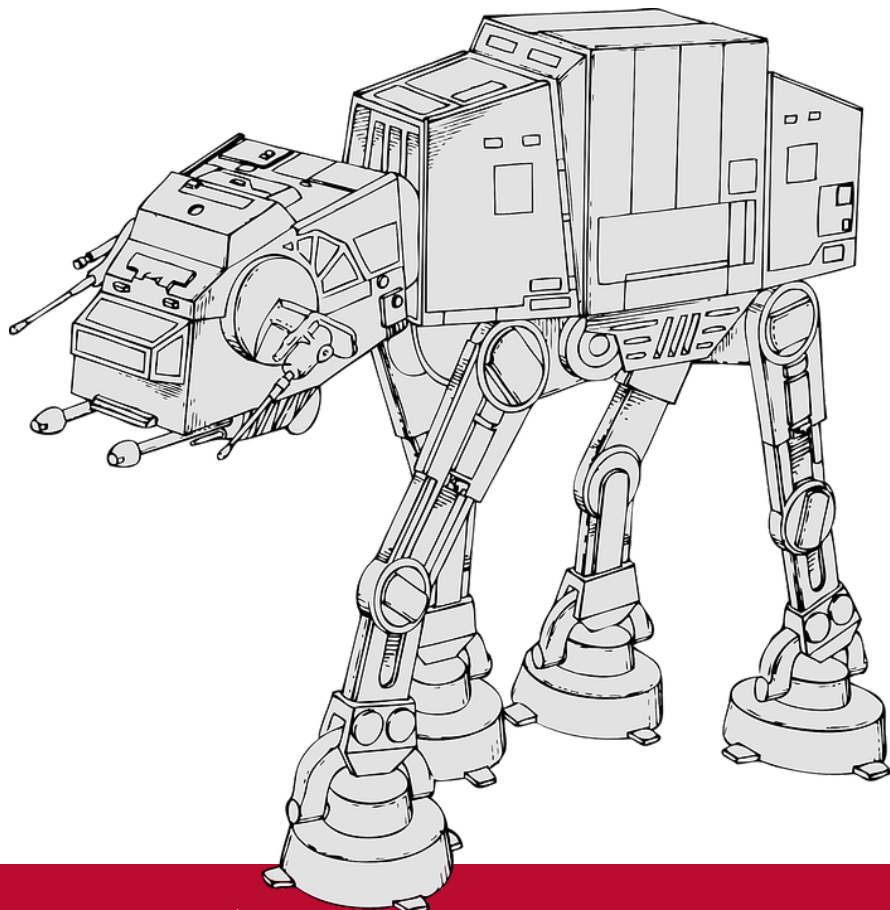


A Machine Learning Study Guide



Machine Learning Handbook

The Definitive Guide

ICS5110, class of 2018/9



L-Università
ta' Malta



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Introduction

This book explains popular Machine Learning terms. We focus to explain each term comprehensively, through the use of examples and diagrams. The description of each term is written by a student sitting in for ICS5110 APPLIED MACHINE LEARNING¹ at the University of Malta (class 2018/2019). This study-unit is part of the MSc. in AI offered by the Department of Artificial Intelligence, Faculty of ICT.

¹ <https://www.um.edu.mt/courses/studyunit/ICS5110>

Activation Functions

Caterini (2018) defined artificial neural networks as “a model that would imitate the function of the human brain—a set of neurons joined together by a set of connections. Neurons, in this context, are composed of a weighted sum of their inputs followed by a nonlinear function, which is also known as an activation function.”

Activation functions are used in artificial neural networks to determine whether the output of the neuron should be considered further or ignored. If the activation function chooses to continue considering the output of a neuron, we say that the neuron has been activated. The output of the activation function is what is passed on to the subsequent layer in a multilayer neural network. To determine whether a neuron should be activated, the activation function takes the output of a neuron and transforms it into a value commonly bound to a specific range, typically from 0 to 1 or -1 to 1 depending on the which activation function is applied.

Step Function

$$f(x) = \begin{cases} 0 & \text{for } x < 0 \\ 1 & \text{for } x \geq 0 \end{cases} \quad (1)$$

$$\frac{d}{dx} f(x) = \begin{cases} 0 & \text{for } x \neq 0 \\ ? & \text{for } x = 0 \end{cases} \quad (2)$$

The Heavside step function, visualised in figure 1 and defined by equation 1, is one of the simplest activation functions that can be used in a neural network. This function returns 0 if the input of a node is less than a predetermined threshold (typically 0), or otherwise it returns 1 if the output of the node is greater than or equal to the threshold. This activation function was first used in a machine learning context by Rosenblatt (1957) in his seminal work describing the perceptron, the precursor to the modern day neural network.

Nowadays, the step function is seldom used in practice as it cannot be used to classify more than one class. Furthermore, since the derivative of this function is 0, as defined by equation 2, gradient descent algorithms are not be able to progressively update the weights of a network that makes use of this function (Snyman, 2005).

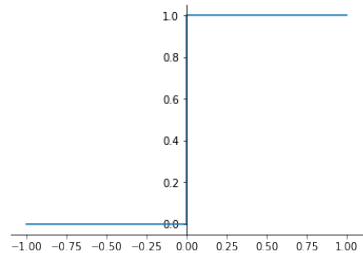


Figure 1: A graph of the step function.

Linear Functions

$$f(x) = ax + b \quad (3)$$

$$\frac{d}{dx} f(x) = a \quad (4)$$

A linear activation function, is any function in the format of equation 3, where $a, b \in \mathbb{R}$. This function seeks to solve some of the shortcomings of the step function. The output produced by a linear activation function is proportional to the input. This property means that linear activation functions can be used for multi-class problems. However, linear functions can only be utilised on problems that are linearly separable and can also run into problems with gradient descent algorithms, as the derivative of a linear function is a constant, as seen in equation 4. Additionally, since the output of the linear function is not bound to any range, it could be susceptible to a common problem when training deep neural networks called the exploding gradient problem, which can make learning unstable (Goodfellow et al., 2016).

Sigmoid Function

$$f(x) = \frac{1}{(1 + e^{-x})} \quad (5)$$

$$\frac{d}{dx} f(x) = f(x)(1 - f(x)) \quad (6)$$

The sigmoid function or logistic function, visualised in figure 2 and represented by equation 5, is one of the most commonly used activation functions in neural networks, because of its simplicity and desirable properties. The use of this function in neural networks was first introduced by Rumelhart et al. (1986), in one of the most important papers in the field of machine learning, which described the back-propagation algorithm and the introduction of hidden layers, giving rise to modern day neural networks. The values produced by the sigmoid function are bound between 0 and 1, both not inclusive, which help manage the exploding gradient problem. The derivative of this function, represented by equation 6, produces a very steep gradient for a relatively small range of values, typically in the range of -2 to 2 . This means that for most inputs that the function receives it will return values that are very close to either 0 or 1.

On the other hand, this last property makes the sigmoid function very susceptible to the vanishing gradient problem (Bengio et al., 1994). When observing the shape of the sigmoid function we see that towards the ends of the curve, the function becomes very unresponsive to changes in the input. In other words, the gradient of the function for large inputs becomes very close to 0. This can become very problematic for neural networks that are very deep in design, such as recurrent neural networks (RNNs). To address this

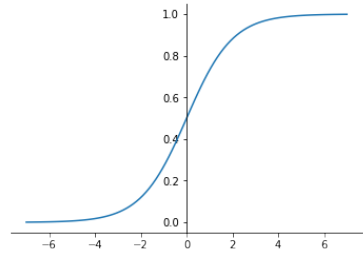


Figure 2: A graph of the sigmoid function.

problems in RNNs Long Short-Term Memory (LSTM) units where introduced as a variant of the traditional RNN architecture (Hochreiter and Schmidhuber, 1997).

