entirety of the human genome can be analysed for candidate regions in less than twenty seconds (with the first few seconds taken reading in all the variant locations in the first place).

...Ignoring empty regions etc.

```
def load_chromosome(self, size, locations):
    chro = np.zeros(size+1, np.int8)

# Populate the chromosome array with 1 for each position a variant exists
for variant_loc in locations:
    chro[variant_loc] = 1
return chro
```

**load\_chromosome** returns a **numpy** array representing the chromosome, given its desired length and a list of variants...

...supervisor suggested bloom filters... ...would be a useful experiment to measure what further performance gains could be acheived with such a method...

## 9.2.4 Searching for Regions

••

**search\_regions** ...possibly a misnomer as this is more of a census than a search... ...responsible for the core objective function as outlined in Section 9.1.1... ..each region of *length*, starting at *stride* is checked for the number of variants contained therein...

...it is here that chromosome six is discarded...

#### 9.2.5 Final Selection

#### initial filter enrich

...provides functions to rank all candidate regions by density of variants ...currently criteria are still hard coded to meet the needs of the project and propose little re-use value unless altered... ...for the purpose of this project we're looking at: ...the median of the GWAS (representative) ...the maximum of the iCHIP (enriching for maximum comparisons between data sets)

...with more time it would be useful to open these functions to input from an end user to provide their own criteria

...in this state does however still allow for repeatable results in future with different data sets or additional groups...

## 9.3 Usage

...

## 9.4 Testing

...particularly difficult. ...required generation of input files that would contain known regions to test were found by the program... ...needed to re-write large aspects of the scripts to allow unit-like testing to take place... ...wanted to avoid a monolithic test class...

...tests are still rather modest and with more time it would be appropriate to improve their function but I have faith in their results...

# **Results**

| WND  | GWAS | iCHIP | CHR | POSITIONS             |
|------|------|-------|-----|-----------------------|
| 234  | 297  | 470   | 1   | 117000001 - 118000000 |
| 1074 | 294  | 1540  | 3   | 46000001 - 4700000    |
| 3125 | 298  | 310   | 10  | 60000001 - 6100000    |
| 5222 | 294  | 336   | 21  | 16500001 - 1750000    |
| 880  | 293  | 344   | 2   | 191500001 - 192500000 |
| 3560 | 299  | 772   | 12  | 9000001 - 1000000     |
| 4407 | 299  | 512   | 15  | 78500001 - 7950000    |
| 1036 | 292  | 300   | 3   | 27000001 - 28000000   |
| 2734 | 300  | 515   | 9   | 5000001 - 6000000     |
| 3426 | 300  | 486   | 11  | 76000001 - 7700000    |
| 15   | 291  | 1029  | 1   | 7500001 - 8500000     |
| 365  | 301  | 487   | 1   | 182500001 - 183500000 |
| 3415 | 301  | 419   | 11  | 70500001 - 7150000    |
| 1581 | 290  | 802   | 4   | 102500001 - 103500000 |
| 3184 | 302  | 449   | 10  | 89500001 - 9050000    |
| 3554 | 290  | 403   | 12  | 6000001 - 700000      |
| 1580 | 289  | 603   | 4   | 102000001 - 103000000 |
| 414  | 288  | 346   | 1   | 207000001 - 208000000 |
| 1948 | 288  | 1297  | 5   | 96000001 - 9700000    |
| 2055 | 304  | 1377  | 5   | 149500001 - 15050000  |
| 2215 | 288  | 622   | 7   | 49500001 - 50500000   |
| 384  | 287  | 827   | 1   | 192000001 - 193000000 |
| 959  | 306  | 406   | 2   | 231000001 - 232000000 |
| 4214 | 286  | 393   | 14  | 88500001 - 8950000    |
| 320  | 307  | 620   | 1   | 160000001 - 161000000 |

Listing 5 : ...top 25...

Figure 10.1 was created quickly in R using **ggplot2** to eyeball the difference in variant distribution across each region (called a 'megabase' due to their length of one million bases).. ...motivated by locating very highly populated regions in the iCHIP group well above the mean in the output...

