Integrated Deep Learning and Bayesian Classification for Prioritization of Functional Genes in Next-Generation Sequencing Data

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DECLARATION

I hereby declare that this thesis is my original work and it has been written by me in its entirety. I have duly acknowledged all the sources of information which have been used in the thesis.

This thesis has also not been submitted for any degree in any university previously.

Chan Khai Ern Edwin 04 April 2017

Acknowledgements

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Abstract

Calling of high confidence variants and ranking them systematically are key steps in enabling personal genomic pipelines for clinical use. In this paper, we describe the use of a deep learning neural network to accurately call variants, and we describe a Bayesian classification method for the ranking of important genes. We show that an optimised neural network can call variants more accurately than single variant callers or concordant callers, with F1 score improvements of 6.5 percent in simulated datasets and 4.5 percent in real datasets over the best concordant methods. We also show that a Bayesian classification system is able to discover critical genes in a Diffuse Large B-Cell Lymphoma patient sample. Ultimately, our work develops a potential analytical pipeline for use in clinics to augment diagnosis and treatment of diseases.

1 Introduction

1.1 Next Generation Sequencing in Personal Genomics Pipelines

Identification of functionally important mutations is a critical step in enabling personal genomic pipelines. Recently, there has been keen interest in using a person's genome to help doctors treat and diagnose disease (Rehm, 2017; Angrist, 2016). The fundamental intuition is that sequencing the individual's genome can help doctors and clinicians narrow down important disease subtypes and progression. For example, in cancer, finding out the key driver mutations can help doctors identify the disease subtype (Stratton, Campbell & Futreal, 2009) and thus provide targeted medication to treat that specific cancer subtype (Janitz, 2011). This is of particular interest now, especially since it is possible to sequence an exome for about \$1,000 - a price similar to most other clinical procedures. This process taps upon advances in Next-Generation Sequencing(NGS), where genomic sequencing is done in parallel to rapidly determine the whole-exome sequence (Metzker, 2010; Mardis, 2008).

However, there are still two critical steps that prevent this data from being used in a clinical setting. Firstly, it is difficult to obtain high-confidence mutations from this sequencing data, and secondly, there needs to be a systematic method to prioritise these mutations for

clinicians and doctors. For the first problem, we have to be able to find out mutations in the person's genome that are different from a reference genome. This step is termed in literature as variant calling as it involves the discovery (calling) of genes (variants) that differ from a reference genome. However, current variant callers are not reliable in accurately and precisely calling mutations (Cornish and Guda, 2015; O'Rawe et al., 2013). As can be seen from Figure 1, single variant callers can differ greatly in effectiveness depending on the type of sequencing machine used, which makes it difficult to be confident of their reliability.

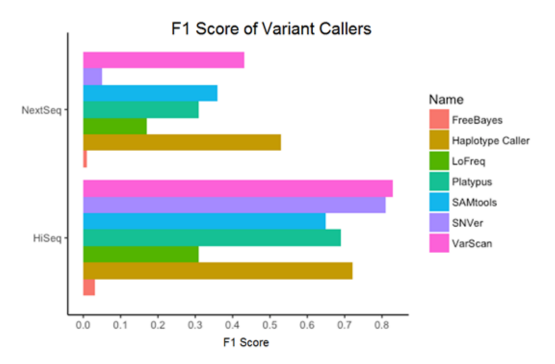


Figure 1: Validation of Variant Calling tools with real patient data on two different Illumina sequencing platforms (HiSeq and NextSeq) using prior known mutations. The F1 score indicates how well a caller can predict true positives (See Appendix 5.3 for more details), and it can be seen that the F1 score for the same variant calling tool can differ greatly. Figure adapted from Sandmann et al. (2017)

The second problem, ranking of mutations, is also important as the importance of each variant has to be determined somehow. With at least 3 million mutations in a typical genome (Shen et al., 2013), there needs to be a systematic way to find out the most important ones in a short amount of time. This is important as a clinician should be able to obtain variants that are of clinical significance without having to sieve through literature to manually pick out genes that important. This would allow them to narrow their search to the most likely candidate genes, and then be able to embark on an ideal treatment pathway.

In this paper, we describe a machine learning approach to solve these two problems to use deep learning to validate high-confidence variant calls and to use Bayesian networks to filter variants and prioritise their importance.

1.2 Variant Calling Methodology

Here, it is useful to analyse variant calling and the problems that make it difficult. Variant calling in NGS data primarily involves the use of various statistical and mathematical methods to discover variants in the genome (Nielson et al., 2011). These variants represent the deviations and differences between the genome of interest and a standard human genome. However, this is not a trivial problem as for every call involves the analysis of a complex amount of information (Zook et al., 2014). This comes from a large amount of information that is created from NGS data - because sequencing is done by mapping a large number of reads together, calling specific mutations becomes a difficult problem. For example, if we look a typical pileup of a variant call involves (Figure 2), we can visualise the complexities that go into a variant call.

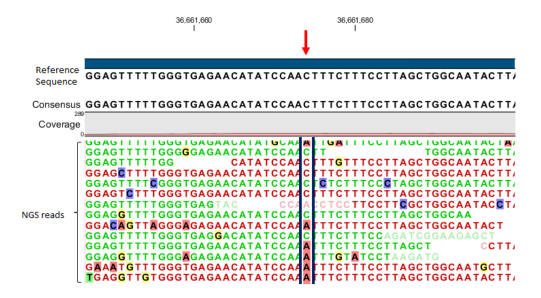


Figure 2: Variant Calling Pileup - Due to noise and errors in sequencing and variant mapping, it is sometimes difficult properly call variants. Figure adapted from CLC Genomics Workbench 9.5, Figure 29.8.

As can be seen from Figure 2, at the location of interest there are a number of reads with competing bases - is there a mutation from C to A? Some of these reads might have been wrongly mapped, while other reads might be sequenced wrongly. The difficulty for the variant caller then is to be able to look at these types of reads and form a decision on whether it should be called as a variant or not (Zook et al., 2014; Davey et al., 2011). This problem fundamentally results from assumptions and implementations of variant callers - certain algorithms are more sensitive and accurate in calling certain classes of mutations but suffer from inaccuracies in calling other variant types and edge cases (O'Rawe et al., 2013). Furthermore, because the variant callers do not differentiate between the specifics of each processing pipeline, such as the types of sequencing machines used or the alignment algorithm used, it is difficult for them to make right decisions in all types of variant calls. There are still areas for improvement in current variant calling methods, including dealing with different classes of mutations, as well as reducing the number of false positives (Mohiyuddin, et al., 2015; Gézsi et al., 2015). Thus, one large hurdle to overcome in enabling personal genomic pipelines is the generation of high-quality variant calls.

1.3 Deep Learning in Variant Calling

One useful way to approach this problem is to consider that each variant caller samples from the same genome but with a different technique. We can then see each variant caller provides us with a unique piece of information on the genome. Thus, we can generate more accurate calls by aggregating the multi-modal data from various callers, allowing us to cross-validate the variants called. The simplest approach to aggregate data is concordance - if multiple variant callers are able to call a variant, it is more likely to be accurate (Lam et al., 2012; Wei et al., 2011). However, the recall of such a tool would be poor due to the differential sensitivity of callers to edge cases, resulting in a lot of false negatives as true variant calls might only be picked up by one or two calling methodologies (O'Rawe et al., 2013). More sophisticated efforts have since been done to use machine learning methods such as Support Vector Machines as a way to integrate variant calling information (Gézsi et al., 2015), and the authors showed that SVMs presented an improvement over concordance based methods. However, with the advent of deep learning techniques and libraries, which

have been shown be able to integrate complex multi-modal information to solve problems (Ng et al., 2015), we hypothesise that deep learning can also be used to integrate the information from variant callers.

Deep learning is a method of machine learning that involves multiple stacks of artificial neural networks (LeCun et al., 2015). These neural networks were inspired by the way our synapses work in the brain and are represented in silico by input/output nodes that fire when a certain threshold is reached. Thus, these neural networks are able to simulate learning - by learning from labelled data correlations between inputs and outputs, these networks can predict outputs if given a new input. Deep learning has been shown to be able to solve other complex non-linear decision boundary problems, including drug molecule solubility (Lusci et al., 2013), facial recognition (Sun et al., 2014) and even predicting the best move in the Japanese board game, Go (Silver et al., 2016). Figure 3 shows a typical neural network with 5 layers, including an input and output later.

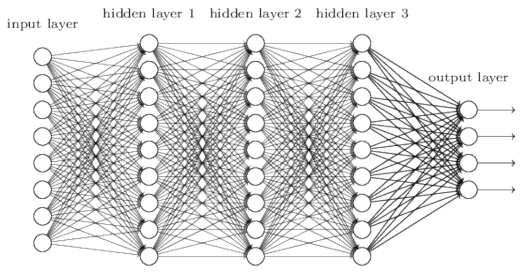


Figure 3: A Neural Network with one input layer, three hidden layers and one output layer. This represents a densely connected neural network, where each node is connected to every node of the preceding and subsequent layers.

Deep learning networks are trained by providing the input data and output data and letting the network learn how to integrate the inputs to create a network of activations that can be used to produce the corresponding output. A more in-depth explanation into their algorithms can be found in Appendix 5.1. In variant calling, we hypothesise that

deep learning will allow us to predict based on variant calling features and data whether a variant is valid and exists, or is erroneous. The deep learning network should be able to draw on the diversity of data with different variant callers and learn which patterns will result in a valid call and which patterns are false positives. It will also allow us to tap on the differential sensitivity of different callers, as the neural network is able to learn which callers work best for which types of mutations. Thus, we propose that deep learning as a combinatorial approach will allow us to improve the accuracy and precision of variant calling.

1.4 Gene Prioritisation and Bayesian Networks

The second problem of enabling personal genomic pipelines is gene prioritisation. The problem of gene prioritisation arises because there is a multitude of data sources we can draw on to analyse how important a gene is (Moreau & Tranchevent, 2012). The possible approaches include studying previously characterised variants and their phenotypic effects on a person as well as how the mutation itself will affect protein function through studying the likelihood of amino acid mutation for conserved regions. These functional annotations can be done with the tool ANNOVAR (Wang, Li, & Hakonarson, 2010) but the fundamental problem here is integrating such information in a systematic manner that is clear and understandable to clinicians. Clinicians may not be so familiar with the tools and functional annotation pipelines, and so for them to trust such a ranking system, they have to be able to understand intuitively how it works.

To solve this problem, we can use Bayesian networks to integrate the information from functional annotations as well as the confidence of a call (how likely it is real) to provide a ranking system for how likely the gene is going to be important. Bayesian networks were chosen for this ranking system as it is understandable and yet have proven stable in terms of solving decision-making problems (Pourret et al., 2008; Jensen et al., 1996). Bayesian networks have been applied in medical treatment decision making (Windecker et al., 2014), ecological studies (Johnson et al., 2014) and even predictive epidemiology (Su et al., 2014). A Bayesian network is a network that records the probabilities of events and based on conditional probabilities and observations it updates the final likelihood of an event. This can

be seen in Figure 2.

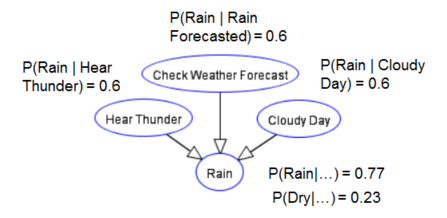


Figure 4: A Sample Bayesian Network for Rain Prediction.

Here, we would like to predict how likely it is to rain. Thus, we record observations (we hear thunder, check the weather forecast, notice it is a cloudy day) and updating the likelihood of rain happening based on the conditional probabilities of P(Rain — Hear Thunder)... and so on. This model of learning was chosen because a Bayesian Network closely mimics the way humans think - we observe events and form co-relational and causative predictions based on those events. This is advantageous over deep learning as in deep learning we are unable to interrogate the system to understand intuitively what the network is learning form - it has high predictive power but is essentially a black-box. The Bayesian network allows clinicians and doctors to see what are the components that went into gene ranking and prioritisation, and even change the probabilities, weights or add more observation nodes based on their own diagnosis and treatment knowledge. Ultimately, this enables the doctors and clinicians to be able to have confidence in the software as they can analyse and understand how it works. This also allows them to be able to explain their methodology and treatment plans clearly to the patient, making the diagnosis and treatment process clear, understandable and transparent.

1.5 Aims and Research Structure

In this paper, we describe the deep learning to validate high-confidence variant calls (focusing on SNVs and short indels) and using Bayesian networks to filter variants and prioritise their importance. Specifically, we will verify the use of deep learning to validate true variants in both real and simulated datasets. Subsequently, we will build a Bayesian network

based on functional annotations to prioritise mutations and test our network on a cancer dataset.

2 Materials and Methods

2.1 Overall Analysis Structure

To enable the deep learning networks and Bayesian network analysis, we first built two main computational pipelines - the first pipeline is used to train and optimise a neural network, and the second pipeline uses a trained neural network to perform variant validation and prediction. Two pipelines are required as one pipeline is needed to train the network, and then the second pipeline can use the trained network to perform predictions. The first pipeline involves using a training dataset (either simulated or obtained from real sequencing data) and then performing processing steps of alignment, variant calling and finally deep learning network training and prediction on the dataset. This pipeline focuses on generate a set of predictions, and subsequently validating this set of predictions with a ground truth. The second analysis pipeline is meant for real datasets with in which the ground truth is not known, and so it uses a pre-trained neural network. For this pipeline, we perform similar steps of alignment and variant calling, and then use a trained neural network to predict which features are true. Finally, we can apply the Bayesian network analysis to predict the important genes in this dataset.

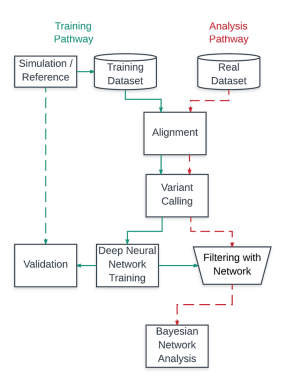


Figure 5: Overall Analytical Pipelines - Pipelines were implemented using the Groovy Domain Specific Language, NextFlow.

2.2 Programming and Pipelining tools

The general programming language used was Python (v2.7). Python was chosen due to its access to various important libraries, including NumPy, SciPy, Pomegranate and PyVCF. NumPy was used to prepare input vectors for deep learning training, SciPy was used to perform Principal Component Analysis and Synthetic Minority Oversampling Technique Methods (See Appendix 5.3 for more information). Finally, PyVCF was used to parse the VCF files into python objects for easy manipulation. Comparison of variants was also performed using python dictionary lookups. This method was chosen due to a high number of lookups required, and a fast O(1) constant time required for each lookup.

General pipelining and chaining of programmes was done using NextFlow and Bash scripts. NextFlow is a Groovy Based Domain Specific Language (DSL) that provides easy writing of parallel pipelines with an accessible unix interface. Nextflow was used to run the overall pipelines and control input and output of abstracted core modules, which are in turn either python scripts or Bash shell scripts. This ensures that results are easily replicable and can

be later implemented as a single analytic pipeline for clinical use.

For our deep learning networks, we used the Keras library (v1.1.1) with a TensorFlow backend (v0.11.0). TensorFlow (Abadi et al., 2015) was chosen due to its distributed computation and queue management system that enabled better performance in training on a CentOS-7 compute cluster compared to other backends. For more explanation on the algorithms underpinning deep learning, see Appendix 5.1 for more information.

Finally, ANNOVAR was used to generate the functional annotations for the Bayesian probabilistic model. The protocols used were snp138, clinvar_20150629 and ljb26_all. Pomegranate (a Python library) was used to generate and compute the probabilistic model and ranking system for our Bayesian Network. For the probabilistic model, The preparation of the probabilistic model and the truth-tables required was done using Python scripts and subsequently used as input in pomegranate.