Hyperbolic Tangent

$$f(x) = \frac{(e^x - e^{-x})}{(e^x + e^{-x})} \quad (7)$$

$$\frac{d}{dx} f(x) = 1 - f(x)^2. \quad (8)$$

The hyperbolic tangent (tanh) function, visualised in figure 3 and represented by equation 7, is another common activation function that is sometimes used instead of sigmoid. The tanh function has the same characteristics of the sigmoid function mentioned above. In fact, when comparing figure 2 to figure 3 one can observe that the tanh function is simply a scaled and translated version of the sigmoid function. As a result of this scaling and translation, the tanh function has a steeper gradient towards the origin, and it returns values between -1 and 1. The derivative of the hyperbolic tangent function is represented by equation 8.

LeCun et al. (2012) analysed various factors that affect the performance of backpropagation, and suggested that tanh may be better suited than sigmoid as an activation function due to its symmetry about the origin, which is more likely to produce outputs that are on average close to zero, resulting in sparser activations. This means that not all nodes in the network need to be computed, leading to better performance. Glorot and Bengio (2010) studied in detail the effects of the sigmoid and tanh activation functions and noted how the sigmoid function in particular is not well suited for deep networks with random initialisation and go on to propose an alternative normalised initialisation scheme which produced better performance in their experiments.

Rectified Linear Unit

$$f(x) = \begin{cases} 0 & \text{for } x < 0 \\ x & \text{for } x \geq 0 \end{cases} \quad (9)$$

$$\frac{d}{dx} f(x) = \begin{cases} 0 & \text{for } x < 0 \\ 1 & \text{for } x \geq 0 \end{cases} \quad (10)$$

The Rectified Linear Unit (ReLU) function, visualised in figure 4 and represented by equation 9, returns 0 if the input of the function is negative, otherwise it outputs the value of the input itself. This function is non-linear in nature even though at first glance it may seem similar to an identity function. The ReLU function is becoming one of the more commonly used activation functions due to its simplicity, performance, and suitability to networks with many layers. Another

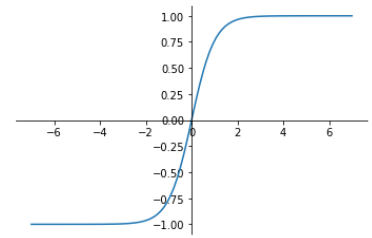


Figure 3: A graph of the hyperbolic tangent (tanh) function.

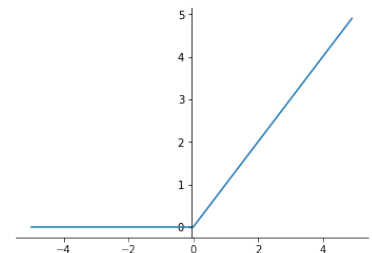


Figure 4: A graph of the ReLU function.

benefit of the ReLU function is that it produces sparse activations unlike many other commonly used functions such as the sigmoid.

The ReLU function has been used in many neural network models to improve their performance. Nair and Hinton (2010) use ReLU to improve the performance of Restricted Boltzmann Machines in object recognition. Krizhevsky et al. (2012a) introduced a breakthrough Convolutional Neural Network (CNN) architecture called AlexNet, which pioneered the use of the ReLU activation function together with dropout layers to minimise over fitting in CNNs.

Unfortunately, because the gradient of the function for inputs that are negative is 0, as seen in equation 10, the ReLU function can still be susceptible to the vanishing gradient problem. To manage this problem a variant of the ReLU function, called Leaky ReLU is sometimes used. Rather than simply returning 0 for negative inputs, the leaky ReLU returns a very small value such as $0.01x$. Maas et al. (2013) compared the performance of Sigmoid, ReLU and Leaky ReLU functions and found that while the the performance of both the ReLU and Leaky ReLU functions was better than the performance achieved with the sigmoid function, the performance of the two ReLU functions was nearly identical.

Bayesian Models in Machine Learning

Many machine learning techniques employ probability theory to make the best decisions given some data. Probabilistic approaches are often also summarised under the term "Bayesian approaches" (Murphy, 2012). The following chapter will provide an overview of the underlying theorem as well as the most popular classifier based on said theorem.

Bayes' Theorem

Bayes' Theorem describes how to calculate the so-called posterior probability of an event A given our observed evidence B , that is $P(A|B)$. It is derived from the definition of conditional probabilities (11). The probability of an event B to occur given that the event A already has occurred is defined as the probability of the intersection of the two events A and B divided by the probability of the event A (Murphy, 2012):

$$P(B|A) = \frac{P(A \cap B)}{P(A)} \quad (11)$$

The probability of the intersection of the two events A and B can be rewritten as (12). However, since $P(A \cap B)$ is equal to $P(B \cap A)$, it can be rewritten as (13), as well (Manning et al., 2009).

$$P(A \cap B) = P(A)P(B|A) \quad (12)$$

$$P(B \cap A) = P(B)P(A|B) \quad (13)$$

Consequently, Bayes' Theorem can be derived as follows:

$$\begin{aligned} P(B|A) &= \frac{P(A \cap B)}{P(A)} = \frac{P(B \cap A)}{P(A)} = \frac{P(B)P(A|B)}{P(A)} \\ \Leftrightarrow P(A|B) &= \frac{P(B|A)P(A)}{P(B)} \end{aligned} \quad (14)$$

$P(A|B)$ is the aforementioned posterior probability, $P(B|A)$ the likelihood of event B given event A , $P(A)$ the prior probability of event A , and $P(B)$ the so-called marginal, that is the likelihood of an independent occurrence of event B (Manning et al., 2009; van Rijsbergen, 1979).

Naïve Bayes Classification Algorithm

Intuitively, Bayes' Theorem can be used to calculate the probability of an hypothesis being true given some observed evidence (Manning et al., 2009; Jurafsky and Martin, 2018; van Rijsbergen, 1979) - it is a structured way of integrating all available information into an estimation of the truth of that hypothesis. However, it can also be used as a classification method (Manning et al., 2009; Jurafsky and Martin, 2018; Murphy, 2012; van Rijsbergen, 1979): The instance to be classified (such as an email, a web page, or a newspaper article) is taken to be the "observed evidence" for that instance to belong to a certain class. For example, the content of an email is taken to be the information available to classify that email as spam or ham. Bayes' Theorem is applied to the conditional probability $P(c|d)$ of a given document d belonging to a certain class c , resulting in (15). The denominator $P(d)$ can be dropped, since the prior probability of a document to occur is the same for all possible classes and does not have any effect on the final classification decision (Jurafsky and Martin, 2018). Ultimately, the result is (16).

$$P(c|d) = \frac{P(d|c)P(c)}{P(d)} \quad (15)$$

$$P(c|d) \propto P(c) \prod_{1 \leq k \leq n_d} P(t_k|c) \quad (16)$$

(16) can be read as the probability of a given document d belonging to a certain class c being proportional to the prior probability of a document occurring in class c multiplied by the probability of that class c having generated a document consisting of all the terms t_k in d . Naïve Bayes classifiers are also called generative classifiers (Murphy, 2012): Given a specific observation d , they return the class c that is most likely to have generated said observation - also called the maximum a posteriori (MAP) class.

The two parameters needed to calculate $P(c|d)$, $P(c)$ and $P(t_k|c)$, are obtained via approximations from the training set, based on maximum likelihood estimations (MLE) (Manning et al., 2009; Jurafsky and Martin, 2018). For the prior probability $P(c)$, the estimate is as follows:

$$\hat{P}(c) = \frac{N_c}{N}, \quad (17)$$

which is the number of documents in the training set belonging to class c divided by the total amount of documents in the training set (Manning et al., 2009; Jurafsky and Martin, 2018). Similarly, the probability of a term t_k occurring in a document of class c is obtained like this:

$$\hat{P}(t_k|c) = \frac{T_{ct}}{\sum_{t' \in V} T_{ct'}}, \quad (18)$$

which is the number of times a term T appears in documents of class c divided by the number of times that term appears in the vocabulary V , i.e. in all documents of the training set (Jurafsky and Martin, 2018; Manning et al., 2009).