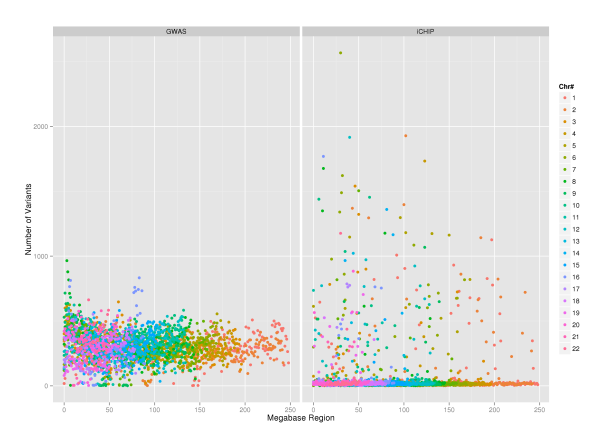


Fig. 10.1 **Megabase Plot**: ...

# **Analysis Pipeline**

Following the extraction of the Goldilocks region for each of the samples... we need to begin putting together the pipeline that would process the work for the leave-one-out analysis...

...involving generation of indexes for each sample ...execute the pipeline and withold one of the Goldilocks samples... ...using **samtools** to pileup the variants across all the samples (excluding one of course) and then performing variant calling... ...must then compare the resulting VCF against the SNP VCF and calculate the concordance to discover whether leaving out that particular sample has caused a change in the accuracy of the variants called...

## 11.1 Implementation

## 11.1.1 Region Extraction

...regions extracted from IRODS ...difficulty gaining user access for both IRODS and "the farm"

#### 11.1.2 samtools index

...index extracted regions

### 11.1.3 samtools mpileup

Following extraction of the Goldilocks region for each full-genome sample, the next stage is to use **samtools mpileup** to ... ...each of the various full genome samples we have for processing. These files are then summarised (and likelihoods are calculated) with samtools mpileup...

**samtools mpileup** is a rather intensive process especially when in a scenario such as ours where there are thousands of input files, one for each of the extracted regions. This not only places considerable strain on the cluster's file system but is incredibly inefficient given the overhead of file handling. As this pipeline will need to be executed once for each leave-one-out experiment, it would be required to reduce the strain on the file server before the job could be submitted. Usefully another member of the **samtools** collection provides functionality to merge a list of given input files such as those extracted Goldilocks regions.

## 11.1.4 samtools merge

### Usage

...undocumented feature to use a file of filenames... allowing more than two files to be specified at a time...

### **Memory Leak**

By default the LSF scheduler at the Sanger Institute will issue 100MB to any submitted job. Given both the number of input files and their total size it was anticipated that this merge job would require more memory. At the very least an estimated 35kB for a 64-bit pointer to each input file and approximately 150MB to house a 32kB buffer to read data from each file also. Yet executing the job with 500MB caused the LSF scheduler to forceably terminate the process for exceeding the maximum allocated memory limit.

Assuming that the intermediate structures for storing and sorting the input data must have been greater than expected, the job's memory limit was generously increased with the syntax demonstrated by Listing 6 only to meet the same fate, even when reserving 16GB of memory...

This is an absurd amount of memory even considering the vast input... It appeared a memory leak had been discovered... that drastically increases in severity as more files are provided as input,

Listing 6: Flags required to raise job memory allocation

...initial memory leak fixed by the author, several large variables not being freed from memory... ...following this, further **samtools merge** jobs were submitted only to also be repeatedly terminated by the LSF scheduler, this time for exceeding the maximum execution time limit for the queue...