2.3 Artificial Datasets

Artificial genomes enable the simulation of NGS data with ground truths to test and validate a neural network. For our simulator, we used Mason, a genome mutation software written in C++ (SeqAn version 2.3.1) to mutate the hg19 reference genome from UCSC (Karolchik et al., 2014). We used indel rates of 0.00002 and SNP rates of 0.00008 to generate sufficient truth variants for analysis, which comprise 229253 SNPs and 57257 indels.

After generating a ground truth model, we simulated sequence reads with error rates and ground truth variants (Figure 5). For error rates, we used published data from Schirmer et al. (2016) as the input to Mason - the general substitution error rate used was 0.0004 per base in the genome, and the insertion and deletion error rate per base were $5 * 10^{-6}$.

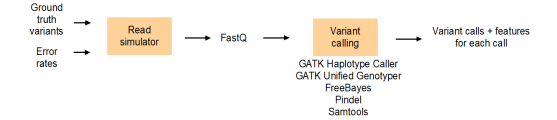


Figure 6: Pipeline for simulation of artificial genome for analysis

2.4 Alignment and Variant Calling

To perform alignment of simulated and real sequences, the Burrows-Wheeler Aligner (Li, 2013), version 0.7.13, was used. Default settings were used, with the mem option which is known to work well with longer sequences. After alignment, variant calling was performed. Variant callers used were FreeBayes (v1.0.2-16), GATK Haplotype Caller (v3.7-0) and Unified Genotyper (v3.7-0), Samtools (v1.3.1) and finally Pindel (v2.3.0)(Garrison & Marth, 2012; McKenna et al. 2010, DePristo et al. 2011; Li H, et al., 2009; Ye et al., 2009). All callers were used at their default settings.

2.5 Feature Engineering

In order to train a neural network, features in the form of numerical vectors must be used an input. We subset our features into three broad sets, which are base-specific information, sequencing error and bias information features, and calling and mapping quality. Here we describe the computation of the features - please see Appendix 5.2 for a more in-depth explanation on their usage and interpretation.

Base Information

Shannon Entropy

Shannon Entropy captures the amount of information contained inside the allele sequences. It is calculated using the equation:

$$H(X) = -\sum_{i=1}^{n} P(x_i) \log_2 P(x_i)$$
 (1)

where $P(x_i)$ is the prior probability of finding each base at each position. This prior probability is calculated in two ways - over the entire genome and over a region of space around the allele (10 bases plus the length of the allele in our calculations).

Kullback Leibler Divergence

The Kullback-Leibler Divergence feature is similar to Shannon entropy, but instead, we use this to measure the informational gain from the reference to the allele sequence. The Kullback-Leibler Divergence is calculated as follows:

$$D_{KL}(P||Q) = -\sum_{i=1}^{n} P(x_i) \log_2 \frac{P(x_i)}{Q(x_i)}$$
 (2)

where $Q(x_i)$ is the prior probability of finding each base at each position based on the genomic region around the allele, while $P(X_i)$ is the posterior probability of finding a specific base inside the allelic sequence.

Base Quality

Base quality refers to the Phred score probability that the called allele is wrong. It is given by the equation:

$$P = 10^{\frac{-Q}{10}}$$

Where P is the Base Quality, and Q is the probability that the allele called is wrong. This is a number computed by the sequencing machine based on the quality of the base samples provided.

Sequencing Biases and Errors

GC content

This feature comprises the percentage GC content of reference genome for at least ten bases around the mutation site.

Longest homozygous run

This feature comprises the longest similar string of bases in the reference genome, for at least ten bases around the mutation site.

Allele Count and Allele Balance

This feature is an output from Haplotype Caller and Unified Genotyper, and describes the

total number of alleles contributing to a call and the balance between reference and alternate alleles reads.

Calling and Mapping Qualities

Genotype Likelihood

The genotype likelihood score provides the Phred-scaled likelihood scores of how confident the caller is in determining that it is a homozygous or heterozygous call, and is provided by all variant callers.

Read Depth

Mapped read depth refers to the total number of bases sequenced and aligned at a given reference base position. It is provided by all variant callers.

Quality by Depth

Quality by Depth is computed by dividing the quality score against allele depth, to obtain an average score of allele quality. This is provided by Haplotype Caller and Unified Genotyper.

Mapping Quality

Mapping quality is a score provided by the alignment method and gives the probability that a read is placed accurately. It is provided by all variant callers except Pindel.

2.6 Patient Derived Xenograft Mouse Model Development and Sequencing

[TO CLARIFY] To test the Bayesian ranking system, we used a patient-derived xenograft mouse model. Athymic mice with the FOX mutation were grown for X number of days, and subsequently, a tumour is grafted onto the mouse's body. Subsequently, the tumour was sequenced on an Illumina MiSeq platform and used for analysis.

3 Results

3.1 Generation of Artificial Datasets

Using a genome mutation software, we generated a mutated genome using the hg19 genome from UCSC (Karolchik et al., 2014) as a reference. The mutated genome contains over 300,000 random mutations spread over the chromosomes as can be seen below in Figures 3 and 4. Artificial genomes are a good method to analyse deep learning networks on as the ground truth, which are the truth variants inside the genome, are already known. This allows accurate verification of prediction schemes and is a commonly used method to test next generation sequencing related software (Escalona, Rocha & Posada, 2016). This is primarily because it is difficult to obtain complete truth datasets for real genomes as due to the inhibitory cost of checking every variant called via Sanger sequencing. Thus, artificial genomes present a simple way to simulate NGS data with perfectly known ground truth variants to test our validation platform.

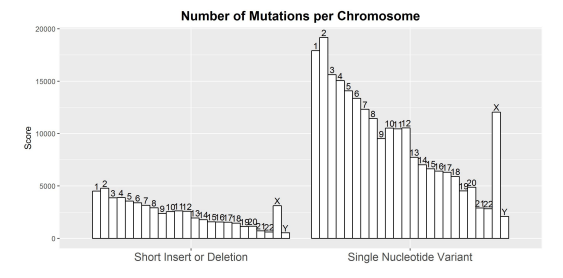


Figure 7: Number of ground truth mutations (variants) created in each chromosome

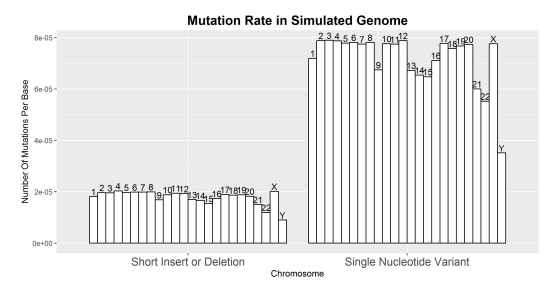


Figure 8: Mutation rate per base in each chromosome

3.2 Feature Engineering

Subsequently, we engineered a set of 19 features to use as input data for our variant callers, using data obtained from the variant callers themselves as well as engineering other features from the dataset. A summary of the features used in training can be found in Table 1, and a description of the full list of features can be found in Appendix 5.2. Features were engineered based on obtaining information on the main aspects of variant calling, which includes the information contained in the sample bases (Base Quality, Entropy, Kullback–Leibler divergence, etc.), the confidence we have in the calling and alignment (Read Depth, Mapping Quality etc) and finally possible biases in the sequencing machine (Allele Balance, Allele Count, GC content).

Table 1: Feature Engineering Table

	Shannon Entropy (Reference, Alternate and	Base Composition (Homopolymer Run,	Read	Mapping	Base	Allele	Quality	Allele	Genotype
Features	KL- Divergence)	GC content)	Depth	Quality	Quality	Balance	by Depth	Count	Likelihoods
Free Bayes	+	+	+	+	+	+			+
Haplotype Caller Unified	+	+	+	+	+		+	+	+
Genotyper	+	+	+	+	+	+	+	+	+
Pindel	+	+	+						+
Samtools	+	+	+	+	+	+			+

3.3 Variant Callers

Variant callers were chosen for our deep learning neural network based on their orthogonal calling and reference methodologies - we wanted to optimise the information that the neural network receives (See Table 2). We used two haplotype-based callers, FreeBayes (Garrison & Marth, 2012) and GATK Haplotype Caller (McKenna et al. 2010, DePristo et al. 2011), two position based callers GATK Unified Genotyper and Samtools (Li H, et al., 2009) and finally Pindel, a pattern growth based caller (Ye et al., 2009). When we analysed the concordance rates of the callers on the simulated dataset, we found a high amount call discordance (Figure 8).

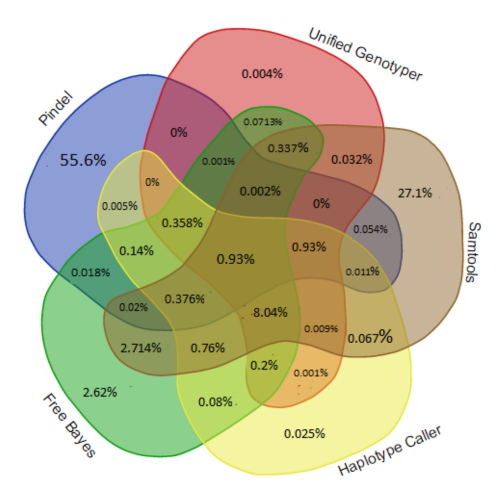


Figure 9: Concordance of callers on simulated dataset, using default settings

Of all the callers, Pindel was the most discordant caller, with over 1.6 million (55.6%) unique calls that are different from other calls. Samtools was also very discordant, with

over 800 thousand unique calls (27.1%) that were unique from the other callers, followed by FreeBayes at 80,000 calls. Interestingly, a high amount of calls (about 100,000) also exists in the intersection of only two callers. Discordance in the variant callers can be explained by the different methodologies that they use to call variants (Table 2). Due to implementation and design choices, as well as statistical methods, each variant caller has a different calling profile. Discordance in the callers provides a strong argument using deep learning to integrate the information from all the callers in a sophisticated manner.

Table 2: Table Comparing Methods and Features of Different variant callers.

	GATK Unified Genotyper	Samtools	GATK Haplotype Caller	Free Bayes	Pindel
Calling Method	Uses a list of mapped reads, calling model is probabilistic with increased priors at regions with known SNPs	Uses a list of mapped reads, calling model is probabilistic. Does not assume sequencing errors are independent and has less hard filters compared to Unified Genotyper	Uses Hidden Markov Models to build a likelihood of haplotypes which are then used to call variants	Uses a posteriori probability model to build a set of haplotypes to represent mutations, calling model is probabilistic with population based priors	Locates regions which were mapped with indels or only one end was mapped, and then performs a pattern growth to find inserts and deletions. Shown to be able to identify medium length indels missed by other callers in real samples (Spencer et al., 2013)
Reference and Mapping Method	Position based caller that realigns fragments and analyses each position to call SNPs and indels	Position based caller that uses mapped sequences to call SNPs and indels.	Analyses regions where there is high likelihood of mutation based on activity score, about the builds a De Bruijn-like graph that reassembles reads (Haplotypes) in that region	Dynamic sliding window based reference frame, using algorithms to determine window size for analysis. Does not require precise alignment, unlike other callers	Focuses on Unmapped regions, regions known to have insert and deletions or regions with only one end mapped.

3.4 Network Architecture

Before training our deep learning network, we tested out various neural network architectures to see which architecture would perform the best for our set of input features. We first explored the flat architecture (Figure 10), which contains stacks of fully connected layers with multiple nodes (initially seven layers, with 80 nodes per layer). This is the simplest architecture, where all the features are loaded onto a single vector, and this entire vector is used as an input to train the neural network. We next explored the PCA + flat architecture which had the same neural network architecture but before the input data was fed into the network, a Principal Components Analysis was done to reduce the dataset to 8 principal components which were then used as input data for the neural network (please see Appendix 5.3 for more details of the PCA analysis). Principal components analysis is a dimensionality reduction technique that enables a compressed representation of data. Each

principal component is a linear summation of the original features (X) in the form

$$PC_1 = \beta_{1,1} * X_1 + \beta_{2,1} X_2 + \dots + \beta_{n,1} X_n$$

$$PC_i = \beta_{1,i} * X_1 + \beta_{2,i} X_2 + \dots + \beta_{n,i} X_n$$

which enables a few principal components to capture a high amount of variance in the dataset. Finally, the last architecture we tested was the merged network this network had a set of layers (initially five layers, 24 nodes per layer) that learns from each caller alone, and then the outputs from each of these layers are subsequently merged and used to make a prediction.

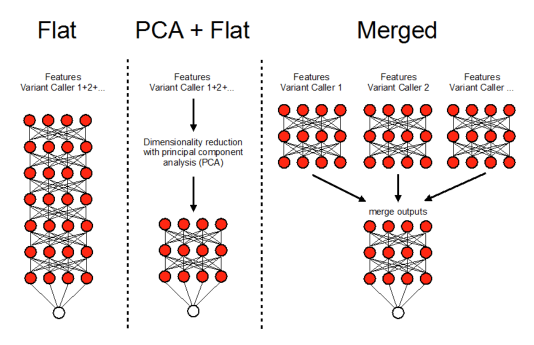


Figure 10: Different Designs for Neural Network Architecture

To study how well each architecture is able to perform, we use the metrics of precision, recall and F1 score. Precision measures how many mistakes the predictor makes (the ratio of true positives over false positives and true positives), recall measures what portion of the truth class can the predictor discover (the ratio of true positives over true positives and false negatives) and finally the F1 score is composite function of both precision and recall.

The derivations of the metrics can be found in Appendix 5.3.

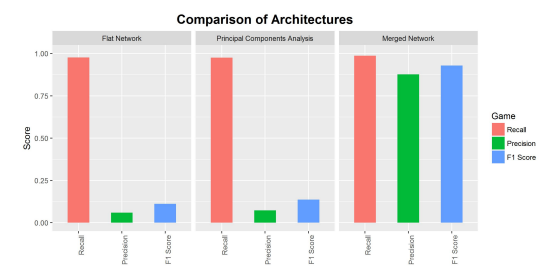


Figure 11: Analysis of Different Neural Network Architecture

Initially, with the flat network, the precision rate was very low at 0.059 with an F1 score of 0.112, indicating that the neural network was unable to learn from the input feature set. We suspected that this was due to high dimensionality in the dataset, which led to our second architecture design, the PCA with flat analysis. Principal components analysis has been shown to be able to successfully improve learning in high-dimensionality datasets (Chen et al., 2014; Van Der Maaten, Postma & Van den Herik, 2009). However, the precision and F1 score for the PCA architecture was also low at 0.0735 and 0.137 respectively. Ultimately both failed to learn, indicating to us that perhaps the features from each of the callers had to be analysed separately before being passed into a separate neural network that did the final score computations. With this merged network, we managed to obtain a precision score(0.877) and an F1 score(0.929) that was far better than the previous two architectures. Interestingly, the recall scores for all three architectures were around the same (±0.01), indicating the main learning point for the neural network was removing false positive calls.

3.5 Network Tuning and Optimisation

Next, we systematically optimised and tuned the deep learning neural network to maximise its predictive ability. In tuning our network, we also sought to study how the various hyperparameters as well as the data structure affected our network's ability to learn from the data. In particular, we focused on four issues - the number of layers, optimiser choice, learning rate choice and finally sample balancing. These four issues are known to be critical in deep learning networks (Ruder et al., 2016; LeCun, Bengio & Hinton, 2015; Yan et al., 2015; Sutskever et al., 2013) and would likely be critical to the success of a neural network.

3.5.1 Number of Layers

Firstly, we studied how many layers should be in the neural network. The number of layers is critical as it determines what kind of information and the representation of data that can be captured by the neural network. Choosing the number of layers is important as sufficient layers are needed to obtain the complex data representation needed for learning, but too many layers might result in the vanishing gradient problem. We started off with a neural network architecture as shown below, and began to vary the number of layers in at each point.