These approximations pose two problems. First of all, the multiplication of all the conditional probabilities $P(t_k|c)$ may result in a so-called floating-point underflow (Manning et al., 2009). In order to avoid this, Naïve Bayes classifiers usually do not calculate the product over $P(t_k|c)$, but the sum over the log values of $P(t_k|c)$ (see (19)). Using the log values instead of the original ones does not interfere with the final classification decision in any way: The class with the highest log probability is still the one most likely to have generated the given document.

$$P(c|d) \propto P(c) \prod_{1 \leq k \leq n_d} \log P(t_k|c) \quad (19)$$

The second problem encountered by the estimations above is data sparseness, resulting in terms not occurring in documents of particular classes in the training set. The preferred solution is to eliminate the resulting zeros by using add-one or Laplace-smoothing (see (20), with $B' = |V|$, the length of the vocabulary), assuming that each and every relevant term occurs at least once in a corpus (Manning et al., 2009).

$$\hat{P}(t_k|c) = \frac{T_{ct} + 1}{\sum_{t' \in V} (T_{ct'} + 1)} = \frac{T_{ct} + 1}{(\sum_{t' \in V} T_{ct'}) + B'} \quad (20)$$

Naïve Assumptions

While the calculations above appear to be straightforward, they pose a problem in terms of their feasibility, especially regarding big sets of data. Relating individual probabilities to all terms of a document – even to those occurring multiple times in different positions – increases the number of parameters dramatically and requires huge sets of training data, which are almost never available. Naïve Bayes offers solutions to this problem via two rather naïve assumptions giving the classifier its name (Manning et al., 2009; Jurafsky and Martin, 2018).

Hence, Naïve Bayes classifiers assume the bag of words model for all terms in a document, that is they assume the probability of a term to be the same, regardless of its position in the document. This is also called the positional independence assumption (Manning et al., 2009; Jurafsky and Martin, 2018).

The second assumption is called the conditional independence assumption. It simplifies the calculation further by proposing that all terms in a document are independent of one another, i.e. the occurrence of a specific term does not make it any more likely for another, possibly related term to appear in the same document. Obviously, the exact opposite is true - in reality, the terms in a document are indeed

dependent on one another. However, Naïve Bayes classifiers still offer accurate classification decisions, regardless of the inaccuracies in their probability estimations (Manning et al., 2009; Jurafsky and Martin, 2018).

Conclusion

The classification method described just now is just one of the versions of Bayesian classifiers, called the multinomial Naïve Bayes classifier. One alternative is the Bernoulli model (Manning et al., 2009). In order to arrive at a classification decision, the Bernoulli model uses only binary information regarding the occurrence (1) or absence (0) of a term in a document. This often leads to errors regarding the classification of long documents, since the actual number of occurrences of a term are not taken into account, which is why the multinomial model is preferred.

Classification

Classification and regression are two techniques which are nowadays being used to extract knowledge and perform predictions on the huge amounts of available data (Sagar, 2017). These techniques predict values which may be either *quantitative* - involving discrete numerical values, or *qualitative* - with values within a particular class or label (James et al., 2013). Predicting marks that students will obtain in their final exams yields a discrete, continuous value. In this case, regression analysis is "used to predict missing or unavailable numerical data values" (Jiawei Han, Micheline Kamber, 2011). On the other hand, predicting the grade (A, B or C) that the students will obtain is considered as a classification problem, since the outcome value is a categorical class label (Jiawei Han, Micheline Kamber, 2011).

Classification is a data analysis technique in which models, also referred to as *classifiers*, are used to "predict categorical (discrete, unordered) class labels". Several scenarios in the real world may require classification models to predict several outcomes. For example, in the health sector, classification can be helpful in predicting whether a person is diagnosed with a particular medical condition or not, based on certain parameters related to experienced symptoms (Venkata Ramana et al., 2011; M. Alzahani et al., 2015). Financial sectors may find classifications helpful when it comes to predicting whether a company is bankrupt or in good financial health (Moradi et al., 2012). Moreover, they may apply classifiers to determine whether a bank should issue a loan to a customer (Thomas, 2000). In these scenarios, classifiers predict labels such as 'positive' or 'negative' for a medical condition, or 'bankrupt' or 'healthy' for the financial status of a company. Since values predicted by a classifier are discrete, any ordering amongst the values is meaningless.

Classification involves two main steps: training a model for classification, and then using the model to classify the data. When *supervised learning* is used for classification, the training stage involves analysing a training data consisting of samples which are labelled with a categorical class. These samples are used to train the model on how to perform classifications. Testing data, also made up of features and their corresponding class labels, is then used to predict the accuracy of the classifier, by comparing the predicted outcome to the actual class label in the testing samples (Neelamegam and Ramaraj, 2013). On the other hand, *unsupervised learning* builds classifiers for predictions

using data which is unlabelled, i.e. the final class is not known. In this case, the model learns how to classify the data by observing the similarities or dissimilarities within the dataset (Jiawei Han, Micheline Kamber, 2011).

Binary Classification

When two classes are available as outputs of a classification problem, then this is called binary classification (Neelamegam and Ramaraj, 2013). Kolo (2011) defines *binary classifiers* as those which determine whether an input should classify as "either in the category or not in the category". This implies that an input which classifies as not part of a class should automatically be considered as a member of the other class.

For example, using a binary classifier to predict 'Pass' or 'Fail' in a subject, by providing it with the marks obtained in the exam, the classifier should determine whether the student 'passed' or 'did not pass' an exam. A 'Fail' label is assumed when the classifier determines that the student 'did not pass' the exam (Kolo, 2011). Decision Trees (Figure 5), Bayesian Networks and Logistic Regression are some of the methods used for binary classification.

Probabilistic Classification

Probabilistic classification can be considered as a subclass of classification, in which apart from assigning categorical classes to unseen data, probabilistic models "use statistical inference to find the best class for a given example" (Aggarwal, 2014). This implies that probabilistic models compute the probability that an unseen example belongs to the possible target classes, and then chooses the final target class by considering the highest probability obtained. One advantage of probabilistic models is that the prediction outcome is composed of both the target label, as well as the confidence with which this class label was obtained.

Initially, probabilistic classification involves the estimation of the posterior probability, $P(C_k | x)$, where C_k denotes the class and x refers to the input variables. This posterior probability estimate can be determined using a *generative model* or a *discriminative model*. Generative models work by obtaining the probability of each class, denoted by $P(C_k)$, as well as the probability of obtaining x when given a class, also known as the class-conditional probabilities, denoted by $P(x | C_k)$ (Aggarwal, 2014). Then, the posterior probability can be determined using Bayes Theorem, shown in Equation 21, or by modelling the joint distribution over x and C_k as in Equation 22.

$$P(C_k | x) = P(C_k) \cdot \frac{P(x | C_k)}{P(x)} \quad (21)$$

$$P(C_k | x) \propto P(C_k) \cdot P(x | C_k) \quad (22)$$

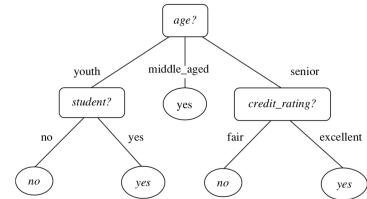


Figure 5: A binary Decision Tree example which determines whether a customer will purchase a computer or not. Reproduced from Jiawei Han, Micheline Kamber (2011).