#### **Memory Leak in Test Harnesses**

```
...getline ...regcomp...
```

...other memory leaks still remained in the tests and wanting to brush up on finding memory leaks with **valgrind**, I volunteered to locate and patch these...

```
==30464==416 bytes in 1 blocks are indirectly lost in loss record 85 of 103
==30464== at 0x4A082F7: realloc (in /usr/lib64/valgrind/vgpreload_memcheck-amd64-linux.so)
==30464== by 0x3FBCCCA725: duplicate_node (in /usr/lib64/libc-2.17.so)
==30464== by 0x3FBCCD3ADA: duplicate_node_closure (in /usr/lib64/libc-2.17.so)
==30464== by 0x3FBCCD415A: calc_eclosure_iter (in /usr/lib64/libc-2.17.so)
==30464== by 0x3FBCCD79F6: re_compile_internal (in /usr/lib64/libc-2.17.so)
            at 0x4A08121: calloc (in /usr/lib64/valgrind/vgpreload_memcheck-amd64-linux.so)
==30464==
==30464==
            by 0x3FBCCCCD88: create_cd_newstate (in /usr/lib64/libc-2.17.so)
==30464==
            by 0x3FBCCCD506: re_acquire_state_context.constprop.41 (in /usr/lib64/libc-2.17.so)
==30464==
            by 0x3FBCC2132E: build_trtable (in /usr/lib64/libc-2.17.so)
==30464==
           by 0x3FBCCD3792: re_search_internal (in /usr/lib64/libc-2.17.so)
==30464== by 0x3FBCCD8E94: regexec@GLIBC_2.3.4 (in /usr/lib64/libc-2.17.so)
==30464==
           by 0x40C6FA: check_test_2 (test_trans_tbl_init.c:124)
            by 0x402CC4: main (test_trans_tbl_init.c:348)
==30464==
```

Listing 7: Example of **valgrind** locating a memory leak in one of the **samtools** test harnesses following the failure to release memory allocated to a compiled regular expression

#### **Poor Time Performance**

...submitting the same job to the "long" (48hr) and "basement" (essentially unlimited) queues, it is clear that the job is taking an extraordinary length of time to complete...

...during this time I took the opportunity to patch various memory leaks in the test harnesses of both merge and split...

valgrind, the tool I used to track down memory leaks in the test harnesses of both samtools merge and samtools split actually consists of more than just memcheck.

callgrind is a profiling tool that keeps track of a program's call stack history, a handy feature built in to some development environments such as QtCreator.

Having constructed a modest test set of files to merge, I called samtools merge from the command line, attaching callgrind and later imported the resulting text file to QtCreator's profiling tool to interpret the results (I usually use KCacheGrind for this but I've been investigating QtCreator's feature set for reasons unrelated

to this project), the result is immediately obvious - millions of calls to functions in zlib; a free compression library.

Further investigations using the QtCreator Analyze interface revealed that these calls all boiled down to one line called not during the process of deflating the input files (as I had expected) but actually during the compression of the output!

Fig. 11.1 callgrind output following merge with default output compression

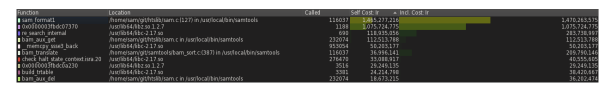


Fig. 11.2 callgrind output following merge with uncompressed output

Looking at a brief explanation of the deflate algorithm, it seems reasonable to conclude the computational cost is rather asymmetric between compressing and uncompressing - in that the effort is locating blocks to compress and in comparison uncompressing is a reversible function on the known blocks.

Indeed, samtools merge specifies a -u option for uncompressed output and the callgrind output (second image) indicates significantly less calls to zlib functionality.

It remains to be seen whether this option will cut down the time needed for the large merge job, perhaps this is merely a red herring and we're yet to discover the true speed trouble. In the meantime let's see if sending this job to the farm will work.

#### The Red Herring

... might be interesting to use gprof which is more geared towards finding functions that spend all your execution time as opposed to callgrind which I believe counts CPU instructions.