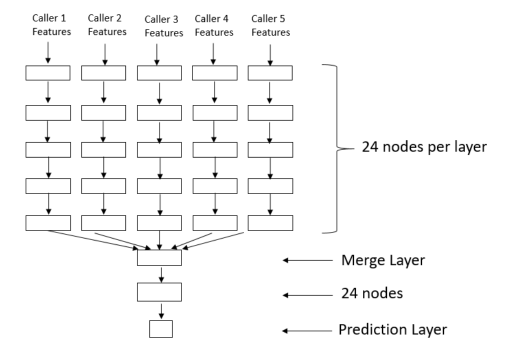


Figure 12: Basic Merge Network Structure

For all layers, the LeakyReLU activation function was used. The LeakyReLU is a refinement of the ReLU activation function which minimises the "dying ReLU" problem,

and both are well-documented activation functions that have been shown to work well in deep neural networks (Anthimopoulos et al., 2016; LeCun, Bengio & Hinton, 2015; Maas, Hannun & Ng, 2013). We noticed that changing the number of layers after the merge layer did not significantly vary the output, and so we focused on changing the number of layers before the merge layer. We studied 6 different neural network structures (4 layers to 9 layers).

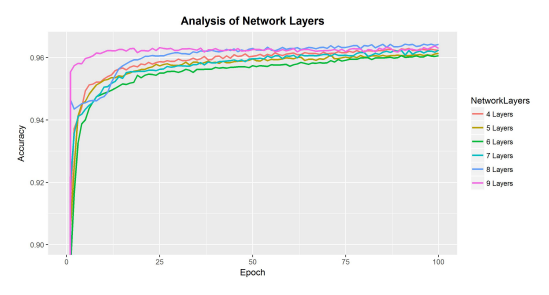


Figure 13: Analysis of Different Number of Layers On Training Accuracy

From analysing the accuracies of the different layers, we find that 8 layers seem the best at learning from the input data. We note that all the layers follow the same rough trend of accuracy, indicating they are all able to learn from the dataset. We decided on the 8 layer network as it seemed best able to learn from the input dataset, with a final accuracy of 0.964 that is about 0.001 higher than other layers. A final design feature used was to add two dropout filters at the last two layers before merging in order to prevent overfitting in data. Dropout filters have been shown to be an effective in preventing overfitting of data (Srivastava et al., 2014). Another active step taken to prevent overfitting was to ensure random separation of test, validation and training datasets.

3.5.2 Optimiser and Learning Rates

Next, we sought to choose the best optimiser and learning rate for our dataset. Both optimisers and learning rates have been well studied and known to be important in neural network training (Ruder et al., 2016; Sutskever et al., 2013). Optimiser choice is critical as the optimisers determine how the weights and gradients are updated in the network, thus playing an integral part in learning. We studied 3 well-known optimisers for use in our network, ADAM, RMSprop and Stochastic Gradient Descent (SGD). ADAM is an adaptive learning rate optimiser that is known to be well suited in large dataset and parameter problems(Kingma & Ba, 2014). RMSprop is another adaptive learning rate optimiser that unpublished, but has been shown to work well for real experimental datasets (Tieleman & Hinton, 2012). SGD is the simplest learning model with no adaptive learning rate but is a useful model because it is the easiest to understand mathematically and has also been shown to solve deep learning problems (Kingma & Ba, 2014). For more information on the mathematical foundations of optimisation and backpropagation, please see Appendix 5.1. For the three optimisers, we ran tests to study the accuracy of the neural network running on each optimiser to predict true variants (Figure 11).

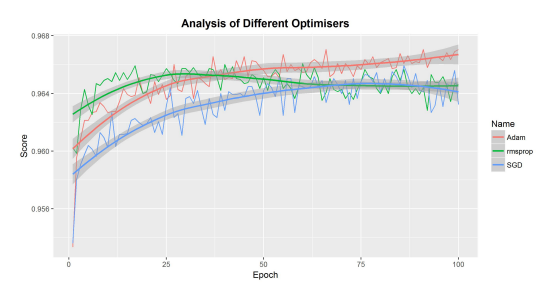


Figure 14: Optimiser accuracies for training at each epoch. Due to the noise in accuracies, the overall momentum of the dataset, calculated as a sliding window average is shown. The 95% confidence interval is also shown.

The adaptive rate optimisers, Adam and RMSprop obtained the highest accuracies of 0.9670 and 0.9660 while SGD reached a maximum accuracy of 0.9569, very close to

that of RMSprop. Interestingly adaptive rate optimisers seemed to have complex learning trajectories, while SGD has a very stable learning rate. This makes sense as adaptive learning rates allow greater gradient descents when the error is high, and decreasing the learning rate at smaller errors (Kingma and Ba, 2014; Zeiler, 2012). This allows Adam and RMSprop to learn at variable rates based on the current gradients. For SGD, it appears that while it takes a while to learn the true minima, it eventually still reaches about the same minima as RMSprop. Ultimately, we chose Adam as our optimiser as the final accuracy discovered by Adam was noted to be higher than RMSprop and SGD, and we note a stable learning curve for Adam, indicating it is able to learn and update the gradients in the neural network to learn from input data at all epochs. Subsequently, we also looked at various initial learning rate for Adam (Figure 12) and found that the most stable learning could be found at a learning rate of 10^{-5} . This initial learning rate is critical as it determines the first few gradient descents which enable stable adaptive learning throughout the epochs (Sutskever et al., 2013). At any larger learning rates (10^{-4} and below), a very high amount of noise was observed, indicating that the learning rate was too high resulting in minima finding errors. At smaller learning rates (10^{-6} and above), the final accuracy after 100 epochs $(0.9639 \text{ for } 10^{-6})$ was lower than the learning rate at 10^{-5} (0.9672). Thus, we chose 10^{-5} to be our learning rate.

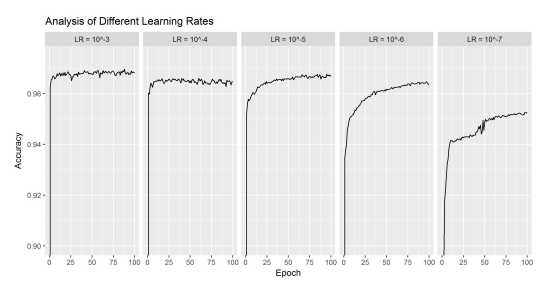


Figure 15: Training Accuracies over Each Epoch for Different Learning Rates

3.5.3 Sample Balancing

Our final concern was sample balancing - the simulated dataset contained an imbalance of positive training examples versus negative training examples. In total, there were 286569 positive training examples and 4547919 negative training examples, which is a 15-fold difference. Such a sample imbalance has been known to affect learning adversely (Yan et al., 2015; López et al., 2012). Thus, we sought to implement two methods of sample balancing, undersampling and oversampling. Undersampling was implemented by removing negative training examples until the number of negative training examples was equal to the number of positive training examples. In oversampling, the Synthetic Minority Oversampling Technique(SMOTE) was done, which uses nearest neighbours to create more data points for the positive training example. Specifically, SMOTE looks at two nearby positive class examples, and creates a new synthetic example in the middle of these two examples (see Appendix 5.3 for more details). Figure 13 shows the metrics for each sampling technique.

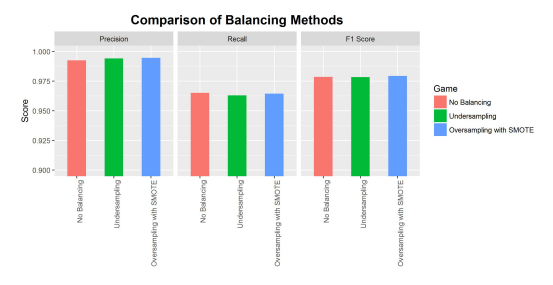


Figure 16: Effect of Sample Balancing on Prediction Ability

Interestingly, we note that overall, undersampling, oversampling and no sampling at all had very small effects on precision, recall and the final F1 score. Specifically, all three metrics were within a range of 0.003 for oversampling, undersampling and no sampling. This could be due to clear boundary separation within positive and negative class examples as well as good representative datapoints within the positive training example class. This prevents the imbalanced data from having too much of an effect on variant prediction and

classification. Still, we note that oversampling techniques resulted in a marginally higher F1 score (0.001 higher than undersampling and no sampling), and since ensuring that datasets are balanced is a recommended protocol to prevent further bias downstream, we used SMOTE oversampling to produce extra positive training class examples for all analysis pipelines.

3.6 Benchmarking of Optimised Network with Mason Dataset

From optimisation steps, we finalised the network architecture as seen in Figure 9, but with 8 layers before the merge layer. We chose the learning rate to be 10^{-5} , and the optimiser used was Adam. With this network, we benchmarked the neural network against the single variant callers, as well as concordance callers, which are an integration of the outputs of the 5 variant callers. Specifically, the n-concordance variant caller is defined as the set of calls that any n callers agree upon - so 1-concordance includes all the calls made by all callers and 4-concordance includes all the calls made by any 4 callers.

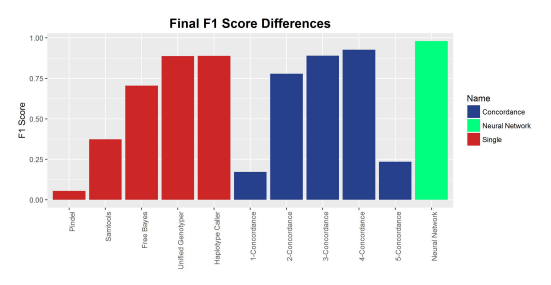


Figure 17: Overall Comparison of Variant Callers

In terms of overall F1 score, we see that the neural network was able to outperform single and concordance-based callers. This provides strong evidence that the neural network is able to learn from the input features whether the variant call is real or not, validating its usage in variant calling. The final F1 score obtained by the best single variant caller was Haplotype caller at 0.888, the best concordance caller had an F1 score of 0.927 while the neural network achieved an F1 score of 0.980. To study whether the increase in F1 score

is due to improvements in precision or recall, we studied the exact precision, recall and F1 scores of the top 2 variant callers as well as the best single variant caller versus the neural network. We find that the neural network is more precise than both, but the recall is rather similar.

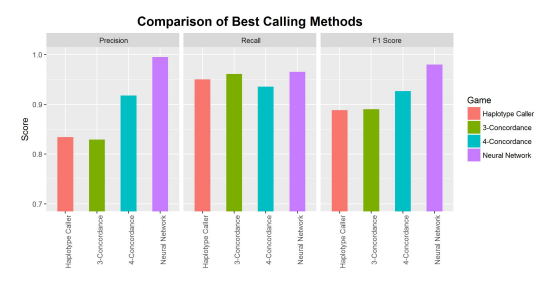


Figure 18: Comparison of Best Variant Callers in terms of Precision, Recall and F1 Score

We find that the neural network is more precise than all four callers, with the neural network had a high precision of 0.995 compared to only 0.917 for 4 concordance. This is a 20 fold decrease in the number of false positives or about 23,000 more false positive calls in the 4 concordance network compared to the neural network. Interestingly, the recall of all the callers was high in the range of 0.90 to 0.95, indicating that while all were able to pick out most of the truth class variables, the main errors came from a high number of false positives that were also called as true mutations. Ultimately, the neural network had an F1 score that was 0.1 above the best single caller and 0.06 above the best concordance caller. Thus, this provides strong evidence that the neural network is able to sieve out false positives within the dataset and stably predict whether a mutation is true.

3.7 Benchmarking of Optimised Network with NA Dataset

After verification of the neural network architecture, optimised parameters and ability to learn with a simulated dataset, we sought to analyse a real dataset to set the validity of the neural network in validating variants. We studied the NA12878 Genome In a Bottle dataset (Zook et al., 2014), which has been used in other variant calling valida-

tion pipelines (Talwalkar et al., 2014; Linderman et al., 2014) and contains a set of high-confidence variant calls which we can use as ground truth for training and validation. This set of high-confidence variant calls is obtained from multiple iterations of orthogonal sequencing methods (using Solid, Illumina platforms, Roche 454 sequencing and Ion Torrent technologies). The usage of multiple platforms enables an intersection of variants that can be considered as the ground truth. We then sought to see if our neural network can predict the ground truth better than single or concordance based variant callers.

We applied the same methodology to the sequences as with the simulated data and then used our neural network to predict the true variants. Validation was done with 47971 high-confidence variant calls.

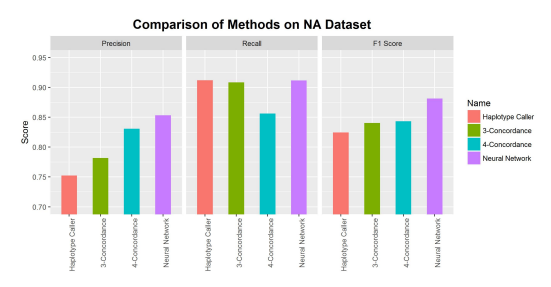


Figure 19: Comparison of Variant Callers

As can be seen from the figure, the neural network was able to predict with the highest precision (0.859) when compared the best single caller, haplotype caller (0.752) and the 2 best concordance callers, 3-concordance (0.782) and 4-concordance (0.830). In terms of recall, the neural network had a higher recall (0.911) compared to the 4 concordance caller (0.856). Thus, we see that in the NA dataset, the neural network compared with the 4 concordance network is able to call 2650 true variants that were missed by 4 concordance and still had 1228 less false positives. This means the neural network was more aggressive in making calls, yet more of the calls were correct. Compared to the three concordance caller, the neural network had 4253 less false positives. Ultimately when we looked at the

F1 score, the neural network was able to outperform concordance variant callers by at least 0.04 and single callers by 0.06. This validates our neural network pipeline in a real genomic dataset and indicates that network is able to learn from the input features.

3.8 Analysis of Gene Importance using Bayesian Ranking systems

After validation of high confidence calls, we sought to enable a clear and understandable ranking of genes. We first build a Bayesian network analysis using known functional annotations from ANNOVAR. These were subsequently used to compute the Bayesian probability ranking, which is shown in Figure 17 below.

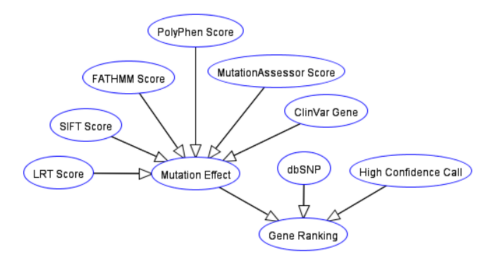


Figure 20: Final Bayesian Network used in Analysis

This network structure was chosen as we wanted to use three different sets of information to update the probability of the gene being important. Firstly, the confidence of the call should matter in how important it is - the more likely a gene is real, the more important it should be. Secondly, the rank should also be determined by how common the variation is, based on studying known SNP polymorphism rates. If it is a common SNP, then the ranking should be downgraded as it is less likely to be a driver mutation (Schork et al., 2009). Finally, we sought to predict the overall effect of mutations via an ensemble of mutation effect predictors. These predictors use different methods to predict the average effect of that mutation - based on statistical methods like position-specific substitution matrixes and Hidden Markov Models to study the effect of a mutation on protein structure and function. We also used the ClinVar database, a curated repository of known Human variants and their resulting phenotypes (Landrum et al., 2014). These scores were then aggregated to update the probability of the mutation effect. To obtain these functional annotations, the informatics tool ANNOVAR (Wang, Li, & Hakonarson, 2010) was used. Table 3 shows the functional annotations obtained from ANNOVAR and how they were

computed.