In these equations C_k denotes the class and x refers to the input variables. Hidden Markov Model and Naive Bayes classifier are two examples of generative models used in probabilistic classification. Discriminative models focus on determining a function that maps the input x directly onto the class C_k .

$$P(C_k|x) = \frac{1}{1 + e^{-\theta^T X}} \quad (23)$$

One popular discriminative model is the Logistic Regression Model, explained by Equation 23, in which θ refers to the parameters to be estimated. Conditional Random Fields is another commonly used discriminative model.

Once the posterior probability is obtained, probabilistic classification then uses decision theory to deduce the class of an unseen example x (Aggarwal, 2014).

Multiclass Classification

Multiclass classification models are able to determine a class for an unseen example from k classes, where $k > 2$. A multiclass classification problem may be solved by extending the aforementioned binary classification algorithms, and others including *Neural Networks* and *Support Vector Machines* (Aly, 2005). For example, a Decision Tree algorithm can naturally handle multiclass classification problems. To predict an outcome using the tree model, the algorithm starts from the root node of the tree and repeatedly traverses branches based on the feature values, until a leaf node is reached (Rokach and Maimon, 2014). The value of the leaf node is considered as the predicted class label of the unseen example, and these values can refer to any of the k classes. Support Vector Machines are another example of classification algorithms which were initially built for binary classification, but eventually extended to be applied in multiclass problems. Extensions of this algorithm handle the separation of the various classes by amending the optimization problem (Aly, 2005).

Aly (2005) discusses approaching a multiclass classification problem by decomposing it into multiple binary classification problems, which can then be solved using binary classifiers.

One-versus-all (OVA)

This approach considers k binary classifiers, where k is the number of classes. Each classifier is trained on data composed of positive examples from the k^{th} class and negative examples from the remaining $k-1$ classes. Unseen examples are assigned the class of the classifier which obtains the highest output (Aly, 2005).

All-versus-all (AVA)

In this approach, $\frac{k(k-1)}{2}$ binary classifiers are built to distinguish between all the possible pairs of classes. When predicting, all classifiers are considered and the class obtaining the highest score is taken as the target class (Aly, 2005).

Error-Correcting Output-Coding (ECOC)

Using this approach for multiclass classification, n binary classifiers are trained to differentiate between K classes. A binary matrix M (Figure 6) is composed of K classes and the corresponding codewords with length N . Thus, binary classifiers are trained using the column values of M .

When predicting labels for unseen examples, a distance measure is used to compare codewords obtained from the N classifiers to the K codewords. The class label is determined by selecting the codeword having the smallest distance (Aly, 2005).

Apart from these methods, *Generalized Coding* (L. Allwein et al., 2001) may also be applied. Finally, multiclass classification problems may be handled using *hierarchical classification*, in which classes are arranged in the form of a tree. Parent nodes of the trees are further split into child nodes, and a binary classifier is used to differentiate between the clusters pertaining to child classes. Eventually, the leaf node constitutes of a single class, thus the model can then be used to classify a new example (Aly, 2005). Binary Hierarchical Classifier (BHS) (Kumar et al., 2002) and Hierarchical SVM (HSVM) (Yangchi Chen et al., 2014) are two hierarchical classification approaches.

	f_1	f_2	f_3	f_4	f_5	f_6	f_7
Class 1	0	0	0	0	0	0	0
Class 2	0	1	1	0	0	1	1
Class 3	0	1	1	1	1	0	0
Class 4	1	0	1	1	0	1	0
Class 5	1	1	0	1	0	0	1

Figure 6: ECOC Binary Matrix. Reproduced from Aly (2005).

Confusion Matrix

A *confusion matrix* (CM), is a contingency table showing how well a model classifies categorical data. By convention (Sammut and Webb, 2017), the CM of an N-class model is an $N \times N$ matrix indexed by the true class in the row dimension and the predicted class in the column dimension (Table 1).

		Predicted Class	
		spam	\neg spam
True Class	spam	10	1
	\neg spam	2	100

Even though CMs are commonly used to evaluate binary classifiers, they are not restricted to 2-class models (Jurafsky and Martin, 2018). A CM of a multi-class model would show the number of times the classes were predicted correctly and which classes were confused with each other (Table 2).

	<i>M&M's</i>	<i>Skittles</i>	<i>Smarties</i>
<i>M&M's</i>	34	3	8
<i>Skittles</i>	1	28	5
<i>Smarties</i>	2	4	22

The CM of the model $h : X \mapsto C$ over the concept $c : X \mapsto C$ using dataset $S \subset X$ is formally defined as a matrix Ξ such that $\Xi_{c,S}(h)[d_1, d_2] = |S_{h=d_1, c=d_2}|$ (Cichosz, 2014). The CM is constructed by incrementing the element corresponding to the true class *vis-a-vis* the predicted class for each object in the dataset (Algorithm 1).

```

 $\Xi \leftarrow 0$ 
for  $x \in S$  do
     $d_1 \leftarrow c(x)$ 
     $d_2 \leftarrow h(x)$ 
     $\Xi_{d_1, d_2} \leftarrow \Xi_{d_1, d_2} + 1$ 

```

In binary classification, the CM consists of 2 specially designated classes called the *positive* class and the *negative* class (Saito and Rehmsmeier, 2015b). As indicated in Table 3, positive outcomes from the true class which are classified correctly are called *true positives* (TP), whilst misclassifications are called *false negatives* (FN). On

Table 1: CM of a hypothetical binary classifier which predicts whether out-of-sample text objects are spam or not. In this example, 10 spam and 100 non-spam objects are classified correctly, whilst 1 spam and 2 non-spam objects are misclassified.

Table 2: CM of a hypothetical sweets classifier. The main diagonal of the CM shows the number of correct predictions, whilst the remaining elements indicate how many sweets were misclassified.

Algorithm 1: The CM is initialised to the zero matrix, and populated by iterating over all the objects x with corresponding true class d_1 and predicted class d_2 and incrementing the element (d_1, d_2) by 1 for each matching outcome.

the other hand, negative true class outcomes which are classified correctly are called *true negatives* (TN), and misclassifications are called *false positives* (FP). In natural sciences, FP are called *Type I* errors and FN are known as *Type II* errors (Fielding and Bell, 1997).

	+ve	-ve
+ve	TP	FN
-ve	FP	TN

The information presented in the CM can be used to evaluate the performance of different binary classifiers (Lu et al., 2004). A number of statistics (Equations 24-30) derived from the CM have been proposed in the literature (Deng et al., 2016) to gain a better understanding of what are the strengths and weaknesses of different classifiers. Caution should be exercised when interpreting metrics (Jeni et al., 2013), since the CM could be misleading if the data is imbalanced and an important subrange of the domain is underrepresented (Raeder et al., 2012). For instance, an albino zebra classifier which always returns negative will achieve high accuracy since albinism is a rare disorder.

These metrics are important in situations in which a particular type of misclassification, i.e. FP or FN, could have worse consequences than the other (Hassanien and Oliva, 2017). For example, FP are more tolerable than FN in classifiers which predict whether a patient has a disease. Both outcomes are undesirable, but in medical applications it is better to err on the side of caution since FN could be fatal.