After re-compiling htslib and samtools with the -pg flag to enable such profiling and executing the same previous merge command on the modest test set, the output as parsed by gprof seems to indicate that the trouble lies with bam\_aux\_get in htslib, with almost 50% of the execution time being spent in this particular function.

```
uint8_t *bam_aux_get(const bam1_t *b, const char tag[2])
{
    uint8_t *s;
    int y = tag[0]<<8 | tag[1];</pre>
```

```
s = bam_get_aux(b);
while (s < b->data + b->l_data) {
   int x = (int)s[0]<<8 | s[1];
   s += 2;
   if (x == y) return s;
   s = skip_aux(s);
}
return 0;
}</pre>
```

At a glance it seems that bam\_aux\_get receives a pointer to a BAM record and a "tag", an array of two characters representing an optional field as defined in Section 1.5 of the SAM file spec.

The function then appears to fetch all these auxiliary tags and iterates over each, comparing a transformation of that tag(x) to a pre-computed transformation on the input tag(y).

This would of course be inherently slow for files with many such tags; especially given that the function is called twice for potentially each line in a BAM file.

#### The Plot Thickens

...as we increase the number of input files, the time taken to read them in becomes non-linearly slower. Currently my money is on the seemingly inefficient trans\_tbl\_init that appears to be called for each file, with the current table of all previous files as an input...

#### 11.1.5 beftools call

...Unfortunately during the initial testing run of this step with all the Goldilocks regions it was discovered that the output only included the standard header information and not a single line for the variants themselves.

...is because the piled up file was not generated with a corresponding reference DNA sequence and so the REF (reference) column is set to N (an ambiguity code which translates to 'any base'). bcftools call does have an -M flag to prevent ignoring rows where the REF base is N (apparently called a "masked reference") however this is currently causing a segmentation fault. Having recompiled htslib, samtools and bcftools I am now able to run bcftools call on my local machine on some test data. I guess I'll need to have someone recompile the source for me on the cluster I'm using....

# Part III

# **Discussions and Conclusions**

# **Current Status**

# **Critical Evaluation**

...Section 3.2.3 The **SciPy Stack** also includes the "Python Data Analysis Library" (**pandas**), which is designed to provide tools to supplement the Python environment, allowing users to perform analysis in Python instead of having to switch to a more analysis focused language such as R.

...at first glance appears to be a far more developed Frontier... ...however it should be considered that Frontier was designed to complement the machine learning experiments and provides

...whilst also providing the **AbstractReader** framework to allow users to quickly define methods to read in their own data regardless of how esoteric or cryptic the file format is...