Table 3: Table of Functional Annotations obtained from ANNOVAR

Annotation Name	Information Type	Method	Scoring Method
Likelihood Ratio Test	Deleterious Mutation Score	Likelihood Ratio Test of each amino acid is evolving neutrally to the alternative model of evolution under negative selection	Score normalised to [0,1] and used directly in Bayesian Network
MutationAssessor	Deleterious Mutation Score	Mutation rate of homologous sequence subfamilies	Score normalised to [0,1] and used directly in Bayesian Network
SIFT	Deleterious Mutation Score	Position Specific Scoring Matrixes with conserved Sequences	Score normalised to [0,1] and used directly in Bayesian Network
PolyPhen2	Deleterious Mutation Score	naïve Bayes classifier on various multiple sequence alignments methods of homologous proteins and protein structure-based features	Score normalised to [0,1] and used directly in Bayesian Network
FATHMM	Deleterious Mutation Score	Hidden Markov Model used to score MSA based on protein homologous sequences	Score normalised to [0,1] and used directly in Bayesian Network
ClinVar Genes	Known Pathogenic Genes	Database lookup of curated set of relationship between variant calls and human phenotype	Higher Probability of Importance if known pathogenic variant
dbSNP138	Common Single Nucleotide Polymorphisms	Database lookup of curated set of known Human SNPs	Lower Probability of Importance if known common variant

Based on scores provided, we report the update the conditional probabilities using the probabilities chain rule - for the first level, this is given as

$$P(Impt|(Del \cap Uncom \cap High\ Conc)) = P(Impt \cap Del \cap Uncom \cap High\ Conc)$$

$$* P(Del \cap Uncom \cap High\ Conc)$$

$$(3)$$

P(Impt) refers to the probability of the gene being important,

P(Del) refers to the probability of the gene being deleterious,

P(Uncom) refers to the probability of the gene being uncommon and

P(High Conc) refers to the probability of the gene being a high confidence call.

Further calculations can be found and derivations can be found in Appendix A

To compute the final probabilities, the software pomegranate was used. This simplifies the node drawing and probabilistic updates of the final ranking scores.

3.9 Validation of Bayesian Network Ranking on PDX dataset

To study the effectiveness of our Bayesian network ranking system, we sequenced and analysed a patient-derived xenograft (PDX) tumour genome. This tumour genome was grafted onto the immunocompromised mouse from a patient with a known cancer subtype - Diffuse Large B-Cell Lymphoma (DLBCL). We chose to analyse lymphoma as it is a well-known and studied disease model with a well-defined disease progression (Knudson et al., 2001; Alizadeh et al., 2000). The patient-derived xenograft model also allows *in vivo* studies of the tumour in its environment and serves as a good model for sequencing and

analysis (Tentler et al., 2012). After sequencing the PDX genome, we put it through our full analysis pipeline, which involves identifying high-confidence mutations using the neural networks and then ranking these genes using the Bayesian network ranking. Figure 19 shows the top 30 genes by probability.

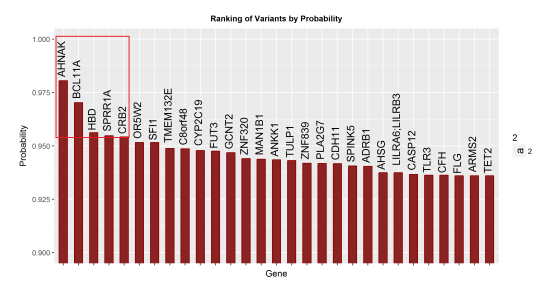


Figure 21: Top 30 genes from Bayesian Ranking Algorithm

Studying the top 5 genes, we found that four of these five genes have been implicated in lymphomas or other cancers. AHNAK is a known tumour suppressor and has been known to be downregulated in lines of Burkitt Lymphoma. BCL11A is a known proto-oncogene in DLBCL and has been found to be overexpressed in 75% of primary mediastinal B-cell Lymphomas, a subset of DLBCL. SPRR1A, the fourth gene ranked in terms of importance, has been shown to be expressed in DLBCL and its expression has been shown to strongly correlate with 5-year survival rate (Figure 21). Finally, development of B-cell lymphoma has been noted in Crb-2 related syndrome, which is a bi-allelic mutation of CRB2. Interestingly, the last of the high ranked genes was noted to be a subunit of Hemoglobin. While there is no strong evidence for the role of Haemoglobin in DLBCL, it has been shown to be expressed in aggressive glioblastomas lines, indicating a possible previously unknown role in cancer. This gives us high confidence that the Bayesian ranking network is able to pick up important and relevant mutations. Without such a ranking system, we would have to look through over 70 thousand genes, without a way to systematically study their likelihood of being important.

Figure 22: Table of Top 5 important genes from Bayesian Ranking

Gene	Full Name	Known Involvement in Lymphoma or Cancer	Evidence	Mutation Location	Predicted Mutation Type
AHNAK	Neuroblast Differentiation- Associated Protein (Desmoyokin)	Known tumour suppressor via modulation of TGFβ/Smad signalling pathway Known to be downregulated in cell lines of Burkitt lymphomas	Lee et al., 2014; Amagai et al., 2004; Shtivelman et al, 1992	chr11 - 62293433 T -> C	non synonymous SNV
BCL11A	B-Cell CLL/Lymphoma 11A	Known proto-oncogene in DLBCL Overexpression of BCL11A was found in 75% of primary mediastinal B-cell lymphomas (a subset of DLBCLs)	Weniger et al., 2006; Schlegelberger et al. 2001; Satterwhite et al., 2001	chr2 - 60688580 C -> G	non synonymous SNV
HBD	Hemoglobin Subunit • Delta	Shown to be expressed by aggressive glioblastoma cell lines	Allalunis-Turner et al., 2013	chr11 - 5255274 G -> A	stop-gain
SPRR1A	Small Proline Rich • Protein 1A (Cornifin-A)	Known to be expressed in DLBCL and expression has been shown to correlate with 5 year survival rate	Liu et al., 2014	chr1 - 152957961 G -> C	non synonymous SNV
CRB2	Crumbs 2, Cell Polarity Complex Component	Cell polarity and cytoskeletal reorganisation is known to affect B-cell lymphoma migration and invasiveness Development of B-cell lymphoma has also been noted in Crb2-related syndrome (bi-allelic mutation of Crb2)	Slavotinek, 2015; Gold et al., 2010	chr9 – 126135887 T -> C	non synonymous SNV

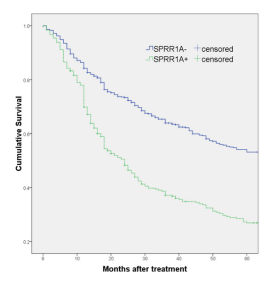


Figure 23: 5 year survival curve of patients with SPRR1A+ and SPRR1A- patients with DLBCL. Source : Zhang et al. (2014), Figure 2.

To aggregate the data from our Bayesian Ranking system, we did a Circos plot for the top 300 genes picked up by our gene ranking system. A Circos enables easy visualisation and analysis of large genome datasets, enabling quick understanding and comprehension of results. From the Circos Plot (Figure 22), we find several interesting gene families that might also be relevant in B-Cell Lymphoma. These include several Toll-Like Receptors(TLRs), TLR3 (chr4,rank 26) and Tlr1(chr4,rank 77) as well as interleukin receptors

IL4R (chr16,rank 37) and IL1 β (chr2,rank 196). TLRs are of significant interest in cancer due to their involvement in the caspase pathway (Kelly et al., 2006), and have been implicated in B-Cell Lymphomas (Marron, Joyce, & Cunningham-Rundles,2012). Interleukins are also important in cancer due to their importance in mediating inflammation and immune response (Balkwill & Mantovani, 2001). Thus, we show that our Bayesian network can be used by clinicians to quickly interrogate the information from functional annotations and database lookups to understand the disease specifics.

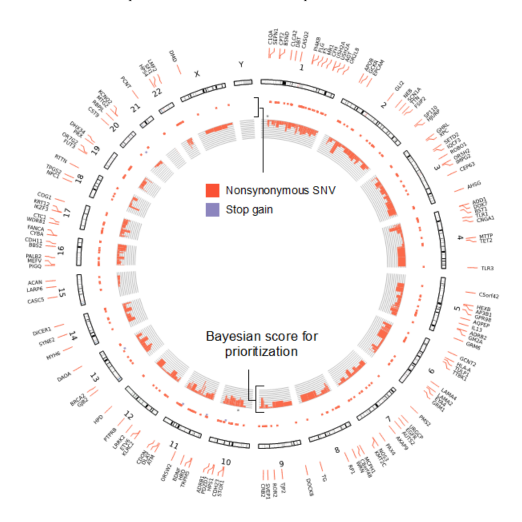


Figure 24: Circos plot of top 300 ranked genes from Bayesian network ranking. In this Circos plot, the outer track indicates the top ranked genes and their positions on the chromosome. The inner track describes the type of mutation that was observed - most mutations were non-synonymous SNVs, with a few stop-gain mutations. The innermost track shows the relative probabilities of each ranked gene.

4 Discussion

We demonstrate the validation of high-confidence variant calls using an optimised deep learning neural network on both real and simulated datasets, and we also show that a Bayesian network can rank and prioritise genes in a systematic way so as to obtain important genes. We show that four of the top five genes had published findings that linked them with lymphoma. Looking at the top 300 genes ranked, we also found interesting families of genes that are known to be involved in Lymphoma progression. To benchmark these results, we compared our variant calling results with other methods like VariantMetaCaller(using Support Vector Machines) and BAYSIC(using Bayesian Latent Analysis) (Gézsi et al., 2015; Cantarel et al., 2014), and we also looked at other methods of gene prioritisation to see how our ranking system compares.

4.1 Comparison of Deep Learning with other Integration Methods

First, we looked at other methods of integrating variant call information, including VariantMetaCaller, which uses Support Vector Machines (a decision making machine learning technique) and BAYSIC, a method that uses a Bayesian probabilistic model to integrate variant call information. Both methods were also used to analyse and predict variants for NA12878 data (Gézsi et al., 2015), but due to differences in metrics, processes and datasets, absolute comparison of precision and recall is difficult. Thus, it is more suitable to study the looking at relative improvements seen when using each method. VariantMetaCaller increased SNP prediction by 0.04 and indel prediction by 0.07 in terms of the Area Under Prediction Recall Curve (AUPRC) metric when compared to their best single variant caller. The AUPRC measures the precision differences at all levels of recall. BAYSIC also noted a 0.03 increase in SNP prediction and 0.05 increase in indel prediction compared to the best single variant caller. Numerically, this seems comparable to our results of a 0.06 increase in both indel and SNP prediction for the NA dataset compared to the best single variant caller. However, since we used the F1 score metric, instead of the AUPRC metric, an relative quantitative comparison is also not so simple. While the AUPRC metric provides evidence of precision and recall improvements at all levels of threshold (Fawcett et al., 2006), it does not provide evidence for a predictors performance at the best threshold. To measure this, the measurement of the F1 score at the best threshold is required - since it is the F1 score that looks at precision and recall for a specific threshold. Instead, what they have shown is that looking at all thresholds, there is an overall increase in precision and performance, but it is unclear what the improvements are at the optimised thresholds. Fawcett (2006) also mentions this problem, as he notes that 'It is possible for a high-AUC classifier to perform worse in a specific region of ROC space than a low-AUC classifier'. Here, AUC refers to the Area Under Curve, another term for the AUPRC, and ROC refers to the Receiver Operator Characteristics graph (Egan, 1976) which is the curve that the AUPRC uses. Thus, a higher AUPRC does not mean that one caller will outperform another when considering only the optimised threshold. Measurement of the F1 score is more relevant in clinical practice as we are mainly interested in the optimal operating conditions where precision and recall are maximised and not the fringe conditions. Our results provide specific evidence that at the optimal recall threshold for each specific type of caller, we can show an F1 score improvement of 0.06.

Thus, one definite step moving forward is to incorporate VariantMetaCaller and BAYSIC into our pipelines as negative controls, and measure using the same dataset and same processes whether deep learning is able to outperform these two methods using the same comparison methods and metrics. Intuitively, we believe that deep learning will be able to edge out improvements as deep learning can form complex representations of the data to learn from that Support Vector Machines are unable to do and ultimately outperform Support Vector Machines in decision problems (LeCun et al., 2015; Schmidhuber, 2015) Furthermore, evidence from our flat and dense network architecture shows that putting all the features in a single vector and using that to performing machine learning might not be the best method as it is difficult to learn features from it.

However, one large limitation in the overall approach of measuring each of the methods against the NA12878 dataset is that the high confidence calls provided is not the ground truth. Zook et al.(2014) themselves estimate a possible false negative or positive that for every 30 million bases in the NA12878 dataset. This is due to true variants that are not inside the high confidence dataset because of errors in one sequencing machine, or genomic regions that cause all sequencing machines to have similar biases and noise. Hence, this would result in misclassification and wrongly called false negative and false positive results,

thus skewing the classification results. To solve this problem, a lot of effort has to be put in to obtain a set of verified truth variants via gold standard Sanger sequencing (Tsiatis et al., 2014), but this might be prohibitively expensive for a large sample of mutations. Still, this would have to be done for us to have a good set of truth variables to test prediction software with before such software can be considered for use in actual treatment and diagnosis.

4.2 Analysis of Bayesian Network

For the Bayesian Network analysis, it is more difficult to numerically benchmark our results for gene prioritisation with current platforms. This is because currently used platforms are qualitative methodologies like gene panels (Olek and Berlin, 2002) or manual literature look-ups of disease-related genes, such as using the ClinVar database. While gene panels work well in a clinical setting, with NGS data it is hoped that as much information about a person's genome as possible can be used in treatment and diagnosis (Meldrum et al., 2011). Using a gene ranking system instead of just looking at a set of implicated genes might allow doctors to find out tease out possible homologs or interacting agents that might be related to the known deleterious genes (perhaps in the gene panel) and integrate that into their treatment and diagnosis.

4.3 Future Directions

One interesting extension we would like to move into in the future is to be able to integrate a druggable genome into the network, enabling the prioritisation of genes which have possible candidate drug targets. This method would look up a drug-gene interaction database, for example, DGldb (Griffith et al., 2013), and use the results to inform the importance of a gene. This would enable doctors to notice further possible drug candidates that would work very well on the gene profile of the patient that they might not have considered previously, thus increasing their scope of possible treatment options and augmenting their skills. Other directions in the future include being able to include extra variant callers which will provide it with even more feature data, enabling it to make better predictions. We also hope to build structural variant calling neural networks, as this is a current set of variants that our neural network does not take into account. Finally, we would also like to move everything onto a web interface such that it is accessible for use in terms of both variant validation and gene prioritisation. This would enable easy access to both the validation and prioritisation

pipelines.

Thus, in this paper, we have shown the use of deep learning neural networks to successfully validate variants in both real and simulated datasets. We also show that using a Bayesian network can identify important genes within a lymphoma disease sample. Ultimately, we hope to be able to put these networks to use in a clinical setting to augment treatment and diagnosis of diseases.

5 Appendixes

5.1 Neural Network Learning

Machine learning is underpinned by two key phases, the feed-forward phase and the backpropagation phase. The feedforward phase describes the computation of a prediction, and during this phase, the input features are used to compute the final output prediction. For a simple network below:

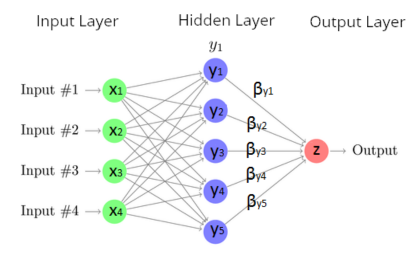


Figure 25: Example neural networks with nodes and weights

The final prediction, z is computed with the equation :

$$z = \beta_{u1} * y_1 + \beta_{u2} * y_2 + \beta_{u3} * y_3 + \beta_{u_4} * y_4 + \beta_{u_5} * y_5$$

$$\tag{4}$$

Where β indicates the weights linking each output to the input of z and each of the y_i terms are computed in the same manner from the x_i layer. At each node (x,y,z), there is also the existence of an activation function that modifies the input of the node to compute an output. Commonly used activation functions include the rectified linear unit (ReLU), sigmoid functions like hyperbolic tangent and logistic function, $S(T) = \frac{1}{1+e^{-t}}$. Thus, the final prediction can be seen as a summation of all weights multiplied by the activation output of each node. In theory, we can expand each of the y_i terms in equation (2) to include the y_i layer activation function as well as rewrite the y_i layer inputs in terms of the sum of outputs and weights from the x_i layers. This complex integration of terms allows for the neural network to form complex continuous decision boundaries as the neural networks can

compute sophisticated non-linear prediction functions despite being a fundamentally linear model.