Accuracy (ACC) is the proportion of correct predictions (Equation 24). It is a class-insensitive metric because it can give a high rating to a model which classifies majority class objects correctly but misclassifies interesting minority class objects (Branco et al., 2016). The other metrics should be preferred since they are more class-sensitive and give better indicators when the dataset is imbalanced.

$$ACC = \frac{|TP \cup TN|}{|TP \cup FP \cup TN \cup FN|} \quad (24)$$

Negative predictive value (NPV) is the ratio of the correct negative predictions from the total negative predictions (Equation 25).

$$NPV = \frac{|TN|}{|TN \cup FN|} \quad (25)$$

True negative rate (TNR), or *specificity*, is the ratio of the correct negative predictions from the total true negatives (Equation 26).

$$TNR = \frac{|TN|}{|TN \cup FP|} \quad (26)$$

True positive rate (TPR), also called *sensitivity* or *recall*, is the ratio of the correct positive predictions from the total true positives (Equation 27).

$$TPR = \frac{|TP|}{|TP \cup FN|} \quad (27)$$

Table 3: CMs of binary classifiers have positive (+ve) and negative (-ve) classes, and elements called *true positives* (TP), *false positives* (FP), *true negatives* (TN) and *false negatives* (FN).

Sensitivity and specificity can be combined into a single metric (Equation 28). These metrics are often used in domains in which minority classes are important (Kuhn and Johnson, 2013). For example, the sensitivity of a medical classifier (El-Dahshan et al., 2010) measures how many patients with the condition tested positive, and specificity measures how many did not have the condition and tested negative.

$$\text{Sensitivity} \times \text{Specificity} = \frac{|TP| \times |TN|}{|TP \cup FN| \times |TN \cup FP|} \quad (28)$$

Positive predictive value (PPV), or *precision*, is the ratio of the correct positive predictions from the total positive predictions (Equation 29). The difference between accuracy and precision is depicted in Figure 7.

$$PPV = \frac{|TP|}{|TP \cup FP|} \quad (29)$$

Precision and recall are borrowed from the discipline of *information extraction* (Sokolova and Lapalme, 2009). A composite metric called *F-score*, *F1-score*, or *F-measure* (Equation 30) can be derived by finding their harmonic mean (Kelleher et al., 2015).

$$F\text{-score} = 2 \times \frac{PPV \times TPR}{PPV + TPR} \quad (30)$$

The complements of ACC, NPV, TNR, TPR and PPV are called, respectively, *error rate*, *false omission rate*, *false positive rate*, *false negative rate* and *false discovery rate*.

The metrics can be adapted for evaluating multi-class models by decomposing an N-class CM into 2-class CMs, and evaluating them individually (Stager et al., 2006). The literature describes two methods for decomposing this kind of CM. In the *1-vs-1* approach, 2-class CMs are constructed for each pairwise class as shown in Table 4.

	+ve	-ve
M&M's	{Skittles, Smarties}	
Skittles	{M&M's, Smarties}	
Smarties	{M&M's, Skittles}	

In the *1-vs-rest* approach, 2-class CMs are constructed for each class and the remaining classes combined together as shown in Table 5.

	+ve	-ve
M&M's	Skittles \cup Smarties	
Skittles	M&M's \cup Smarties	
Smarties	Skittles \cup M&M's	

Using all metrics could be counterproductive due to information redundancy, but none of the metrics is enough on its own (Ma and Cukic, 2007). For instance, recall is class-sensitive but it would give a perfect score to an inept model which simply returns the positive

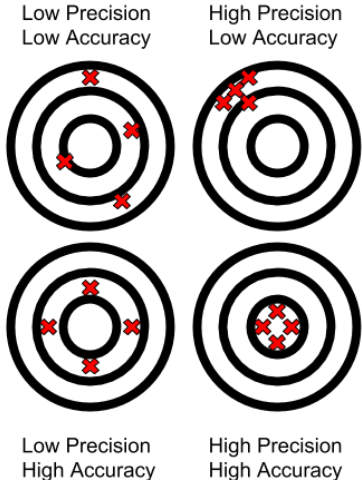


Figure 7: Accuracy vs Precision.

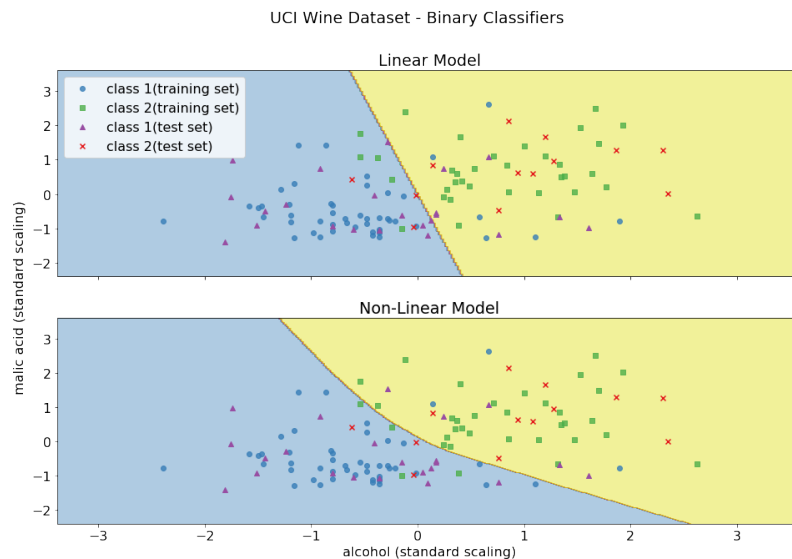
Table 4: 2-class CMs derived from the classes in Table 2. The +ve classes are paired separately with each -ve class.

Table 5: 2-class CMs derived through decomposition of the 3-class CM from Table 2 using the 1-vs-rest approach.

class. Thus, the best approach is to evaluate with complementary pairs (Gu et al., 2009) such as sensitivity vs specificity, or precision vs recall; or a combined measure such as the F-score.

Taking into account the above, CMs are suitable for visualising, evaluating, and comparing the performance of binary or multi-class classifiers. They should be used in conjunction with metrics such as the F-measure to avoid bias, especially if the dataset is unbalanced. For further details on the theoretical aspects of CMs and for practical examples in R refer to (Cichosz, 2014); for examples in Python refer to (Müller et al., 2016).

The following example is motivated by the samples in the *Scikit-Learn* documentation and the work of (Géron, 2017). The models in Figure 8 were trained on the *wines* dataset included with Scikit-Learn.



As it can be deduced from Figure 8, the decision boundary of the non-linear model is a better fit than the linear model. The CMs in Figure 9 also show that non-linear model performs better with a higher TP, and consequently lower TN. The biggest advantage of the non-linear model is the higher sensitivity resulting in a better F-score.

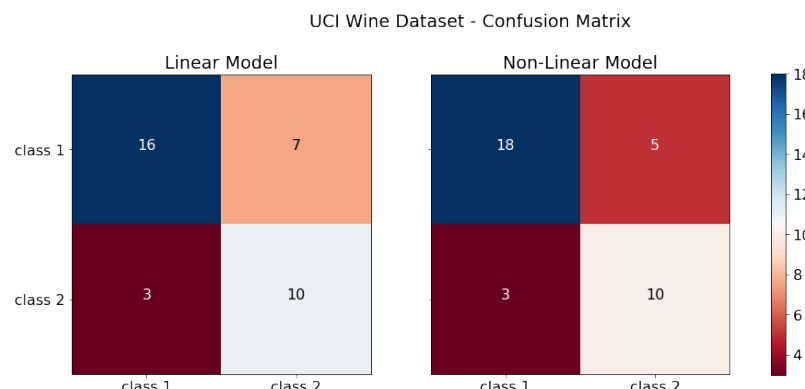


Figure 8: Decision boundary learned by a linear and non-linear binary classifier.

	Linear	Non-Linear
Accuracy	0.72	0.78
Specificity	0.77	0.77
Sensitivity	0.70	0.78
Precision	0.84	0.86
F-score	0.76	0.82

Table 6: Statistics derived from the CMs in Figure 9.

Figure 9: The linear classifier has 16 TP, 10 TN, 7 FN and 3 FP, whilst the non-linear classifier has 18 TP, 10 TN, 5 FN and 3 FP.