# **Conclusions**

# References

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# Appendix A

# **Input Examples**

## A.1 BAMcheckR'd Example Output

```
\# Summary Numbers. Use 'grep ^SN \mid cut -f 2-' to extract this part.
         raw total sequences:
SN
         filtered sequences:
         sequences: 41400090
                        1
SN
         is paired:
         1 1st fragments:
SN
SN
                              20700045
         last fragments:
SN
                               20700045
SN
         reads mapped:
                           41291484
                           108606
60000
SN
         reads unmapped:
\mathtt{SN}
         reads unpaired:
         reads paired:
                             41231484
SN
         reads duplicated: 5756822
         reads MQ0: 1038644
SN
SN
         reads QC failed: 0
SN
         non-primary alignments:
         total length: 3105006750
SN
         bases mapped:
                             3096861300
         bases mapped (cigar): 3090885143
SN
         bases trimmed: 0
bases duplicated: 431761650
SN
SN
         mismatches: 9107833
error rate: 0.002946
                          0.002946675
SN
\mathtt{SN}
         average length:
SN
         maximum length:
                               75
         average quality:
SN
                               36
SN
         insert size average: 178.7
SN
         insert size standard deviation:
                                               44.1
         inward oriented pairs: 20577242 outward oriented pairs: 3140
SN
SN
         pairs with other orientation: 3711
pairs on different chromosomes: 31535
SN
         fwd percent insertions above baseline: 1.43135383851191
SN
         fwd percent insertions below baseline:
                                                     0.686265539012562
         fwd percent deletions above baseline: 0.38326380878871
         fwd percent deletions below baseline:
                                                   0.44923551909251
SN
                                                   1.08264446659241
SN
         rev percent insertions above baseline:
         rev percent insertions below baseline:
                                                      0.457290262062496
```

```
SN
          rev percent deletions above baseline:
                                                        1.15931214598243
SN
          rev percent deletions below baseline:
                                                        0.413119424753248
SN
          contiguous cycle dropoff count:
SN
          fwd.percent.insertions.above.baseline:
                                                         1.43135383851191
SN
                                                         0.686265539012562
          fwd.percent.insertions.below.baseline:
SN
                                                        1.38326380878871
          fwd.percent.deletions.above.baseline:
SN
          fwd.percent.deletions.below.baseline:
                                                        0.44923551909251
SN
          rev.percent.insertions.above.baseline:
                                                         1.08264446659241
SN
          rev.percent.insertions.below.baseline:
                                                         0 457290262062496
SN
          rev.percent.deletions.above.baseline:
                                                        1.15931214598243
SN
          rev.percent.deletions.below.baseline:
                                                        0.413119424753248
SN
          quality.dropoff.fwd.high.iqr.start.read.cycle:
                                                                 0
SN
          quality.dropoff.fwd.high.iqr.end.read.cycle:
SN
          quality.dropoff.fwd.high.iqr.max.contiguous.read.cycles:
                                                                           0
                                                                            20
SN
          quality.dropoff.fwd.mean.runmed.decline.start.read.cycle:
SN
          quality.dropoff.fwd.mean.runmed.decline.end.read.cycle:
                                                                           51
SN
          quality.dropoff.fwd.mean.runmed.decline.max.contiguous.read.cycles:
                                                                      36.9775883578997
SN
          quality.dropoff.fwd.mean.runmed.decline.high.value:
                                                                     36.301749247405
SN
          quality.dropoff.fwd.mean.runmed.decline.low.value:
SN
          quality.dropoff.rev.high.iqr.start.read.cycle:
SN
          quality.dropoff.rev.high.iqr.end.read.cycle:
SN
          quality.dropoff.rev.high.iqr.max.contiguous.read.cycles:
                                                                           0
SN
          quality.dropoff.rev.mean.runmed.decline.start.read.cycle:
                                                                            18
SN
          quality.dropoff.rev.mean.runmed.decline.end.read.cycle:
                                                                           56
SN
          quality.dropoff.rev.mean.runmed.decline.max.contiguous.read.cvcles:
                                                                                       39
SN
          quality.dropoff.rev.mean.runmed.decline.high.value:
                                                                      36.1517621338504
SN
          quality.dropoff.rev.mean.runmed.decline.low.value:
                                                                     35.3152133727245
SN
          quality.dropoff.high.iqr.threshold:
SN
          quality.dropoff.runmed.k:
SN
          quality.dropoff.ignore.edge.cycles:
                                                      3
SN
          A.percent.mean.above.baseline:
                                                 0.0991164444444441
SN
          C.percent.mean.above.baseline:
                                                 0.12737955555556
SN
          G.percent.mean.above.baseline:
                                                 0.0603679999999997
SN
                                                 0.086800000000005
          T.percent.mean.above.baseline:
SN
          A.percent.mean.below.baseline:
                                                 0.099116444444451
SN
          C.percent.mean.below.baseline:
                                                 0.127379555555555
SN
          G.percent.mean.below.baseline:
                                                 0.0603680000000002
SN
          T.percent.mean.below.baseline:
                                                 0.086799999999993
SN
          A.percent.max.above.baseline:
                                                0.6017333333333332
SN
          C.percent.max.above.baseline:
                                                0.394266666666667
SN
          G.percent.max.above.baseline:
                                                0.2956
SM
          T.percent.max.above.baseline:
                                                0.768000000000001
SN
                                                0.31826666666666
          A.percent.max.below.baseline:
SN
          C.percent.max.below.baseline:
                                                0.8257333333333333
SN
          G.percent.max.below.baseline:
                                                0.554400000000001
SN
          T.percent.max.below.baseline:
                                                0.251999999999999
SN
          A.percent.max.baseline.deviation:
                                                    0.6017333333333332
SN
          C.percent.max.baseline.deviation:
                                                    0.8257333333333332
SN
          G.percent.max.baseline.deviation:
                                                    0.554400000000001
SN
          T.percent.max.baseline.deviation:
                                                    0.768000000000001
SN
          A.percent.total.mean.baseline.deviation:
                                                           0.198232888888889
SN
                                                           0.254759111111111
          C.percent.total.mean.baseline.deviation:
GM
          G.percent.total.mean.baseline.deviation:
                                                           0.120736
SN
          T.percent.total.mean.baseline.deviation:
                                                           0.1736
# First Fragment Qualitites. Use 'grep ^FFQ | cut -f 2-' to extract this part.
# Columns correspond to qualities and rows to cycles. First column is the cycle number.
FFQ
                    8968
                                3619
                                             9863
                                                         747
                                                                    5094
                                                                                0
                                                                                          6642
                                                                                                      1609
                                                                                                                   4673
FFO
           2
                    21676
                                 0
                                           0
                                                             0
                                                                      0
                                                                               0