After a prediction is made, we then have to check whether it is correct, and change our weights if an erroneous prediction was made (Figure 24).

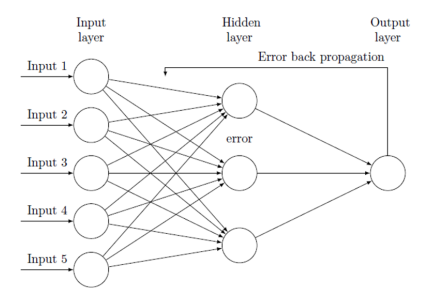


Figure 26: Backpropagation of Error Terms

This is the backpropagation step, which involves backpropagating the error terms from the output layers to the input layer and updating the weights at each node based on the differential relationship between the error and each specific gradient. Specifically, this is governed by the optimiser functions which have been mentioned earlier - one example of such an function is the Stochastic Gradient Descent function, which is

$$\beta_{yi}^{n} = \beta_{yi}^{n-1} - \alpha \frac{\partial E_n(\beta)}{\partial \beta_i} \tag{5}$$

Here, each β term indicates a gradient, α is a constant for the learning rate and $\frac{\partial E_n(\beta)}{\partial \beta_i}$ is the term used to modify the weight of the gradient based on the cost function $E_n(\beta)$. The idea used in all backpropagation functions is gradient descent, where the contribution of the gradient term to the error is computed and the gradient is changed by an amount in order to reduce the future contribution of the gradient to that error. Here it is useful to consider what the cost function $E_n(\beta)$ is. It is essentially the error rate when a set of gradients is

used to perform predictions, as it measures how many accurate predictions were made and how many wrong predictions were made. For a binary class predictor (which is what we are using, only true and false), this is given by the equation

$$E(\beta) = -\frac{1}{n} \sum_{i=1}^{n} \sum_{j=1}^{2} y_{ij} log(p_{ij})$$
(6)

where y_{ij} indicates the empirical observed probabilities of each class label while $log(p_{ij})$ is the theoretical probabilities of each class label. This is also known as binary cross-entropy, which is derived from Shannon's entropy (See Appendix 5.2.1). From this term, we see that if the neural network predicts something with a high probability (y_{ij} is high) and it is false (p_{ij} is low) so then $log(p_{ij})$ is a big negative number, and so the cost function will very high. On the other hand, if y_{ij} and p_{ij} is high then the entropy will be close to zero, indicating a correct prediction. Since each of the prediction terms can be rewritten in terms of the gradient(rewrite z in terms βy_i and so on, we can theoretically compute the contribution of each gradient to the cost function to see how the cost function changes as the gradient changes. Thus, this is what gradient descent does - it tries to see how the cost function that minimises the error term.

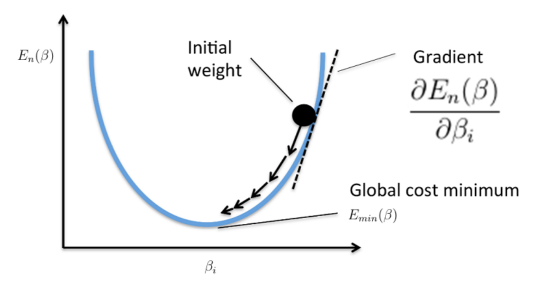


Figure 27: Gradient Descent, which attempts to find the gradient at which the cost function is minimised (since the cost function depends on the gradient).

This is best seen in Figure 27 above, where the gradient, or specifically the partial

differentiation of the cost function with regards to each gradient is used to move the gradient to a new position so as to minimise the error term. Thus, machine learning is in essence a minimisation problem - we want to find a set of weights that minimises the cost function, and because the cost function describes how many predictions we made correctly, this is also training our network to accurately predict outputs from inputs.

5.2 Feature Engineering

We subset our features into three broad sets, which are base-specific information, sequencing error and bias information features, and calling and mapping quality. Base information tells us base specific properties, including information contained in the base as well as the quality of sequenced bases in the samples. Sequencing error and bias features attempt to tease out potential biases in sequencing, including features such as GC content, longest homopolymer run and as well as allele balances and counts. Finally, calling and mapping quality provides information on the mapping and calling confidence of the variant callers, and includes features such as genotype confidence and mapping quality. In all, these sets of information provide information on the key aspects of variant calling - specifically the properties of the bases in the samples, the characteristics of the sequencing process and finally the variant calling and mapping algorithms.

5.2.1 Base Information

Shannon Entropy

Shannon Entropy captures the amount of information contained in the allele sequences. It is calculated using the equation :

$$H(X) = -\sum_{i=1}^{n} P(x_i) \log_2 P(x_i)$$
(7)

where $P(x_i)$ is the probability of finding each base at each position. Thus, we calculate the entropy by summing up the probabilities/log(probabilities) at each position. This prior probability is calculated in two ways and both are used as features - firstly, the overall genome base probabilities are calculated over the entire genome, and thus the entropy is related to the probability of finding a base at any position in the genome. The second way prior probability is calculated is to take a region of space around the allele (10 bases plus the length of the allele in our calculations) and use those probabilities to calculate the entropy of the allelic sequence. Intuitively, it attempts to find out the amount of information contained within the allelic sequence and hopefully the neural network is able to use the information to determine the validity of a mutation.

Kullback Leibler Divergence

The Kullback-Leibler Divergence feature is similar to Shannon entropy, but instead we use this to measure the informational change converting from the reference to the allele sequence. The Kullback-Leibler Divergence is calculated as follows:

$$D_{KL}(P||Q) = -\sum_{i=1}^{n} P(x_i) \log_2 \frac{P(x_i)}{Q(x_i)}$$
(8)

where $Q(x_i)$ is the prior probability of finding each base at each position based on the genomic region around the allele, while $P(X_i)$ is the posterior probability of finding a specific base inside the allelic sequence. Thus, the KL divergence describes the informational gain when the probabilities from Q is used to describe P. Intuitively, since we know the base probabilities of the region, we can then study the probabilities observed in the reference allelic sequence and see how well $Q(X_i)$ probabilities is able to approximate $P(X_i)$ probabilities.

Base Quality

Base quality refers to the Phred score probability that the called allele is wrong. It is given by the equation:

$$P = 10^{\frac{-Q}{10}}$$

Where P is the Base Quality and Q is the probability that the allele called is wrong. This is a number computed by the sequencing machine based on the quality of the base samples provided, and tells us how much confidence the sequencing machine has in calling that base.

5.2.2 Sequencing Biases and Errors

GC content

This feature computes the calculated GC content of reference genome, which may affect sequencing results and accuracy as regions with a GC content are known to be more difficult to sequence. This is because of the greater strength of GC bonds, resulting in errors and biases in sequencing (Benjamini & Speed, 2012).

Longest homozygous run

Homopolymer runs (AAAAAAA) are known to cause sequencer errors (Quail et al.,2012), and might be a factor in determining whether a variant is true. This because long homopolymers provide the same type of signal to the sequencing machine, resulting in a difficult in estimating the magnitude of the signal or rather how many bases are in that homopolymer, resulting in errors and wrongly called variants. The reference sequence region including the allele was checked for homopolymer runs.

Allele Count and Allele Balance

Allele count gives the total number of alleles in called phenotypes, while allele balance gives the ratio of final allele called over all other alleles called (reference allele for heterozygous calls, or other alleles for homozygous calls). Both these features give us information of possible biases in the sequencing machine.

5.2.3 Calling and Mapping Qualities

Genotype Likelihood

The genotype likelihood provides the Phred-scaled likelihood scores of how confident the caller is in determining that it is a homozygous or heterozygous call, and for the homozygous calls whether it is a more likely to be a bi-allelic mutation or no mutation at all. This feature thus gives us the confidence of the caller in determining if one or two alleles have mutated.

Read Depth

Mapped read depth refers to the total number of bases sequenced and aligned at a given reference base position. The read depth tells us how many reads contributed to a specific call, and thus provides information on how much evidence there is for the variant call

Quality by Depth

Quality by Depth is computed by dividing the quality score against allele depth, to obtain

an average score of allele quality. This composite feature provides information on the information provided by each read supporting the call

Mapping Quality

Mapping quality is originally a score provided by the alignment method and gives the probability that a read is placed accurately. The variant callers compute a overall mapping quality of the reads that provide evidence for a variant call which is given in this feature. A low mapping quality means that there are multiple positions where the reads contributing to this variant call could have gone, and thus providing evidence that this might not be an accurate call due to poor mapping.

5.3 Mathematical and Statistical Tools

5.3.1 Derivation of F1 Score

The F1 score is a useful measure as it can measure both the precision as well as the recall of a predictor. For a binary predictor with a binary truth class(Figure 26), we can obtain 4 types of results - true positives, true negatives, false positives and false negatives.

		Predicted Class		
		Yes	No	
Actual Class	Yes	True Positive	False Negative	
	No	False Positive	True Negative	

Figure 28: Confusion Matrix

True positives are positive predictions that are made that are actually positive class labels, while false positives are positive predictions that are made that have negative class labels. Similarly, true negatives are negative predictions that have negative class labels, while false negatives are negative predictions that are actually positive class labels. From this, we can define two equations, precision and recall. Precision is defined as (8) while recall is defined as (9).

$$Precision = \frac{True\ Positive}{True\ Positive + False\ Positive} \tag{9}$$

$$Recall = \frac{True\ Positive}{True\ Positive + False\ Negative}$$
 (10)

Precision tells us how likely a positive prediction made will be true, while recall tells us how much of the truth class positive predictions the predictor is able to encompass. Thus, a predictor can have a high precision but low recall (makes few predictions but are very accurate) or a high recall and low precision(makes a lot of predictions that capture all truth variables, but have a lot of false positives as well). In genomics, both types of errors are not desired - we would want all the predictions to be true (precision), while not losing out on any important mutations (recall). Thus, we use the composite metric, the F1 score, that

looks at the overall precision and recall of a predictor. It is defined as follows :

$$F1 \ Score = \frac{2 * Precision * Recall}{Precision + Recall}$$
 (11)

5.3.2 Principal Components Analysis (PCA)

Principal Components Analysis (PCA) is a commonly used tool for dimensionality reduction. It was first proposed by Pearson in 1901 (Pearson, 1901) and has been commonplace in many data analytics and signal processing methodologies (Jolliffe, 2002). PCA works by attempting to discover orthogonal principal components (PCs) that are able to represent the original data. Specifically, this means that the PCs are able to capture variance in the datasets. This is done by finding the Eigenvalues and Eigenvectors of the dataset, with the eigenvectors representing a linear combination of all input variables and the eigenvalues representing the amount of variance that that eigenvector is able to represent. Ultimately, we select n eigenvectors that is able to represent a percentage of variance in our dataset. Because each eigenvector is orthogonal, they are able to capture the variance in the dataset. For our analysis, we decided to use 8 principal components - we took the limit as the last principal components that was able to represent at least 0.5% of variance in the dataset.

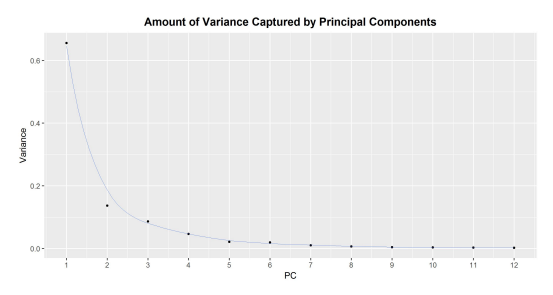


Figure 29: Variance captured by first 12 principal components

To carry out PCA, we used the preprocessing step SciPy to normalise all the input vectors to mean 0 and standard deviation 1. Subsequently, we perform principal components decomposition to obtain the eigenvector transformed representation of the dataset and their corresponding eigenvalues. We then fit 8 of the principal components that explained the largest amount of variance into the neural network to study if it can learn from the compressed representation of the input features.

5.3.3 Synthetic Minority Overrepresentation Technique (SMOTE)

SMOTE is a statistical technique described in by Chawla et al. (2002) to overcome problems with imbalanced datasets that are common in machine learning. SMOTE oversamples the training class with fewer variables in a way that tries not to replicate data points (that makes certain data points over-represented) without creating new invalid training examples. It does this by taking the intersection of two nearest data points of the same training class. This can be seen in Figure 28.

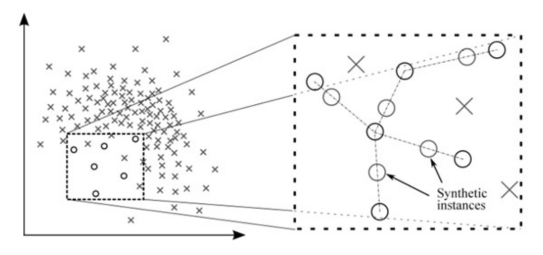


Figure 30: SMOTE oversampling algorithm

In doing so, it creates a more generalised representation of the sample class with less training examples, without replicating certain datapoints and without creating invalid data. This enables intelligent oversampling of the dataset to balance out the positive and negative feature classes. SMOTE has been shown to be valid for other datasets including sentence boundary detection (Liu et al., 2006) and data mining (Chawla, 2005).

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7 Relevant Code

3 code segments are provided to clarify the implementation of generating the matrixes for deep learning, deep learning networks and finally bayesian networks. Other code segments not shown include the code base for parsing vcf input into features, concordance generators, NextFlow and Bash code used to simulate and process genomic data as well as to control deep learning and analytic pipelines, and other python helper scripts (e.g. comparing two VCF files, analysis with pre-trained network).

7.1 generate_matrixes.py

29

```
#This python script generates the set of matrixes to be used in deep learning, and then calls the main
         method that trains the deep learning network.
    #Input: a directory that contains all the vcf files for processing, as well as a truth file.
    #Output : np.arrays of features from generate_matrixes with accompanying truth labels, feature set lengths,
     → a dictionary of vcf object records, as well as the list of relevant sample features for easy reference
    #Notes :
    #Vcf files should have the "vcf" string in their name and truth file should have a "truth" string in its
     \rightarrow name.
    #No other file should be present in the folder
    #Overall Strategy :
    #Generate a dictionary of lists, where the keys are mutations, and the value is contains a matrix
        containing information of all five callers
    #Secondly, for each mutation label, check if it is inside the truth file or not. The truth is preloaded
     #Finally, pass the set of features with accompanying truth labels to the neural network
10
    #The main datastructure used are python dictionaries, which allows O(1) dictionary lookup times
11
12
    import os
13
    import time
    from ANNgenerateresults import * #this file contains all the main methods for actual neural network
15
     \hookrightarrow training
    from methods import * #this file contains all the methods for parsing each VCF entry into a numerical list
         of features
    #declare names of useful files that contains processed data to be saved
18
    LIST_OF_INPUTS_NAME = '/ANN/samplelist.p'
19
    TRUTH_DICTIONARY_NAME = '/ANN/truthdict.p'
20
    CALLER_LENGTH_FILE_NAME = '/ANN/callerlengths.txt'
21
    VCF_LIST_FILE_NAME = '/ANN/vcf_list.p'
22
    SCORES_NAME = '/ANN/scores.txt'
23
    Y_DATA_NAME = '/ANN/myydata.txt'
24
    X_DATA_NAME = '/ANN/myXdata.txt'
25
26
    #Initialise NUMBER_OF_CALLERS
    NUMBER_OF_CALLERS = 5
28
```

```
30
    # This method follows the typical input output processing pipeline
31
    # It takes in the user input, and loads it into local variables.
32
     \hbox{\it\# It then executes another method, main\_analyse\_samples\_and\_truth on the loaded variables } \\
33
    # Finally, it then saves files into a directory determined by the final variables, and calls the next step
34
     \hookrightarrow of the pipeline
    {\it \# the neural network training, which is } {\it main\_gather\_input\_execute\_prep\_output}
35
36
    def load_and_save_data(user_input):
37
        user_input = vars(user_input)
38
        input_samples, referencepath, output_location = load_references(user_input)
                                                                                         # load user input
        my_x_dataset, my_y_dataset, list_of_samples, truth_dictionary, length_of_caller_outputs, \
40
        vcf_record_list = main_analyse_samples_and_truth(input_samples, referencepath)
41
        save_files(output_location, my_x_dataset, length_of_caller_outputs,
42
                    list_of_samples, truth_dictionary, vcf_record_list, my_y_dataset)
43
        orig_stdout = sys.stdout #save print statements into stdout
44
        f = file(str(output_location) + SCORES_NAME, 'w')
45
        sys.stdout = f
46
        main_gather_input_execute_prep_output(length_of_caller_outputs, truth_dictionary, my_x_dataset,
            my_y_dataset, list_of_samples, output_location, vcf_record_list)
48
    # This method first prepares a dictionary of truth to be checked against. It then initialises
    # a dictionary of samples with all the keys, each key being a variant call, and then fills it up each key
     → with data from each caller
    # subsequently, it removes dictionary entries that are the wrong size, and then checks whether
    # each entry in the dictionary is true or not by looking up the truth dictionary
52
    # subsequently it performs array balancing, and converts the data to np.array, as well as the dictionary of
     \hookrightarrow truth
    # and list of called samples
54
    def main_analyse_samples_and_truth(path, referencepath):
56
        os.chdir(path)
57
        truthdict = generate_truth_list(path)
58
        print "truth dictionary generated at time :", time.time() - start
59
        callerlengths, list_of_called_samples, vcf_list = generate_input(path, referencepath)
60
        print "samples generated at time :", time.time() - start
61
        clean_truth_array, cleaned_sample_array = check_predicted_with_truth(list_of_called_samples,
62

→ truthdict)

        print "samples checked with truth at time :", time.time() - start
63
        cleaned_sample_array = np.array(cleaned_sample_array, np.float64)
64
        clean_truth_array = np.array(clean_truth_array)
65
        return cleaned_sample_array, clean_truth_array, list_of_called_samples, truthdict, callerlengths,
66
         \hookrightarrow vcf_list
```