Convolution

Convolution in Mathematics

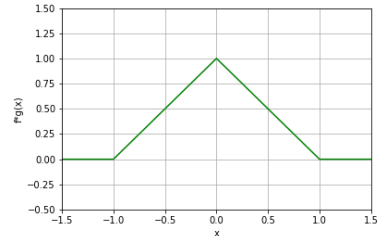
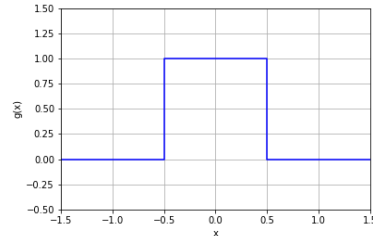
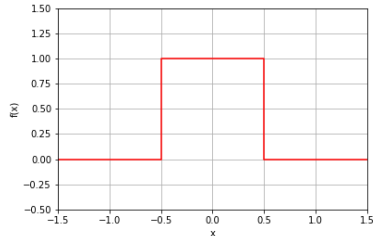
Definition

Convolution is a mathematical operation on two functions to produce a result that reflects how one of the input functions is modified by the other input function.

The convolution of functions $f(x)$ and $g(x)$ (denoted $f * g(x)$) is defined (Bracewell, 2000), for continuous functions f and g , as²

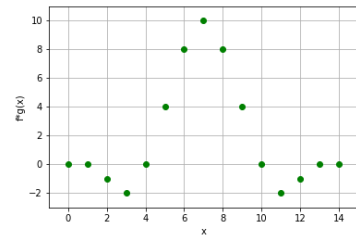
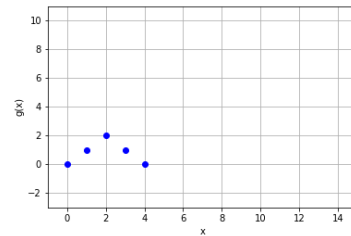
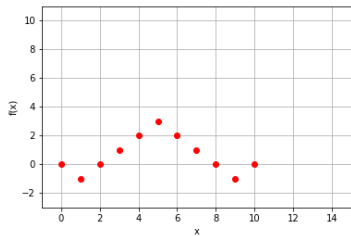
$$(f * g)(t) \equiv \int_{-\infty}^{\infty} f(\tau)g(t - \tau)d\tau \quad (31)$$

² In order for the continuous convolution operator $f * g$ to be defined, both f and g must be integrable functions.



and, for discrete functions $f(n)$ and $g(n)$, as

$$(f * g)(n) \equiv \sum_{m=-\infty}^{\infty} f(m)g(n-m) \quad (32)$$



Properties

Since convolution is defined as a product of integrable functions on the linear space, the following algebraic properties are satis-

Figure 10: These plots show two continuous functions ($f(x)$ and $g(x)$) and their convolution $f * g(x)$

Figure 11: These plots show two discrete functions ($f(n)$ and $g(n)$) and their convolution $f * g(n)$

fied (Bracewell, 2000):

Commutativity

$$(f * g) = (g * f) \quad (33)$$

Associativity

$$(f * (g * h)) = ((f * g) * h) \quad (34)$$

Distributivity

$$(f * (g + h)) = (f * g) + (f * h) \quad (35)$$

Multiplication by a scalar value

$$a(f * g) = (af) * g \quad (36)$$

Multiplicative identity, where δ denotes the delta distribution

$$f * \delta = f \quad (37)$$

Differentiation

$$\frac{d}{dx}(f * g) = \frac{df}{dx} * g = f * \frac{dg}{dx} \quad (38)$$

Integration

$$\int_{R^d} (f * g)(x) dx = \left(\int_{R^d} f(x) dx \right) \left(\int_{R^d} g(x) dx \right) \quad (39)$$

Applications

Listed below are some of the main applications of the convolution operator in various fields of knowledge (Srivastava and Buschman, 2013):

- **Image Processing** - Different operations are performed on images, in which the original image (larger matrix) and the filter to be applied (smaller matrix, also known as 2D kernel) are treated as 2-dimensional arrays. The kernel size and the values of its elements determine the effect on the original image
- **Signal Filtering** - Provided the filter function is the same as the impulse response function used in signal filtering, the two operations are equivalent
- As a handy tool for **Polynomial Multiplication** - If we consider two polynomials being multiplied, we can use a convolution process to obtain the coefficients of the resulting polynomial
- **Audio Processing** - Reverberation is a desired effect in auditoriums, music halls, cinemas, and similar constructions. Convolution is used to digitally simulate reverberation in such structures, providing architects with information about the acoustic quality of a building prior to its construction
- **Artificial Intelligence** - Convolutional Neural Networks use convolution in one or more of its internal layers in order to process input data and enhance specific features.

- **Probability Theory** - The Probability Density Function (PDF) of the sum of two independent random variables can be obtained by the convolution of the PDFs of the two variables

Convolution in Neural Networks

Definition

Convolutional Neural Networks (CNNs) are a subclass of multi-layer neural networks in which some of its hidden layers are convolutional layers.

Like most neural networks, CNNs can be trained using back propagation algorithms and are mostly used to recognize visual patterns with minimal or no preprocessing ([Lawrence et al., 1997](#)).

Origins and Evolution

CNNs were initially developed as an attempt to replicate the processing of sight in living organisms.

A seminal paper published in 1968 ([Hubel and Wiesel, 1968](#)) noted that two types of neurons took part in the process of identifying images: simple cells (responsible for the detection of straight edges and their orientation), and complex cells (with larger receptive fields, but not affected by the exact position of the edges in the input image). During the 1980s, CNNs evolved, followed by the introduction of Time Delay Neural Networks (TDNNs) in 1987 ([Waibel et al., 1989](#)). In the early 1990s, CNNs were modified and used for medical image processing and for automatic detection of breast cancer in mammograms ([Zang et al., 1994](#)).

In 1998 LeCun lead a team that created LeNet-5, a CNN used by banks to identify handwritten digits in checks.

With the introduction of Graphic Processing Units (GPUs), training of CNNs was greatly improved, allowing for the implementation of efficient Deep Learning Neural Networks with impressive results in image processing and other applications ([Ciresan et al., 2013](#)).

Layers

A typical CNN has sets of layers with specific functions, such as:

- **Convolutional layer** - This is the main feature of a CNN.

In this layer, a set of filters (also known as kernels) is convoluted over the entire input data. The result of this operation (the activation map of the filter), extracts or enhances specific features in the input data that are passed to the following layers in the CNN.

Since the result of each convolution operation is affected by only a small part of the input data (due to the reduced size of the filter if compared to the size of the input data), this can be interpreted

as each point of the activation map being affected by only a small subset of input points.

- **Pooling layer** - In this layer, simple operations are applied in order to reduce the size of the input data, such as maximum pooling or average pooling.
In 2×2 average pooling, for example, each result point is the average of 4 input points, reducing the size of the input field by a factor of 4.
- **Rectified Linear Unit (ReLU) layer** - This layer applies the $f(x) = \max(0, x)$ activation function to its input, removing negative values from the input field.
- **Fully Connected layer** - In this layer, usually applied after several convolutional and pooling layers, neurons are connected to all activations in the previous layer, as in regular neural networks.
- **Loss layer** - This is the last layer in a typical CNN. In this layer, training is based on the divergence between the desired and predicted labels.

Training

One important feature of CNNs is that, with the exception of the Loss layer, all other layers (Convolutional, Pooling, ReLU, and Fully Connected layers) can be trained using unsupervised training ([Arel et al., 2010](#)).

Also, usually each layer or set of layers is trained individually, which also improves the speed and the overall result of the training.