                                                                                         0
                                                                                                  43
                                                                                                            1885
FFO
[...]
FFQ
           74
                     3697
                                 39
                                           0
                                                     919
                                                                4933
                                                                            0
                                                                                      0
                                                                                               56866
                                                                                                            1524
FFQ
           75
                     4542
                                           0
                                                    0
                                                             0
                                                                      0
                                                                                4634
                                                                                            77822
                                                                                                         0
                                  0
FFQ
           76
                                                                                      0
                                                                                               0
```

```
# Last Fragment Qualitites. Use 'grep ^LFQ | cut -f 2-' to extract this part.
\mbox{\tt\#} Columns correspond to qualities and rows to cycles. First column is the cycle number.
LFQ
          1
                   8869
                               0
                                        0
                                                 0
                                                          0
                                                                  0
                                                                           63 0
                                                                                              0
                                                                                                       1156
LFQ
           2
                    3300
                               0
                                        0
                                                 0
                                                          0
                                                                   0
                                                                            0
                                                                                     0
                                                                                              0
                                                                                                       0
                    6816
                                                          573
                                                                                                 7011
LFO
                               0
                                        0
                                                 0
                                                                             83
           3
[...]
LFQ
           74
                    5980
                                3
                                         91
                                                  0
                                                            0
                                                                     0
                                                                              1340
                                                                                          9696
                                                                                                     72939
LFQ
           75
                    4314
                                                  168
                                                                                 8591
                                                                                            0
                                                                                                     70358
LFO
           76
                    0
                             0
                                      0
                                              0
                                                       0
                                                                0
                                                                     0
                                                                                  0
                                                                                            0
# Mismatches per cycle and quality. Use 'grep ^MPC | cut -f 2-' to extract this part.
# Columns correspond to qualities, rows to cycles. First column is the cycle number, second
\mbox{\tt\#} is the number of N's and the rest is the number of mismatches
                   14078
                               0
                                        2594
                                                     6777
                                                                 416
                                                                            1919
                                                                                        0
                                                                                                 2222
MPC
                   21407
                                                  0
                                                           0
                                                                                      0
           2
                                0
                                         0
                                                                   0
                                                                             0
MPC
           3
                    3205
                               0
                                        0
                                                 37
                                                           0
                                                                    43
                                                                              0
                                                                                       12
Γ...1
                                                          131
MPC
           74
                     779
                               0
                                        3
                                                 0
                                                                                         93
                                                                                                   7485
                                                                                                              . . .
                                                                           47
MPC
          75
                    136
                               0
                                        0
                                                 0
                                                          3
                                                                                     704
                                                                                                 9302
           76
                                      0
                                               0
                                                        0
                                                                          0
\# GC Content of first fragments. Use 'grep ^GCF \mid cut -f 2-' to extract this part.
GCF
          0.5
                     56
GCF
           1.76
                      60
GCF
          3.02
                      126
GCF
           4.27
                      212
GCF
           5.78
                      347
[...]
GCF
           93.72
                       378
           95.23
GCF
           96.48
                       87
GCF
GCF
           97.74
                       55
GCF
           99.25
                       17
# GC Content of last fragments. Use 'grep `GCL | cut -f 2-' to extract this part.
GCL
          0.5
                     118
GCL
          1.76
                      175
GCL
           3.02
                      230
GCL
           4.27
                      354
GCL
           5.78
[...]
GCL
           93.72
                       613
GCL.
           95.23
                       430
GCL
           96.48
                       274
GCI.
           97.74
                       185
GCL
           99.25
                       110
# ACGT content per cycle. Use 'grep 'GCC | cut -f 2-' to extract this part. The columns are: cycle, and A,C,G,T counts [%]
                                                          27.2
GCC
                   26.93
                                23.09
                                             22.77
          1
GCC
           2
                    26.78
                                23.24
                                             22.97