67

```
# This method generates the truth dictionary, by iterating through the vcf file, parsing all the vcf
         entries and appending them all as keys in the dictionary
69
     def create_truth_dictionary(generated_truth_dictionary, truth_file):
70
         vcf_reader = vcf.Reader(open(truth_file, 'r'))
71
         for record in vcf_reader:
             if "GL" in record.CHROM:
                                          #Ignore non-regular chromosomes in our dataset
73
                 continue
             templist = []
75
             for item in record.ALT:
76
                 templist.append(str(item).upper())
                                                               #Alternates might be a list, so they have to be
                  \hookrightarrow saved as a immutable tuple
             generated_truth_dictionary[(str(record.CHROM), str(record.POS), str(record.REF).upper())] =
78
              79
     # This method generates the input dictionary, by first initialising the keys of the dictionary by iterating
      \hookrightarrow through the vcf file once, and then
     # Iterating through the vcf file again and parsing all the entries as input vectors
81
     def generate_input(path, referencepath):
83
         reference_dictionary = get_reference_dictionary_for_entropy(referencepath)
84
         base_entropy = get_ref_entropy(referencepath)
         full_dictionary = get_dictionary_keys(path)
86
         list_of_called_samples, callerlengths, vcf_list = fill_sample_dictionary(base_entropy,
87

    full_dictionary, path, reference_dictionary)

         return callerlengths, list_of_called_samples, vcf_list
88
90
     # This method goes through all the training variant calling files and extracts unique calls as keys in the
91
         sample dictionary
92
     def get_dictionary_keys(path):
93
         sample_dictionary = {}
94
         for vcf_file in os.listdir(path):
95
             if ignore_file(vcf_file):
96
                 continue
97
             vcf_reader = vcf.Reader(open(vcf_file, 'r'))
98
             sample_dictionary = create_dictionary_keys(vcf_reader, sample_dictionary)
         return sample_dictionary
100
101
     #This method ensures the feature vector is in the right order - the entries must always be in the order fb,
102
      \rightarrow hc, ug, pindel and st.
103
     def create_list_of_paths(path):
104
```

```
105
         list_of_paths = [0] * NUMBER_OF_CALLERS
         for vcf_file in os.listdir(path):
106
             if ignore_file(vcf_file):
107
                 continue
108
             if "fb" in vcf_file:
109
                 list_of_paths[0] = vcf_file
             if "hc" in vcf_file:
111
                 list_of_paths[1] = vcf_file
112
             if "ug" in vcf_file:
113
                 list_of_paths[2] = vcf_file
114
             if "pind" in vcf_file:
                 list_of_paths[3] = vcf_file
116
             if "st" in vcf_file:
117
                 list_of_paths[4] = vcf_file
         return list_of_paths
119
120
     # This method goes through all the training variant calling files and fills each entry in a sample
121
      # with data. If it is empty, it returns an array of length n, where n is the number of variables
     # that same caller would have provided.
123
     # Each caller has a different amount of variables because it contains different datasets
124
125
     def fill_sample_dictionary(base_entropy, sample_dictionary, path, reference_dictionary):
126
         callerlengths = [0] * number_of_callers
127
         index = 0
128
         total_mode_value = 0
129
         list_of_paths = create_list_of_paths(path)
130
         for vcf_file in list_of_paths:
131
             index += 1
132
             opened_vcf_file = vcf.Reader(open(vcf_file, 'r'))
             removaldict = iterate_over_file_to_extract_data(base_entropy, sample_dictionary,
134
                                                               reference_dictionary, opened_vcf_file, vcf_file)
135
             mode_value = get_mode_value(removaldict)
136
             add_length_to_caller_lengths_based_on_file_name(vcf_file, mode_value, callerlengths)
137
             refill_dictionary_with_zero_arrays_for_each_file(sample_dictionary, index, mode_value)
138
             total_mode_value += mode_value
139
         list_of_passed_samples, vcf_list = add_mode_values_into_list_of_samples(sample_dictionary,
140
          \hookrightarrow total_mode_value)
         return list_of_passed_samples, callerlengths, vcf_list
141
142
143
     # this method fills the dictionary with empty arrays with the same length as the ones that were supposed to
144
      \hookrightarrow be added
145
```

```
def refill_dictionary_with_zero_arrays_for_each_file(full_dictionary, index, length_of_data_array):
146
         empty_set = []
147
         for i in range(length_of_data_array):
148
             empty_set.append(0)
149
         for item in full_dictionary:
150
             checksum = len(full_dictionary[item][0])
151
             if checksum < index:
152
                  arbinfo = empty_set
153
                  full_dictionary[item][0].append(arbinfo)
154
155
     # this method iterates through all the files to extract data from each sample. It uses methods from the
157
     # methods.py function, which parses each record for data.
158
     def iterate_over_file_to_extract_data(base_entropy, sample_dictionary, recorddictionary, vcf_reader1,
160

    vcf_file):

         removaldict = {}
161
         for record in vcf_reader1:
162
             if "GL" in str(record.CHROM):
163
                  continue
164
             sample_name = get_sample_name_from_record(record)
165
             sample_data = getallvalues(record, recorddictionary, base_entropy, vcf_file)
166
             sample_dictionary[sample_name][0].append(sample_data)
167
             sample_dictionary[sample_name][1] = record
168
             create_removal_dict(sample_data, removaldict)
169
         return removaldict
170
     # this method counts the mode number of entries in the dictionary. Due to certain vcf files having multiple
172
      → possible number of entries for a field, this will create an error
     # as the size of the input arrays should always be constant. Thus, any sample that does not fit the array
173
      \hookrightarrow should be removed.
     \# TO-DO See if a better implementation can be done that doesn't reduce data available
174
175
     def create_removal_dict(sample_data, removaldict):
176
         count = 0
         count += len(sample_data)
178
         if count not in removaldict:
179
             removaldict[count] = 1
180
         else:
181
             removaldict[count] += 1
182
183
184
     # this method prepares the reference genome dictionary for use in entropy calculations
186
```

```
def get_reference_dictionary_for_entropy(reference_path):
         record_dictionary = SeqIO.to_dict(SeqIO.parse(reference_path, "fasta"), key_function=get_chr)
188
         return record_dictionary
189
190
     # this method ensures that the files inputed are correct
191
192
     def ignore_file(vcf_file):
193
         if "vcf" not in vcf_file or "truth" in vcf_file:
194
             return True
195
         return False
196
     # this method creates the set of keys for the dictionary
198
199
     def create_dictionary_keys(vcf_reader, sample_dictionary):
200
         for record in vcf_reader:
201
             if "GL" in str(record.CHROM):
202
203
                  continue
             sample_name = get_sample_name_from_record(record)
204
205
             sample_dictionary[sample_name] = [[], []] # fullname has become a key in fulldictionary
         return sample_dictionary
206
207
     # standard method that returns a tuple of the variant call object with the chromosome, position, reference
         and tuple of alternates
209
     def get_sample_name_from_record(record):
210
         templist = []
211
         for item in record.ALT:
             templist.append(str(item).upper())
213
         sample_name = (str(record.CHROM), str(record.POS), str(record.REF).upper(), tuple(templist))
214
         return sample_name
215
216
     # this method sets the length of the input neural networks
217
218
     def add_length_to_caller_lengths_based_on_file_name(vcf_file, caller_length, callerlengths):
219
         if "fb" in vcf_file:
220
             callerlengths[0] = caller_length
221
         if "hc" in vcf_file:
222
             callerlengths[1] = caller_length
223
         if "ug" in vcf_file:
224
             callerlengths[2] = caller_length
225
         if "pind" in vcf_file:
226
             callerlengths[3] = caller_length
227
         if "st" in vcf_file:
             callerlengths[4] = caller_length
229
```

```
230
     # this method wraps the create truth dictionary method and is used to checking that the dictionary file has
231
         the correct name
232
     def generate_truth_list(path):
233
         generated_truth_dictionary = {}
         for truth_file in os.listdir(path):
235
             if "truth" not in truth_file:
236
                  continue
237
             create_truth_dictionary(generated_truth_dictionary, truth_file)
238
         return generated_truth_dictionary
240
     # this method takes in the mutation (in a tuple) and checks if that mutation exists in the truth dictionary
241
     # A mutation exists if the chromosome, reference and position of the variant call is correct, AND one of
      \hookrightarrow the alternate alleles it contains
     # is also an alternate allele in the truth dataset
243
244
     def check_sample_against_truth_dictionary(tuple_name, final_truth_list, truth_dictionary):
245
246
         temp_tuple = (tuple_name[0], tuple_name[1], tuple_name[2])
         if temp_tuple in truth_dictionary:
247
             for alternate in tuple_name[3]:
248
                  if alternate in truth_dictionary[temp_tuple]:
249
                      final_truth_list.append(1)
250
                      return
251
         final_truth_list.append(0)
252
         return
253
254
     # This method loads the paths of the files into local variables
255
256
     def load_references(user_input):
257
         file1 = user_input['input'][0]
258
         referencepath = user_input['reference']
259
         output_location = user_input['output']
260
         return file1, referencepath, output_location
261
262
     # This method saves all the processed data into files that can be used for other purposes later or loaded
263
         natively instead of doing the processing again
264
     def save_files(output_location, x_array, length_of_caller_outputs, sample_list, truth_dict,
265
          vcf_dictionary_file,
                     y_array=[]):
266
         file2 = output_location
267
         x_data_file_name = str(file2) + str(X_DATA_NAME)
         np.save(x_data_file_name, x_array)
269
```

```
270
         vcf_file_name = str(file2) + str(VCF_LIST_FILE_NAME)
         caller_length_file_name = str(file2) + str(CALLER_LENGTH_FILE_NAME)
271
         truth_dictionary_name = str(file2) + str(TRUTH_DICTIONARY_NAME)
         list_of_inputs_name = str(file2) + str(LIST_OF_INPUTS_NAME)
273
         np.save(caller_length_file_name, length_of_caller_outputs)
274
         with open(list_of_inputs_name, 'wb') as samplesave1:
275
             pickle.dump(sample_list, samplesave1)
276
         with open(truth_dictionary_name, 'wb') as samplesave2:
             pickle.dump(truth_dict, samplesave2)
278
         with open(vcf_file_name, 'wb') as samplesave3:
279
             pickle.dump(vcf_dictionary_file, samplesave3)
         if y_array != []:
281
             y_data_file_name = str(file2) + str(Y_DATA_NAME)
282
             np.save(y_data_file_name, y_array)
284
     # This method takes in two dictionaries, a dictionary of truth mutations and a dictionary of sample
285
      → mutations,
     # checks whether each of the sample variables are inside the truth dictionary
286
     # and returns 2 arrays, an array of samples and an array of accompanying truth labels
287
288
     def check_predicted_with_truth(passed_list_of_samples, dictionary_of_truth=[]):
289
         final_array_of_samples = []
290
         final_truth_list = []
291
         for item in passed_list_of_samples:
292
             if dictionary_of_truth:
293
                 check_sample_against_truth_dictionary(item[0], final_truth_list, dictionary_of_truth)
294
             temp_array = []
             for row in item[1]:
296
                 temp_array.extend(row)
297
             final_array_of_samples.append(temp_array)
298
         if dictionary_of_truth:
299
             return final_truth_list, final_array_of_samples
300
         return final_array_of_samples
301
302
     # This method ensures that only the variables that have the modal number of features are used
303
     # in neural network training to ensure all array sizes are the same
304
305
     def add_mode_values_into_list_of_samples(full_dictionary, mode_value):
306
         list_of_passed_samples = []
307
         vcf_list = []
308
         for key in full_dictionary:
309
             second_count = 0
310
             for item in full_dictionary[key][0]:
                 second_count += len(item)
312
```

```
313
              if second_count != mode_value:
                  continue
314
              list_of_passed_samples.append([key, full_dictionary[key][0]])
315
              vcf_list.append(full_dictionary[key][1])
316
         return list_of_passed_samples, vcf_list
317
     # This method gets the modal number of features from a modal dictionary
319
320
     def get_mode_value(removaldict):
321
         curr = 0
322
         mode_value = 0
         for new_key in removaldict:
324
              if removaldict[new_key] > curr:
325
                  curr = removaldict[new_key]
326
                  mode_value = new_key
327
         return mode_value
328
329
     # This method iterates through the dataset to create a modal dictionary which contains a key-value pair of
330
      \hookrightarrow (number of features - number of times seen).
     # The mode number of features is kept
331
332
333
     def iterate_through_dictionary_to_find_mode_size(full_dictionary):
         removaldict = {}
334
         samples = 0
335
         for key in full_dictionary:
336
              samples += 1
337
              if samples == sample_limit:
338
                  break
339
              count = 0
340
             for item in full_dictionary[key]:
                  count += len(item)
342
              if count not in removaldict:
343
                  removaldict[count] = 1
344
345
                  removaldict[count] += 1
346
         return removaldict
347
348
     if __name__ == "__main__":
350
         np.seterr(divide='raise', invalid='raise')
351
         parser = argparse.ArgumentParser(description="train neural net")
352
         parser.add_argument('-i', '--input', help="give directories with files", nargs='+')
353
         parser.add_argument('-d', '--debug', help="look at matrixes built")
354
         parser.add_argument('-r', '--reference', help="")
355
```

```
parser.add_argument('-o', '--output', help="")

paths = parser.parse_args()

start = time.time()

load_and_save_data(paths)
```