With the introduction of Graphic Processing Units (GPUs), speed of training and processing in the Convolutional layer has been greatly improved ([Steinkrau et al., 2005](#))

Applications

CNNs are currently applied to a wide range of areas, such as ([Schmidhuber, 2015](#)):

- **Image recognition and Video analysis** - Image Recognition Systems using CNNs are very efficient, especially if applied to facial recognition. Very good results have been obtained in challenges, and in the ImageNetNet tests performed almost as good as humans. If compared to image recognition, video analysis adds a level of complexity, since, in addition to the analysis of each frame, a time dimension needs to be added to the problem and convolution needs to be performed both on each image as well as on the time domain.
- **Game strategy** - A number of board games have been implemented using CNNs, most notably Checkers, in Chess and in Go, the first time Artificial Intelligence beat an expert human player in this game.

- **Drug discovery** - In this area, CNNs are used to predict the behaviour of drug molecules and human proteins.
- **Natural language processing** - For NLP, CNNs are used extensively, including semantic parsing, search query retrieval, sentence modeling, classification, and prediction.

Convolutional Filters

Definition

Convolutional filters (also known as kernels), when applied to image and video processing, are small square matrices with an odd number of rows and columns and used in the Convolutional layer of CNNs in order to detect or enhance specific features in the input image.

The convolution operation performed by CNNs is a special two dimensional case of the discrete convolution operator.

$$R(m, n) = (K * I)(m, n) \equiv \sum_{s=-a}^a \sum_{t=-b}^b K(s, t) I(m-s, n-t) \quad (40)$$

where $R(m, n)$ is the convoluted result at point m, n , K is the kernel matrix being applied to the input matrix, I is the input matrix, and a and b depend on the dimensions of the kernel used.

Practical example: Edge Detection kernel applied

The figure below shows a practical example of the application of the Edge Detection kernel:

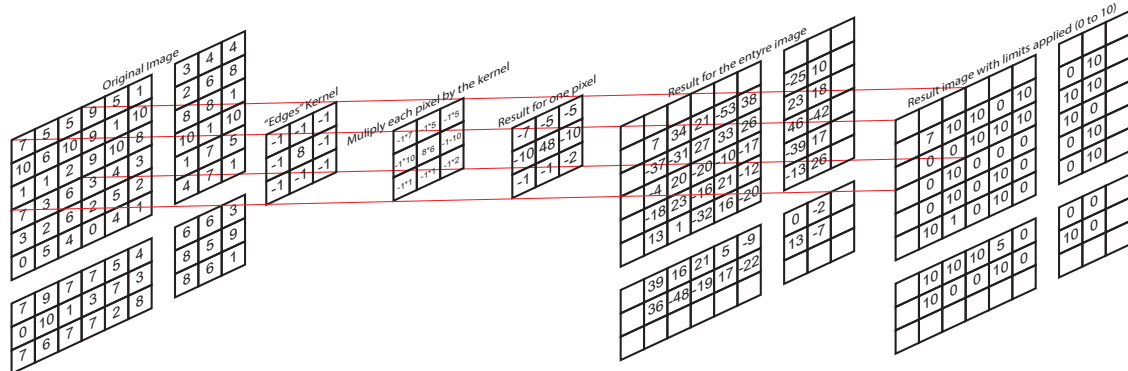


Figure 12: Edge Detection kernel applied

- The kernel is moved over the entire image, pixel by pixel
- At each position, the value for each pixel in the original image is multiplied by the corresponding value in the kernel ($7 * -1 = -7$; $5 * -1 = -5$; $5 * -1 = -5$; $10 * -1 = -10$; $6 * 8 = 48$; $10 * -1 = -10$; $1 * -1 = -1$; $1 * -1 = -1$; $2 * -1 = -2$)
- All results are added $((-7) + (-5) + (-5) + (-10) + 48 + (-10) + (-1) + (-1) + (-2)) = 7$

- Optionally, the result of the summation may be limited to valid values (colors in images typically range from 0 to 255. In this example we have limited the values to $[0 - 10]$)

Common Convolutional Filters

Convolutional filters can be defined to achieve a large range of effects in the input image.

A small sample of possible kernels is provided below:

$$\begin{bmatrix} 0 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 0 \end{bmatrix}$$



Figure 13: **Identity Kernel** and the original image (unchanged, no convolution applied)

$$\frac{1}{16} \begin{bmatrix} 1 & 2 & 1 \\ 2 & 4 & 2 \\ 1 & 2 & 1 \end{bmatrix}$$



Figure 14: **Gaussian Blur Kernel** and the resulting image

$$\frac{1}{9} \begin{bmatrix} 1 & 1 & 1 \\ 1 & 1 & 1 \\ 1 & 1 & 1 \end{bmatrix}$$



Figure 15: **Box Blur Kernel** and the resulting image

$$\frac{-1}{256} \begin{bmatrix} 1 & 4 & 6 & 4 & 1 \\ 4 & 16 & 24 & 16 & 4 \\ 6 & 24 & -476 & 24 & 6 \\ 4 & 16 & 24 & 16 & 4 \\ 1 & 4 & 6 & 4 & 1 \end{bmatrix}$$



Figure 16: **Unsharp** Kernel and the resulting image

$$\begin{bmatrix} 0 & -1 & 0 \\ -1 & 5 & -1 \\ 0 & -1 & 0 \end{bmatrix}$$



Figure 17: **Sharpen** Kernel and the resulting image

$$\begin{bmatrix} -1 & -1 & -1 \\ -1 & 8 & -1 \\ -1 & -1 & -1 \end{bmatrix}$$



Figure 18: **Edge Detection** Kernel and the resulting image

Cross-Validation

Cross-validation (CV) is an estimation method used on supervised learning algorithms to assess their ability to predict the output of unseen data (Varma and Simon, 2006; Kohavi, 1995). Supervised learning algorithms are computational tasks like classification or regression, that learn an input-output function based on a set of samples. Such samples are also known as the labeled training data where each example consists of an input vector and its correct output value. After the training phase, a supervised learning algorithm should be able to use the inferred function in order to map new input unseen instances, known as testing data, to their correct output values (Caruana and Niculescu-Mizil, 2006). When the algorithm incorporates supervised feature selection, cross-validation should always be done external to the selection (feature-selection performed within every CV iteration) so as to ensure the test data remains unseen, reducing bias (Ambroise and McLachlan, 2002; Hastie et al., 2001). Therefore, cross-validation, also known as out-of-sample testing, tests the function's ability to generalize to unseen situations (Varma and Simon, 2006; Kohavi, 1995).

Cross-validation has two types of approaches, being i) the exhaustive cross validation approach which divides all the original samples in every possible way, forming training and test sets to train and test the model, and ii) the non-exhaustive cross validation approach which does not consider all the possible ways of splitting the original samples (Arlot et al., 2010).

The above mentioned approaches are further divided into different cross-validation methods, as explained below.

Exhaustive cross-validation

Leave-p-out (LpO)

This method takes p samples from the data set as the test set and keeps the remaining as the training set, as shown in Fig. 20a. This is repeated for every combination of test and training set formed from the original data set and the average error is obtained. Therefore, this method trains and tests the algorithm $\binom{n}{p}$ times when the number of samples in the original data set is n , becoming inapplicable when $p > 1$ (Arlot et al., 2010).

Leave-one-out (LOO)

This method is a specific case of the LpO method having $p = 1$. It requires less computation efforts than LpO since the process is only repeated $n\text{choose}1 = n$ times, however might still be inapplicable for large values of n (Arlot et al., 2010).

Non-exhaustive cross-validation

Holdout method

This method randomly splits the original data set into two sets being the training set and the test set. Usually, the test set is smaller than the training set so that the algorithm has more data to train on. This method involves a single run and so must be used carefully to avoid misleading results. It is therefore sometimes not considered a CV method (Kohavi, 1995).

k -fold

This method randomly splits the original data set into k equally sized subsets, as shown in Fig. 21. The function is then trained and validated k times, each time taking a different subset as the test data and the remaining $(k - 1)$ subsets as the training data, using each of the k subsets as the test set once. The k results are averaged to produce a single estimation. Stratified k -fold cross validation is a refinement of the k -fold method, which splits the original samples into equally sized and distributed subsets, having the same proportions of the different target labels (Kohavi, 1995).