                                                          27.02
GCC
           3
                   26.46
                                23.59
                                             23.3
                                                         26.66
GCC
           4
                    26.29
                                23.79
                                             23.45
                                                          26.46
GCC
           5
                   26.47
                                23.61
                                             23.3
                                                         26.62
[...]
GCC
           70
                    26.09
                                 24.26
                                              23.45
                                                           26.2
GCC
           71
                    26.07
                                 24.25
                                              23.46
                                                           26.22
GCC
           72
                    26.04
                                 24.27
                                              23.49
                                                           26.2
GCC
           73
                    26.07
                                 24.25
                                              23,47
                                                           26.22
GCC
           74
                     26.08
                                 24.24
                                              23.45
                                                           26.23
GCC
           75
                    26.01
                                              23.51
                                 24.31
                                                           26.18
# Insert sizes. Use 'grep ^IS | cut -f 2-' to extract this part.
# The columns are: pairs total, inward oriented pairs, outward oriented pairs, other pairs
IS
         0
                  10
                            0
                                    1
                                              9
TS
         1
                  3
                           0
                                    3
                                             0
IS
         2
                  4
                           0
                                    4
                                             0
IS
         3
                  5
                           0
                                    5
                                             0
                                             0
IS
                                    2
```

```
IS
                                     3
[...]
IS
          110
                     33952
                                  33952
                                               0
                                                         0
IS
          111
                     38433
                                  38433
                                                0
                                                         0
          112
                     43373
                                  43370
IS
                                               0
                                                         3
IS
          113
                     48160
                                  48159
IS
          114
                     53175
                                  53171
                                               0
IS
          115
                     59504
                                  59502
                                               0
IS
          116
                     64668
                                  64668
                                               0
                                                         0
IS
          117
                     71107
                                  71105
IS
          118
                     77157
                                  77156
                                               0
IS
          119
                     84044
                                  84044
                                               0
                                                         0
IS
          120
                     90116
                                  90110
                                               3
                                                         3
[...]
IS
          327
                     6546
                                 6546
                                             0
                                                       0
IS
          328
                     6483
                                 6483
                                             0
                                                       0
IS
          329
                     6201
                                 6201
                                             0
                                                       0
                     6228
TS
          330
                                 6228
                                             0
                                                       0
                     5852
                                 5852
# Read lengths. Use 'grep ^RL | cut -f 2-' to extract this part. The columns are: read length, count
RL
          75
                   41400090
# Indel distribution. Use 'grep ^ID | cut -f 2-' to extract this part.
# The columns are: length, number of insertions, number of deletions
ID
                   128650
                                 183418
                   26409
                                39770
ID
          2
ID
          3
                   10213
                                16046
ID
          4
                   7756
                               11444
                   1746
                               3455
ID
[...]
ID
          35
                    0
                             8
ID
          36
                    0
                             1
ID
          37
                    0
                             1
ID
          38
                    0
                             1
ID
                    0
          40
                             2
\mbox{\tt\#} Indels per cycle. Use 'grep ^IC \mid cut -f 2-' to extract this part.
\# The columns are: cycle, number of insertions (fwd), .. (rev) , number of deletions (fwd), .. (rev)
                   0
                                                  179
IC
          2
                   24
                             15
                                       150
IC
          3
                   129
                              138
                                         441
IC
          4
                   253
                              310
                                         623
                                                     829
IC
                   557
                              724
                                         786
                                                    1164
          5
[...]
IC
          70
                    571
                               710
                                          638
                                                     761
IC
          71
                    350
                               428
                                           309
                                                      434
IC
          72
                    154
                               150
                                          38
                                                     45
IC
          73
                    60
                              61
                                        15
                                                   23
                    20
          74
                              19
                                        11
IC
                                                  12
# Coverage distribution. Use
                             'grep ^COV | cut -f 2-' to extract this part.
COV
           [1-1]
                       1
                                 332980694
                                 105004580
COV
           [2-2]
COV
           [3-3]
                        3
                                 29112182
COV
           [4-4]
                        4
                                 13415014
COV
           [5-5]
                        5
                                 6716815
[...]
COV
           [996-996]
                            996
                                       2
COV
           Г997-9971
                            997
                                       2
COV
           [998-998]
                            998
COV
           [1000-1000]
                             1000
           [1000<]
                          1000
                                      116
\mbox{\tt\# GC-depth}. Use 'grep ^GCD \mid cut -f 2-' to extract this part.
# The columns are: GC%, unique sequence percentiles, 10th, 25th, 50th, 75th and 90th depth percentile
GCD
           0
                    0.001
                                 0 0 0
                                                            0
GCD
                      0.002
                                   0.101
                                                              0.101
           0.4
                                                 0.101
                                                                           0.101
                                                                                        0.101
```

