

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

**Chan Khai Ern, Edwin**

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# 1 Introduction

Variant calling is a critical step in the identification of mutations in genomes, and is important in downstream applications such as the annotation of mutant genes and the analysis of physiological consequences. However, current variant callers still tend to have low concordances for variants called (O’Rawe et al., 2013; Cornish and Guda, 2015), primarily due to differences in variant calling algorithms and assumptions. Furthermore, these variant callers do not take into account the importance of each variant in the Human metabolic and biochemical pathways. This is critical for clinicians as a clinician should be able to obtain variants that are of clinical significance, enabling them to embark on the best treatment pathway. In this paper, we describe a tool, INSERTNAMEHERE, to integrate data from multiple variant callers, filter true variants and prioritise their importance. This tool uses deep learning for filtering variants, and bayesian updating to determine variants that are the most important.

Variant calling primarily involves the use of various statistical and mathematical methods to discover variants, or mutations, in the genome. Calling variants allows the analysis of deviations and differences between the genome of interest and a standard human genome. However, 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; Gzsi et al., 2015). Both these problems fundamentally result 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. Probabilistic haplotype generating callers (such as GATK’s haplotype caller and FreeBayes) tend to be more accurate for SNPs and indels (McKenna et al. 2010; Garrison & Marth, 2012). They perform de-novo local assembly, where they rebuild small portions of the genome, and subsequently use bayesian analysis to determine the existence of variants. Specifically, they generate short haplotypes of local regions from sampled sequences, and determine (based on the haplotypes and prior probabilities in the reference genome) whether a variant should be called. However, these methods can only handle limited window sizes, preventing the detection of larger structural variants. For these we have to rely on other tools that ex-

amine larger segments of the genome (Ning et al., 2009) or use libraries of known mutation regions, and study these breakpoints to check if any mutations have occurred (Gerstein et al., 2015). Due to the heterogeneity in mutations, no single caller works best for all classes of mutations, pointing towards a variant calling framework that aggregates data from multiple callers.

Indeed, studies have shown low concordances between variant callers themselves, due to their specific implementations and algorithms (Mohiyuddin, et al., 2015; Gzsi et al., 2015). If we consider that each variant caller samples from the same genome but with a different statistical technique, then we can see each variant caller as a mode of data that 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 using multiple techniques.

The simplest approach to aggregate data is concordance - if multiple variant callers are able to call a variant, it is most likely to be accurate. However, the precision of such a tool would be poor due to the differential sensitivity of callers to edge cases. This would defeat the purpose of using multiple callers in the first place, as the strength of a combinatorial approach lies in tapping into the sensitivities of different callers. 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 (Gzsi 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 hypothesize that deep learning can also be used to integrate the information from variant callers.

Deep learning is a method of machine learning that involves deep stacks of artificial neural networks. 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 are able to predict outputs if given a

new input.

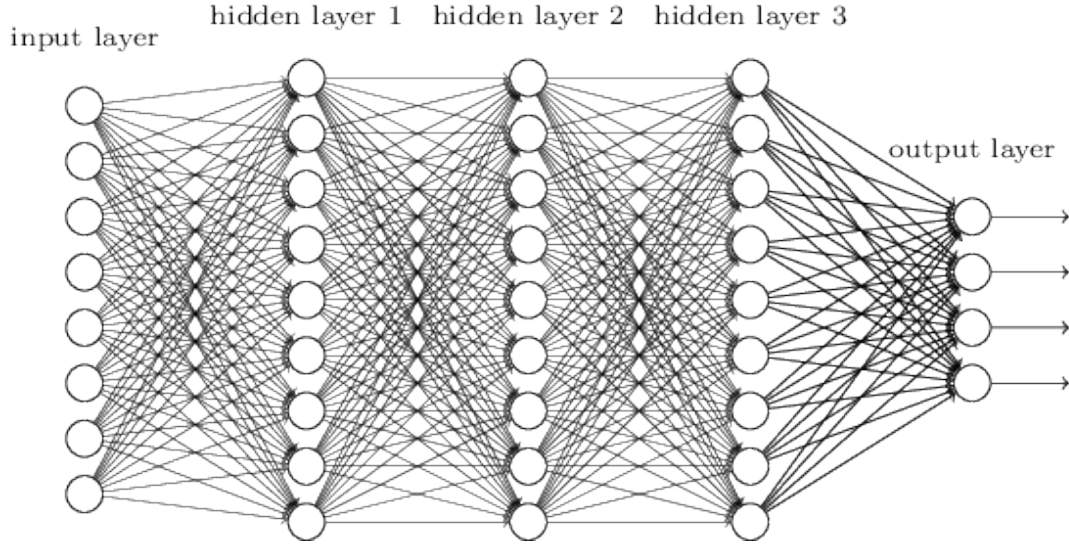


Figure 1: A Neural Network with 1 input layer, 3 hidden layers and 1 output layer. This represents a densely connected neural network, where each node is connected to every node of the preceding and subsequent layers. At each node, linking functions can be had