7.2 train_network.py

33

```
#This script is called by the generate_matrixes.py script and contains the implementation of the neural
     \hookrightarrow network.
    #Input : np.arrays of features from generate_matrixes with accompanying truth labels, feature set lengths,
     \hookrightarrow a dictionary of vcf object records, as well as the list of sample features
    #Output : A VCF file containing all the filtered entries by the neural network, as well the list of

    accompanying scores

    #Overall Strategy :
    #Perform SMOTE oversampling of the input features, and then use the features to train the neural network
    #After training, perform validation on test dataset, and subsequently prepare a vcf file with filtered
     \hookrightarrow entries
     #import all necessary components
    import argparse
    import cPickle as pickle
11
    import sys
12
    import numpy as np
    import vcf
14
    from imblearn.over_sampling import SMOTE
    from keras.callbacks import *
16
    from keras.layers import Dense, Dropout, Activation
17
    from keras.layers.advanced_activations import LeakyReLU
    from keras.layers.normalization import BatchNormalization
19
20
    from keras.models import Sequential
    from keras.models import load_model
21
    from keras.optimizers import RMSprop
22
    from sklearn.metrics import *
23
    from sklearn.model_selection import train_test_split
25
    #set constants
27
    PCA_COMPONENTS = 8
28
    STEP_INCREMENT = 10
    RECURSION_LIMIT = 0.0002
30
    VERBOSE = 1
    seed = 1337
32
```

```
# Initialise random seed for reproducibility
    np.random.seed(seed)
35
36
    #Prepare file names for saving
37
    vcf_file_name = "/ANN/truevcf.vcf"
38
    keras_model_name = "/ANN/model"
    model truth name = "/ANN/modeltruths.txt"
40
    model_predictions_name = "/ANN/modelpredictions.txt"
41
    original_vcf_reader = "/data/backup/metacaller/stage/data/version6.3a/hc.vcf.normalisedtrain.vcf"
43
    # this method takes in a path and returns training matrixes for the ANN
    # The path should contain n caller vcf files and 1 truth file
    # vcf files should be labelled with vcf and truth file should be labelled with truth
46
    # no other file should be present in the folder
    def main_gather_input_execute_prep_output(array_sizes, dict_of_truth_input, fullmatrix_sample,
     calculated_prediction_actual, calculated_truth_actual = train_neural_net(20, 10, fullmatrix_sample,
49
        \hookrightarrow \quad \texttt{save\_location, array\_sizes)}
       get_all_relevant_scores(calculated_prediction_actual, calculated_truth_actual, dict_of_truth_input,
50
        → list_of_samples_input, vcf_dictionary, save_location)
    # This method counts the number of false negatives inside the input sample
52
53
    def count_false_negative(calculated_prediction_actual, calculated_truth_actual):
        count_false_negative = 0
55
        for i in range(len(calculated_prediction_actual)):
56
           if calculated_prediction_actual[i] == 0 and calculated_truth_actual[i] == 1:
57
               count_false_negative += 1
58
       return count_false_negative
60
    # this is the wrapper function for the recursive hill climbing algorithm to get the best f1 score
61
    # It starts from a low threshold value, and marginally increases the threshold until it is unable to find
62
    # any better F1 scores. It then reports the threshold, F1 score and produces the filtered callset
63
64
    def get_all_relevant_scores(calculated_prediction_actual, calculated_truth_actual, dict_of_truth_input,
65
                              list_of_samples_input, vcf_list, outputpath):
66
       print "Here are some predictions", calculated_prediction_actual[:100]
       print "here are some truths", calculated_prediction_actual[:100]
68
       f1_score_left = get_scores(calculated_prediction_actual, calculated_truth_actual, 0.0,
69
        → list_of_samples_input, dict_of_truth_input)
       guess_f1_final_score, guess_f1_final = recursive_best_f1_score(calculated_prediction_actual,
70
```

```
get_scores(calculated_prediction_actual, calculated_truth_actual, guess_f1_final,
         produce_vcf_file(calculated_prediction_actual, guess_f1_final, list_of_samples_input, vcf_list,

→ outputpath)

73
     # This method produces the vcf file through filtering with the neural network threshold calls
75
     def produce_vcf_file(calculated_prediction_actual, guess_f1_final, list_of_samples_input, vcf_list,
76

→ outputpath):

        prediction = []
77
         for item in calculated_prediction_actual:
            if item > guess_f1_final:
                prediction.append(1)
80
            else:
                prediction.append(0)
82
        list_of_records = []
83
         for i in range(len(list_of_samples_input)):
            if prediction[i] == 1:
85
                 list_of_records.append(vcf_list[i])
        vcf_reader = vcf.Reader(filename=original_vcf_reader)
        vcf_writer = vcf.Writer(open(outputpath + vcf_file_name, 'w'), vcf_reader)
         for record in list_of_records:
            vcf writer write record(record)
90
91
     # This method is the recursive function that attempts to find the threshold that produces the best f1
     # by iterating through steps of thresholds (0.2, 0.02 and 0.002) until no better F1 score can be found for

→ a marginal increase in threshold.

     # It then returns the best F1 score and the threshold
94
     def recursive_best_f1_score(calculated_prediction_actual, calculated_truth_actual, dict_of_truth_input,
96
                                list_of_samples_input, guess, guess_score, step):
97
         if step <= RECURSION_LIMIT:</pre>
98
            return guess_score, guess
99
        new_guess = guess + step
100
        new_guess_score = get_scores(calculated_prediction_actual, calculated_truth_actual, new_guess,
101
                                     list_of_samples_input, dict_of_truth_input)
102
         if new_guess_score > guess_score:
103
            return recursive_best_f1_score(calculated_prediction_actual, calculated_truth_actual,
104

    dict_of_truth_input,
105
                                           list_of_samples_input, new_guess, new_guess_score, step)
        return recursive_best_f1_score(calculated_prediction_actual, calculated_truth_actual,
106
            dict_of_truth_input,
                                       list_of_samples_input, guess, guess_score, step / STEP_INCREMENT)
107
```

```
108
     # this method uses pre-loaded data to train the neural network. It is optional and only used when this
109
         python script is called natively and not imported
110
     def load_references(input_paths):
111
112
         input_paths = vars(input_paths)
         fullmatrix_sample = np.load(input_paths['input'][0])
113
         fullmatrix_truth = np.load(input_paths['input'][1])
114
         with open(input_paths['input'][3], 'rb') as fp1:
115
             list_of_samples_input = pickle.load(fp1)
116
         with open(input_paths['input'][4], 'rb') as fp2:
             dict_of_truth_input = pickle.load(fp2)
118
         array_sizes = np.load(input_paths['input'][5])
119
         with open(input_paths['input'][6], 'rb') as fp3:
             vcf_dictionary = pickle.load(fp3)
121
         orig_stdout = sys.stdout
122
123
         f = file(str(input_paths['input'][3]) + '.txt', 'w')
         sys.stdout = f
124
125
         return array_sizes, dict_of_truth_input, fullmatrix_sample, fullmatrix_truth, list_of_samples_input,
             input_paths, vcf_dictionary
126
     # this method solves the double false negative problem that is created due to the neural network prediction
         scheme
128
     def remove_duplicated_false_negative(prediction_list, truth_list, false_negatives):
129
         count = 0
130
         removal_list = []
131
         for i in range(len(prediction_list) - 1, -1, -1):
132
             if count == false_negatives:
133
                 break
             if prediction_list[i] == 0 and truth_list[i] == 1:
135
                 removal_list.insert(0, i)
136
                 count += 1
137
         for index in removal_list:
138
             prediction_list.pop(index)
139
140
             truth_list.pop(index)
         return prediction_list, truth_list
141
     # this method takes in the binary truth and predicted samples and calculates the true positive rate, false
143
         positive rate, recall, precision and f1 score
144
     def get_scores(actual_predictions, actual_truth, value, sample_list, truth_dictionary, verbose=0):
145
         temp_actual_truth = list(actual_truth)
146
         prediction = []
147
```

```
for item in actual_predictions:
148
             if item > value:
149
                 prediction.append(1)
150
             else:
151
                 prediction.append(0)
152
         false_negatives = count_false_negative(actual_predictions, actual_truth)
153
         finalpredictionnumbers, finaltruthnumbers = add_negative_data(sample_list, truth_dictionary,
154
              prediction, temp_actual_truth)
         finalpredictionnumbers, finaltruthnumbers = remove_duplicated_false_negative(finalpredictionnumbers,
155
          \hookrightarrow finaltruthnumbers, false_negatives)
         final_f1_score = f1_score(finaltruthnumbers, finalpredictionnumbers)
         if verbose:
157
             print_scores(actual_truth, final_f1_score, finalpredictionnumbers, finaltruthnumbers, prediction,
158
              \hookrightarrow value)
         return final_f1_score
159
160
     # default method for printing all relevant scores
161
162
     def print_scores(actual_truth, final_f1_score, finalpredictionnumbers, finaltruthnumbers, prediction,
163

    value):

         final_false_positive, final_true_negative = perf_measure(finaltruthnumbers, finalpredictionnumbers)
164
         print "final false positive rate is :", final_false_positive
165
         print "final true negative rate is :", final_true_negative
166
         print "final precision score is :", precision_score(finaltruthnumbers, finalpredictionnumbers)
167
         print "final recall score is :", recall_score(finaltruthnumbers, finalpredictionnumbers)
         print "threshold is", value
169
         print "final F1 score is : ", final_f1_score
170
171
     # This method looks at the set of predicted samples and the set of truths and adds the false negatives to
172
         the predicted sample.
173
     def add_negative_data(list_of_samples, dict_of_truth, array_of_predicted, array_of_truth):
174
         dict_of_samples = generate_sample_dictionary(array_of_predicted, list_of_samples)
175
         list_of_truth = generate_list_of_truth(dict_of_truth)
176
         new_array_of_predicted = list(array_of_predicted)
         new_array_of_truth = list(array_of_truth)
178
         original_length = len(new_array_of_predicted)
179
         for item in list_of_truth:
             fillnegative(item, dict_of_samples, new_array_of_predicted, new_array_of_truth)
181
         print "number of false data samples are", (len(new_array_of_predicted) - original_length)
182
         return new_array_of_predicted, new_array_of_truth
183
184
     # This method generates a list of truth variant calls from a dictionary of truth variant calls.
```