| GCD | 19 | 0.003  | 0.049 | 0.049 | 0.049 | 0.049  | 0.049  |
|-----|----|--------|-------|-------|-------|--------|--------|
| GCD | 20 | 0.004  | 0.06  | 0.06  | 0.06  | 0.06   | 0.06   |
| GCD | 21 | 0.004  | 0.045 | 0.045 | 0.045 | 0.045  | 0.045  |
| []  |    |        |       |       |       |        |        |
| GCD | 66 | 99.99  | 0.244 | 2.693 | 6.746 | 11.794 | 15.885 |
| GCD | 67 | 99.994 | 1.279 | 1.279 | 4.305 | 9.667  | 11.483 |
| GCD | 68 | 99.997 | 4.148 | 4.148 | 4.463 | 5.741  | 7.354  |
| GCD | 69 | 99.999 | 0.499 | 0.499 | 0.499 | 1.935  | 1.935  |
| GCD | 72 | 100    | 0.476 | 0.476 | 0.476 | 1.219  | 1.219  |

## A.2 auto\_qc Decision Matrix

| lanelet  | sample    | study | npg     | aqc  |         |  |
|----------|-----------|-------|---------|------|---------|--|
| 9999_9#1 | AQCTest00 | 0000  | AQCTest | pass | passed  |  |
| 9999_9#3 | AQCTest00 | 0002  | AQCTest | fail | failed  |  |
| 9999_9#2 | AQCTest00 | 0001  | AQCTest | pass | pass    |  |
| 9999_9#5 | AQCTest00 | 0004  | AQCTest | warn | warn    |  |
| 9999_9#4 | AQCTest00 | 0003  | AQCTest | warn | warning |  |
| 9999_9#7 | AQCTest00 | 0006  | AQCTest | fail | fail    |  |
| 9999_9#6 | AQCTest00 | 0005  | AQCTest | pass | passed  |  |
| 9999_9#9 | AQCTest00 | 8000  | AQCTest | warn | warning |  |
| 9999_9#8 | AQCTest00 | 0007  | AQCTest | pass | passed  |  |

## A.3 Goldilocks Pathsfile

| # gwas   |   |
|----------|---|
| cd-gwas  | /pools/encrypted/sanger/vcf/cd-seq.vcf.gz.q   |
| uc-gwas  | /pools/encrypted/sanger/vcf/uc-seq.vcf.gz.q   |
| # ichip  |   |
| cd-ichip | /pools/encrypted/sanger/vcf/cd.ichip.vcf.gz.q |
| uc-ichip | /pools/encrypted/sanger/vcf/uc.ichip.vcf.gz.q |