Figure 1 depicts a sample neural network with 5 layers, with 8 data points as input, and 3 data points as output. In such a network, we would train it 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. This in part mimics the way we learn - experience teaches us that certain stimuli will result in specific effects (when we see a lightning bolt, we can expect the sound of thunder), and thus when new input comes in (a lightning bolt is seen), we can predict that the output that would arise (thunder is heard). In variant calling, deep learning will allow us to predict based on variant calling patterns and data whether a variant is valid and exists, or is erroneous. This will allow us to draw of the diversity of data with different variant callers, through letting the network learn which patterns will result in a valid call and which patterns are actually 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, such a combinatorial approach will allow us to improve the accuracy and precision of variant calling.

$$P(MutationisImportant|MutationisclinVar) = \frac{P(MutationisclinVar|mutationisimportant) * P(Mutationisimp}{P(MutationisClinVar)} \quad (1)$$

$$\prod P(token_i|spam) = P("free"|spam) * P("viagra"|spam) * P(nexttoken|spam).... \quad (2)$$

$$P(clinVar|important) \quad (3)$$

$$P(mutationTaster|important) \quad (4)$$

$$P(SIFT|important) \quad (5)$$

## 2 Materials and Methods

### 2.1 Artificial Datasets

To generate datasets to train variant calling networks, VarSim was used with ART in order to generate artificial datasets. Datasets generated has

### 2.2 Variant Callers

Variant callers were chosen for our neural network based on their orthogonal calling and reference methodologies - we wanted to maximise the range of variant callers in order to optimise the information that the neural network receives (See Table N). All callers

REFERENCES :

Samtools : Li H, et al., 2009

GATK UG/HC : McKenna et al. 2010, DePristo et al. 2011

FB : Garrison & Marth, 2012

Table 1: Table Comparing Methods and Features of Different variant callers.

	<b>GATK Unified Genotyper</b>	<b>GATK Haplotype Caller</b>	<b>Free Bayes</b>	<b>Samtools</b>	<b>Pindel</b>
<b>Calling Method</b>	Uses a list of mapped reads, calling model is probabilistic with increased priors at regions with known SNPs	Uses Hidden Markov Models to build a likelihood of haplotypes which are then used to call variants	Uses haplotypes to represent mutations, calling model is probabilistic with population based priors	Uses a list of mapped reads, calling model is probabilistic model with increased priors at regions with known SNPs	Locates regions which were mapped with indels or only one end was mapped, and then performs a pattern growth to find inserts and deletions.
<b>Reference and Mapping Method</b>	Position based caller that realigns fragments and analyses each position to call SNPs and indels	Only analyses regions where there is high likelihood of mutation based on activity score, and builds a De Bruijn-like graph that reassembles reads in that region (Haplotypes)	Sliding window based caller, using reference matches instead of position	Position based caller that uses mapped sequences to call SNPs and indels	Focuses on Unmapped regions, regions known to have insert and deletions or regions with only one end mapped.
<b>Unique Features</b>	Looks at each position and determines if there is a variant or not based on a model. Tends to be aggressive when calling indels to due priors	Shown in literature to be one of the best variant callers for predicting variants and indels in real datasets (Sandmann et al., 2017; Hwang et al., 2015)	Does not require precise alignment, unlike other callers	Does not assume sequencing errors are independent unlike Unified Genotyper, and has less hard filters.	Known to be able to identify medium length indels due to pattern growth method

## 2.3 Feature Selection

## 2.4 Neural Networks

## 2.5 Technologies

For our deep learning networks, we used the Keras library with a TensorFlow backend. TensorFlow was chosen due to its superior performance on single machines with multiple cores. On our Ubuntu compute cluster, TensorFlow’s distributed CPU computation and queue management system enabled better performance in network training compared to other backend machine learning technologies technologies. For more explanation on the algorithms underpinning deep learning, see Appendix (INSERT) for more information

The general programming platform used was Python. Python was chosen due to its access to various important libraries, including NumPy, SciPy, Pomegrenate 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 Apped-ndix INSERT) for more information. Pomegranate was used to generate and compute the probabilistic model and ranking system for our Bayesian Network (INSERT see MM methods section N for more information). Finally, PyVCF was used to parse the VCF files into python objects for easy manipulation.

## **2.6 Artificial Datasets**

## **2.7 Processing tools**

## **2.8 PDX datasets**

# **3 Results and Discussion**

## **3.1 Generation of Artificial Genome**

In order to begin training

### **3.2 Feature Engineering**

### **3.3 Network Training and Optimisation**

### **3.4 Benchmarking of Network with Mason Datasets**

### **3.5 Benchmarking of Network with NA Datasets**

### **3.6 Analysis of PDX dataset using Bayesian Ranking systems**

## **4 Future Directions**

## **5 Appendixes**

### **5.1 Key Technologies**

## **6 Acknowledgements**

## **7 Bibilography**

## **References**

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