186

```
def generate_list_of_truth(dict_of_truth):
         list_of_truth = []
188
         for key in dict_of_truth:
189
              mytuple = dict_of_truth[key]
190
             temptuple = []
191
             for item in mytuple:
192
                  temptuple.append(item)
193
              list_of_truth.append([key[0], key[1], key[2], temptuple])
194
         return list_of_truth
195
196
     # This method generates a dictionary of sample variant calls from a list of sample variant calls.
198
     def generate_sample_dictionary(array_of_predicted, list_of_samples):
199
         dict_of_samples = {}
200
         for i in range(len(list_of_samples)):
201
              item = list_of_samples[i]
202
203
              if array_of_predicted[i] == 0:
                  continue
204
205
              new_key = (item[0][0], item[0][1], item[0][2])
              new_value = item[0][3]
206
              if new_key not in dict_of_samples:
207
                  dict_of_samples[new_key] = new_value
              else:
209
                  dict_of_samples[new_key] = list(dict_of_samples[new_key])
210
                  dict_of_samples[new_key].extend(new_value)
211
                  dict_of_samples[new_key] = tuple(dict_of_samples[new_key])
212
                  # print dict_of_samples[new_key]
213
         return dict_of_samples
214
215
     # Actual method to calculated false positive, false negative rates
216
217
     def perf_measure(y_actual, y_hat):
218
         true_positive = 0
219
         false_positive = 0
220
         false_negative = 0
221
         true_negative = 0
222
223
         for i in range(len(y_hat)):
224
              if y_actual[i] == 1 and y_hat[i] == 1:
225
                  true_positive += 1
226
         for i in range(len(y_hat)):
227
              if y_hat[i] == 1 and y_actual[i] == 0:
228
                  false_positive += 1
         for i in range(len(y_hat)):
230
```

```
231
             if y_actual[i] == 1 and y_hat[i] == 0:
                 false_negative += 1
232
         for i in range(len(y_hat)):
233
             if y_hat[i] == 0 and y_actual[i] == 0:
234
                 true_negative += 1
235
236
         print "true positives :", true_positive
237
         print "false positives :", false_positive
238
         print "false negatives :", false_negative
239
         print "true negatives :", true_negative
240
         true_positive = float(true_positive)
242
         false_positive = float(false_positive)
243
         false_negative = float(false_negative)
         if false_positive == 0 and true_positive == 0:
245
             false_positive_rate = 0
246
247
         else:
             false_positive_rate = false_positive / (false_positive + true_positive)
248
249
         if false_negative == 0 and true_positive == 0:
             true_negative_rate = 0
250
         else:
251
             true_negative_rate = false_negative / (false_negative + true_positive)
252
253
         return false_positive_rate, true_negative_rate
254
255
256
     # comparator method that takes a tuple and checks whether it is in the dictionary of samples, if it is not,
257
         then add a false negative call to the dataset
258
     def fillnegative(tuple1, sampledict, arrayofsamples, arrayoftruths):
259
         tuple2 = (tuple1[0], tuple1[1], tuple1[2])
260
         if tuple2 in sampledict:
261
             for ALT in tuple1[3]:
262
                 if ALT in sampledict[tuple2]:
263
                      return
264
         arrayofsamples.append(0)
265
         arrayoftruths.append(1)
266
     # main method that performs neural network training. This method takes in the sample matrixes, the truth
268
      → variables, a save file location, number of epochs,
     # size of input arrays and the minibatch training size. It first performs SMOTE on the input dataset, then
      → splits it into training and test dataset. It then
     # initialises the deep learning layers, compiles the neural network and uses the input data to fit the
      → network. The best set of weights at any point is saved
```

```
# to a file and reloaded at the end of the fitting. After training, the neural network is used to predict
      \hookrightarrow the original un-oversampled dataset
     def train_neural_net(mybatch_size, mynb_epoch, myX_train, myy_train, location, arraysize):
273
         fb_size, hc_size, ug_size, pindel_size, st_size = get_sizes(array_sizes)
274
         X_resampled, y_resampled = do_smote_resampling(myX_train, myy_train)
275
         X_train, X_test, y_train, y_test = train_test_split(X_resampled, y_resampled,
276
                                                               test_size=0.33, random_state=seed)
         X_fb, X_hc, X_ug, X_pindel, X_st = prep_input_samples(array_sizes, X_train)
278
         X_fb_test, X_hc_test, X_ug_test, X_pindel_test, X_st_test = prep_input_samples(array_sizes, X_test)
279
         batch_size = mybatch_size
         nb_epoch = mynb_epoch
281
282
         fb_branch = Sequential()
         develop_first_layer_matrixes(fb_branch, fb_size)
284
285
         hc_branch = Sequential()
286
         develop_first_layer_matrixes(hc_branch, hc_size)
287
288
289
         ug_branch = Sequential()
         develop_first_layer_matrixes(ug_branch, ug_size)
290
291
         pindel_branch = Sequential()
292
         develop_first_layer_matrixes(pindel_branch, pindel_size)
293
294
         st_branch = Sequential()
295
         develop_first_layer_matrixes(st_branch, st_size)
296
297
         final_model = Sequential()
298
         final_model.add(Merge([fb_branch, hc_branch, ug_branch, pindel_branch, st_branch], mode='concat',
299
          final_model.add(Dense(24, activation='linear'))
300
         final_model.add(LeakyReLU(alpha=0.05))
301
         final_model.add(Dense(6, activation='linear'))
302
         final_model.add(LeakyReLU(alpha=0.05))
303
         final_model.add(Dense(1, activation='linear'))
304
         final_model.add(Activation('sigmoid'))
305
         print (final_model.summary())
         adam = Adam(lr=0.00001, rho=0.9, epsilon=1e-08, decay=0.0)
307
         final_model.compile(loss='binary_crossentropy',
308
309
                              optimizer=adam,
                              metrics=['accuracy'])
310
311
         filepath = location + "/best_weights.hdf5"
312
```

```
313
         checkpoint = ModelCheckpoint(filepath, monitor='val_acc', verbose=1, save_best_only=True, mode='max')
         callbacks_list = [checkpoint]
314
         model_history = final_model.fit([X_train], y_train, batch_size=batch_size, nb_epoch=nb_epoch,
315
                                          validation_split=0.2, verbose=2, callbacks=callbacks_list)
316
         final_model = load_model(location + "/best_weights.hdf5")
317
         print model_history.history['val_acc'], model_history.history['val_acc']
         print model_history.history['val_loss'], model_history.history['val_loss']
319
         np.save(location + "/best_weights.hdf5", model_history.history['val_acc'])
320
         np.save(location + "/best_weights.hdf5", model_history.history['val_loss'])
321
         scores = final_model.evaluate([X_test], y_test)
322
         print scores
         final_prediction_array_probabilities = final_model.predict([myX_train])
324
         final_prediction_array_probabilities = np.squeeze(final_prediction_array_probabilities)
325
         save_model_details(final_model, final_prediction_array_probabilities, myy_train, location)
326
327
         return final_prediction_array_probabilities, myy_train
328
329
     # Method to perform SMOTE oversampling
330
331
     def do_smote_resampling(myX_train, myy_train):
332
         sm = SMOTE(kind='regular')
333
         where_are_NaNs = np.isnan(myX_train)
334
         myX_train[where_are_NaNs] = 0
335
         X_resampled, y_resampled = sm.fit_sample(myX_train, myy_train)
336
         return X_resampled, y_resampled
338
     # this method saves the details of the neural network
339
340
     def save_model_details(final_model, save_model_probabilities, trutharray, location):
341
         name1 = location + model_predictions_name
342
         name2 = location + model_truth_name
343
         name3 = location + keras_model_name
344
         np.save(name1, save_model_probabilities)
345
         np.save(name2, trutharray)
346
         final_model.save(name3)
347
348
     # this method gets the array size of the features used
349
     def get_sizes(array_sizes):
351
         fb_size = array_sizes[0]
352
         hc_size = array_sizes[1]
353
         ug_size = array_sizes[2]
354
         pindel_size = array_sizes[3]
         st_size = array_sizes[4]
356
```

```
return fb_size + hc_size + ug_size + pindel_size + st_size
357
358
359
     # this method uses a map function to filter data such that each merge layer gets the correct set of data
360
361
     def prep_input_samples(array_sizes, x_training_data):
362
363
         X_fb = np.array(map(lambda x: x[count:array_sizes[0]], x_training_data))
364
         count += array_sizes[0]
365
         X_hc = np.array(map(lambda x: x[count:count + array_sizes[1]], x_training_data))
366
         count += array_sizes[1]
         X_ug = np.array(map(lambda x: x[count:count + array_sizes[2]], x_training_data))
368
         count += array_sizes[2]
369
         X_pindel = np.array(map(lambda x: x[count:count + array_sizes[3]], x_training_data))
         count += array_sizes[3]
371
         X_st = np.array(map(lambda x: x[count:count + array_sizes[4]], x_training_data))
372
         count += array_sizes[4]
373
         return X_fb, X_hc, X_ug, X_pindel, X_st
374
375
376
     if __name__ == "__main__":
377
         parser = argparse.ArgumentParser(description="train neural net")
378
         parser.add_argument('-i', '--input', help="give directories with files", nargs='+')
379
         input_path = parser.parse_args()
380
         array_sizes, dict_of_truth_input, fullmatrix_sample, fullmatrix_truth, \
381
         list_of_samples_input, paths, vcf_dictionary = load_references(input_path)
382
         main_gather_input_execute_prep_output(array_sizes, dict_of_truth_input, fullmatrix_sample,

→ fullmatrix_truth, list_of_samples_input, paths, vcf_dictionary)

     7.3
            compute_bayesian.py
```

```
#This script takes in a VCF file with functional annotation already done, and computes the bayesian network

→ using the annotations. It produces a sorted list of vcf entries in a text file, with accompanying

→ annotation scores

#Input: VCF file with functional annotation

#Output: A sorted list of vcf entries with accompanying annotation scores, redirected from stdout to a

→ file

#Overall Strategy:

#First extract all the features from the vcf files and then perform feature-wise normalisation.

#Subsequently, prepare the bayesian network by creating edges, nodes, preparing prior distritions

#Finally use features to update the bayesian network to obtain final probabilities for importance

#Report a list of sorted probabilites for easy ranking

import matplotlib

import vcf
```

```
12
    matplotlib.use('Agg')
13
14
    from pomegranate import *
15
16
    #main method for loading references into local variables
18
    def load_reference(paths):
19
        paths = vars(paths)
20
         input = paths['input']
21
         opened_vcf_file = vcf.Reader(open(input, 'r'))
        name3 = input + "finalscores.txt"
23
         # orig_stdout = sys.stdout
24
         # f = file(name3 + '.txt', 'w')
         # sys.stdout = f
26
        return opened_vcf_file
27
     #method for getting functional annotation scores
29
30
    def get_scores(record):
31
         list_of_important_mutations = [record.INFO['SIFT_score'], record.INFO['LRT_score'],
32
                                         record.INFO['MutationAssessor_score'],
33
                                         record.INFO['Polyphen2_HVAR_score'], record.INFO['FATHMM_score']]
34
         if 'NN_prediction' in record.INFO:
35
             NN_prediction = record.INFO['NN_prediction'][0]
        else:
37
             NN_prediction = -1
        list_of_important_mutations = map(lambda x: x[0], list_of_important_mutations)
39
        list_of_important_mutations = map(lambda x: None if x == None else float(x),
40
         \hookrightarrow list_of_important_mutations)
        return NN_prediction, list_of_important_mutations
41
    \#main method that controls I/O - it gets the input, applies the main function and then prepares the output
43
44
    def main(paths):
45
        vcf_object = load_reference(paths)
46
        full_list_of_scores = analyse_main(vcf_object)
47
        prepare_output(full_list_of_scores)
49
    #this method controls the processes applied to the vcf file - for each record, it extract the list of

→ scores.

    # normalises it, compute probabilities, sorts it and then return output
51
    def analyse_main(vcf_object):
```

```
full_list_of_scores = extract_list_of_scores(vcf_object)
        apply_feature_wise_normalisation(full_list_of_scores)
55
        compute_network_and_probabilities(full_list_of_scores)
56
        full_list_of_scores.sort(key=lambda x: x[4], reverse=True)
57
        return full_list_of_scores
58
    # since print is redirected to stdoutput, print function is used to store output
60
61
    def prepare_output(full_list_of_scores):
62
        for item in full_list_of_scores:
63
            print item[2], item, item[2].INFO['Gene.refGene']
64
65
    # wrapper function used to create bayesian network for all records
66
    def compute_network_and_probabilities(full_list_of_scores):
68
        for record in full_list_of_scores:
69
            network = create_network_and_compute_probabilities(record)
70
            compute_record(network, record)
71
    # this function applies a featurewise normalisation of all features to a range of 0-1, and flip scores
73
    # for certain features
74
75
    def apply_feature_wise_normalisation(full_list_of_scores):
76
        for i in range(6):
            min_num = 1000000
            max_num = -1000000
79
            for item in full_list_of_scores:
                 if item[1][i] != None:
81
                     min_num = min(min_num, item[1][i])
82
                     max_num = max(max_num, item[1][i])
            for item in full_list_of_scores:
84
                 if item[1][i] != None:
                     value = ((item[1][i] - min_num) / (max_num - min_num) + 0.2) / 1.3
86
                     item[1][i] = value
87
                 else:
                    item[1][i] = 0.5
89
            if i == 0 or i == 5:
90
                 for item in full_list_of_scores:
                     if item[1][i] != None:
92
                         item[1][i] = -item[1][i]
93
    # extract list of of scores from each record, including all functional annotations, clinvar scores and
95
        dbsnp
```

96

```
def extract_list_of_scores(vcf_object):
         count = 0
98
         full_list_of_scores = []
99
         for record in vcf_object:
100
             count += 1
101
             nn_prediction, list_of_scores = get_scores(record)
102
             if not list(filter(lambda x: x != None, list_of_scores)):
103
                  continue
104
             get_clinvar_scores(list_of_scores, record)
105
             snp_present = get_db_snp_scores(record)
106
             full_list_of_scores.append([float(nn_prediction), list_of_scores, record, snp_present])
         return full_list_of_scores
108
109
     # Compute the Bayesian Network by assuming observations and attaching mapped probabilities (0,1) to
      \rightarrow P(X=True \mid Y=True)
111
112
     def compute_record(network, record):
         beliefs = network.predict_proba({'Real Gene': 'True', 'ClinVar': 'True', 'PolyPhen': 'True', 'LRT':
113
          → 'True', 'MutationAssessor': 'True', 'SIFT': 'True', 'FATHMM_gene': 'True', 'rs_gene': 'True'})
         # print "\n".join("{}\t{}".format(state.name, belief) for state, belief in zip(network.states,
114
          \hookrightarrow beliefs))
115
         # get the probability that the gene is important
         prob_gene_important = beliefs[2].values()[1]
116
         beliefs = map(str, beliefs)
117
         record.append(prob_gene_important)
118
         record.append(record[2].INFO['snp138'])
119
         record.append(record[3])
121
     # If snp is present in db-snp, attach probability of importance to 0.3, else 0.7
122
123
     def get_db_snp_scores(record):
124
         snp_present = 0.7
125
         if record.INFO['snp138'][0] != None:
126
             snp_present = 0.3
127
         return snp_present
128
129
     # If snp is present in clinvar, attach probability of importance to 0.7, else 0.3
130
131
     def get_clinvar_scores(list_of_scores, record):
132
         if record.INFO['clinvar_20150629'][0] != None:
133
             list_of_scores.append(0.7)
134
         else:
135
             list_of_scores.append(0.3)
137
```

```
# wrapper method to create the bayesian network and compute probabilities
139
     def create_network_and_compute_probabilities(record):
140
         ClinVar_gene, FATHMM_gene, LRT_gene, MutationAssessor_gene, MutationTaster_gene, PolyPhen2_gene,
141

→ SIFT_gene, functional_gene, importgene, real_gene, rs_gene = initialise_distributions()

142
             record)
         # set up states
143
         s1, s10, s11, s2, s3, s4, s5, s6, s7, s8, s9 = generate_states(ClinVar_gene, FATHMM_gene, LRT_gene,
144
          → MutationAssessor_gene, MutationTaster_gene, PolyPhen2_gene, SIFT_gene, functional_gene,
             importgene, real_gene, rs_gene)
         # set up network
         network = add_edges_bake_network(s1, s10, s11, s2, s3, s4, s5, s6, s7, s8, s9)
146
         return network
147
     # method to create the edges in the network
149
150
     def add_edges_bake_network(s1, s10, s11, s2, s3, s4, s5, s6, s7, s8, s9):
151
         network = BayesianNetwork("Gene Prediction")
152
153
         network.add_states(s1, s2, s3, s4, s5, s6, s8, s9, s10, s11)
         network.add_edge(s1, s3)
154
         network.add_edge(s2, s3)
155
         network.add_edge(s4, s2)
156
         network.add_edge(s5, s2)
157
         network.add_edge(s6, s2)
158
         network.add_edge(s7, s2)
159
         network.add_edge(s8, s2)
160
         network.add_edge(s9, s2)
         network.add_edge(s10, s2)
162
         network.add_edge(s11, s3)
163
         network.bake()
         return network
165
166
167
     # method that generates the nodes in the bayesian network
168
     def generate_states(ClinVar_gene, FATHMM_gene, LRT_gene, MutationAssessor_gene, MutationTaster_gene,
169
         PolyPhen2_gene,
                         SIFT_gene, functional_gene, importgene, real_gene, rs_gene):
170
171
         s1 = State(real_gene, name="Real Gene")
         s2 = State(functional_gene, name="Functional Gene")
172
         s3 = State(importgene, name="Important Gene")
         s4 = State(ClinVar_gene, name="ClinVar")
174
         s5 = State(PolyPhen2_gene, name="PolyPhen")
175
         s6 = State(LRT_gene, name="LRT")
         s7 = State(MutationTaster_gene, name="MutationTaster")
177
```

```
178
         s8 = State(MutationAssessor_gene, name="MutationAssessor")
         s9 = State(SIFT_gene, name="SIFT")
179
         s10 = State(FATHMM_gene, name="FATHMM_gene")
180
         s11 = State(rs_gene, name="rs_gene")
181
         return s1, s10, s11, s2, s3, s4, s5, s6, s7, s8, s9
182
183
     #methods to initialise prior distributions in bayesian network
184
185
     def initialise_distributions(record):
186
         ClinVar_gene = DiscreteDistribution({'True': 0.5, 'False': 0.5})
187
         PolyPhen2_gene = DiscreteDistribution({'True': 0.5, 'False': 0.5})
188
         LRT_gene = DiscreteDistribution({'True': 0.5, 'False': 0.5})
189
         MutationTaster_gene = DiscreteDistribution({'True': 0.5, 'False': 0.5})
190
         MutationAssessor_gene = DiscreteDistribution({'True': 0.5, 'False': 0.5})
         SIFT_gene = DiscreteDistribution({'True': 0.5, 'False': 0.5})
192
         FATHMM_gene = DiscreteDistribution({'True': 0.5, 'False': 0.5})
193
         rs_gene = DiscreteDistribution({'True': 0.5, 'False': 0.5})
194
         import_cdp = get_cdp(3, [(record[0] + 0.2) / 1.3, record[3], 0.8])
195
196
         functional_cdp = get_cdp(6, record[1])
         functional_gene = ConditionalProbabilityTable(functional_cdp, [ClinVar_gene, PolyPhen2_gene, LRT_gene,
197
                                                                          MutationAssessor_gene,
198
                                                                          SIFT_gene, FATHMM_gene])
199
         real_gene = DiscreteDistribution({'True': 0.5, 'False': 0.5})
200
         importgene = ConditionalProbabilityTable(import_cdp, [real_gene, rs_gene, functional_gene])
201
         return ClinVar_gene, FATHMM_gene, LRT_gene, MutationAssessor_gene, MutationTaster_gene,
202
          → PolyPhen2_gene, SIFT_gene, functional_gene, importgene, real_gene, rs_gene
203
204
     # method that builds the cdp table. n is the number of input variables, probability list gives the
205

→ probability

     # that the i-th X variable is true P(Xi=True).
206
207
     def get_cdp(n, prob_list):
208
         temp_list = create_true_false_matrix(n)
209
         calculate_probabilities(n, prob_list, temp_list)
210
         return temp_list
211
212
213
     # Generates a True False matrix using binary counting logic, critical for input in bayesian network
214
215
     def create_true_false_matrix(n):
216
         temp_list = []
217
         for i in range(0, 2 ** n):
             temp_row = []
219
```

```
220
             for j in range(n):
                  number_2 = i // (2 ** (n - j - 1))
221
                  number_1 = number_2 \% 2
                  if number_1 == 0:
223
                      temp_row.append('False')
224
225
                  else:
                      temp_row.append('True')
226
              temp_list.insert(0, temp_row + ['False'])
227
              temp_list.insert(0, temp_row + ['True'])
228
         return temp_list
229
231
     # calculates the probabilities, taking in the true list as well as a list of probabilities. The key here is
232
     # the probability that the mutation is true is related to the scores given by mutation taster etc..
     # ie\ P(X\ is\ impt\ /\ X\ is\ Clinvar) = P(X\ is\ Clinvar)
234
235
236
     def calculate_probabilities(n, prob_list, temp_list):
         for i in range(0, 2 ** (n + 1), 2):
237
238
              true_row = temp_list[i]
             true_probability = 1
239
             false\_probability = 1
240
241
             for k in range(0, n, 1):
                  if true_row[k] == 'True':
242
                      true_probability *= prob_list[k]
243
                      false_probability *= 1 - prob_list[k] # probability that mutation is false is 1 minus
                       \hookrightarrow mutation is true
                  else:
                      true_probability *= 1 - prob_list[k]
246
                      false_probability *= prob_list[k]
247
              final_true_probability = true_probability / (true_probability + false_probability)
              final_false_probability = false_probability / (true_probability + false_probability)
249
              temp_list[i].append(final_true_probability)
250
              temp_list[i + 1].append(final_false_probability)
251
252
     if __name__ == "__main__":
253
         parser = argparse.ArgumentParser(description="train neural net")
254
         parser.add_argument('-i', '--input', help="give directories with files")
255
         paths = parser.parse_args()
         main(paths)
257
```