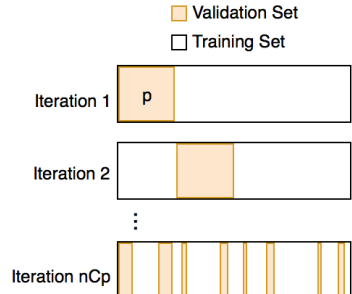


Figure 19: Leave-p-Out Exhaustive Cross Validation

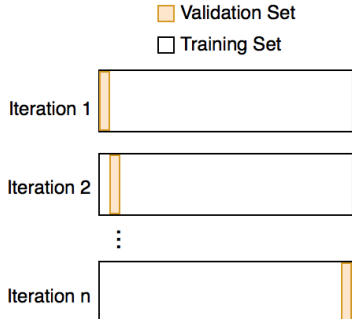


Figure 20: Leave-One-Out Exhaustive Cross Validation

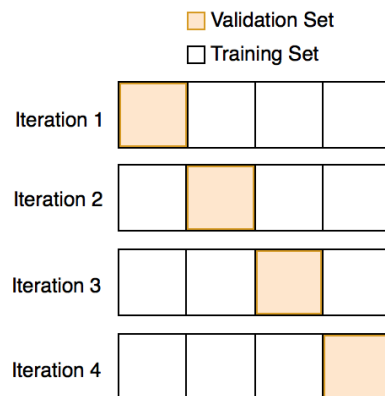


Figure 21: k -Fold Cross Validation where $k=4$

Repeated random sub-sampling

This method is also known as the Monte Carlo CV. It splits the data set randomly with replacement into training and test subsets using some predefined split percentage, for every run. Therefore, this generates new training and test data for each run but the test data

of the different runs might contain repeated samples, unlike that of k -fold (Xu and Liang, 2001).

All of the above cross-validation methods are used to check whether the model has been overfitted or underfitted and hence estimating the model's ability of fitting to independent data. Such ability is measured using quantitative metrics appropriate for the model and data (Kohavi, 1995; Arlot et al., 2010). In the case of classification problems, the misclassification error rate is usually used whilst for regression problems, the mean squared error (MSE) is usually used. MSE is represented by Eq. 41, where n is the total number of test samples, Y_i is the true value of the i^{th} instance and \hat{Y}_i is the predicted value of the i^{th} instance.

$$MSE = \frac{1}{n} \sum_{i=1}^n (Y_i - \hat{Y}_i)^2 \quad (41)$$

Underfitting is when the model has a low degree (e.g. $y = x$, where the degree is 1) and so is not flexible enough to fit the data making the model have a low variance and high bias (Baumann, 2003), as seen in Fig. 23a. Variance is the model's dependence on the training data and bias is model's assumption about the shape of the data (Arlot et al., 2010). On the other hand, as seen in Fig. 23b, overfitting is when the model has a too high degree (e.g. $y = x^{30}$, where the degree is 30) causing it to exactly fit the data as well as the noise and so lacks the ability to generalize (Baumann, 2003), making the model have a high variance. Cross-validation helps reduce this bias and variance since it uses most of the data for both fitting and testing and so helps the model learn the actual relationship within the data. This makes cross-validation a good technique for models to acquire a good bias-variance tradeoff (Arlot et al., 2010).

As stated in (Kohavi, 1995), the LOO method gives a 0% accuracy on the test set when the number of target labels are equal to the number of instances in the dataset. It is shown that the k -fold CV method gives much better results, due to its lower variance, especially when $k = 10, 20$. Furthermore, R. Kohavi et al. state that the best accuracy is achieved when using the stratified cross-validation method, since this has the least bias.

Therefore, let's take an example using the stratified k -fold cross-validation method with $k = 10$. Let's say that we are trying to solve age group classification, using eight non-overlapping age groups being 0-5, 6-10, 11-20, 21-30, 31-40, 41-50, 51-60, and 61+. We are using the FG-NET labelled data set, which contains around 1000 images of individuals aged between 0 and 69. Before we can start training our model (e.g. CNN), we must divide our data set into training and test subsets and this is where cross validation comes in. Therefore, we start by taking the 1000 images of our data set and splitting them according to their target class. Let us assume we have an equal amount of 125 (1000/8) images per class³. As depicted in Fig. 24, we can now start forming our 10 folds by taking 10% of each age-group bucket, randomly without replacement. Hence, we will

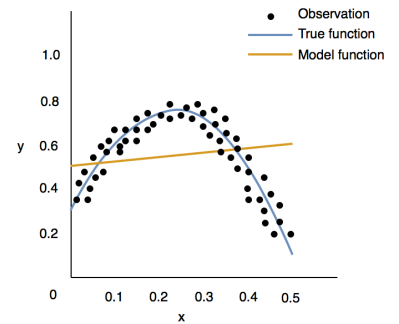


Figure 22: Model Underfitting

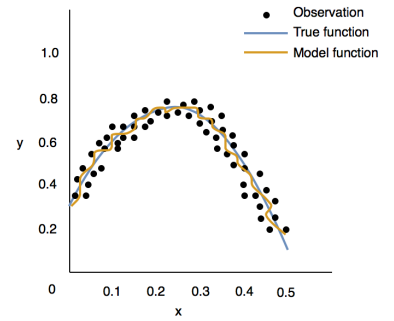
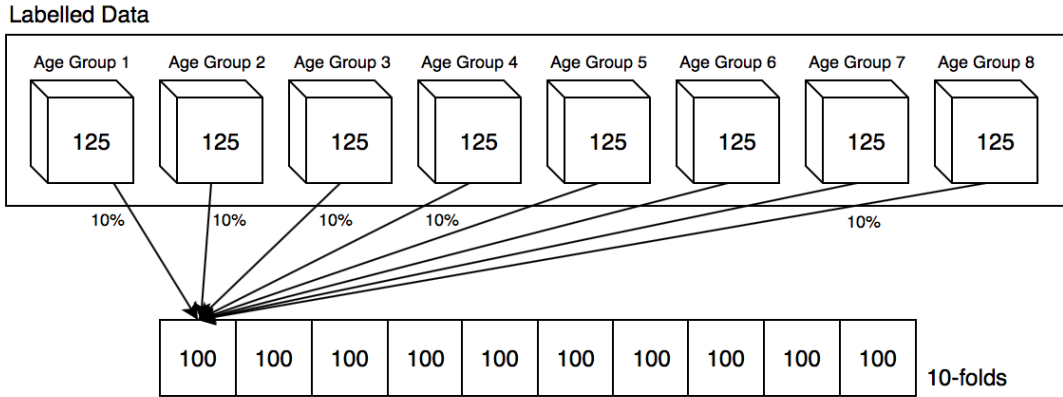


Figure 23: Model Overfitting

³ Down-sampling or up-sampling are common techniques used when there is an unequal amount of samples for the different classes.

end up with 10 subsets of 100 images that are equally distributed along all age-groups. With these subsets, we can estimate our model's accuracy with a lower bias-variance tradeoff. Since we are using 10-fold CV, we will train and test our model 10 times. For the first iteration, we shall use subset 1 as the validation set and subsets 2 to 10 as the training set, for the second iteration we use subset 2 as the test set and subsets 1 plus 3 to 10 as our training set, and so on (as shown in Fig. 21). For each iteration we use the misclassification error rate to obtain an accuracy value and we finally average the 10 accuracy rates to obtain the global accuracy of our model when solving age group classification, given the FG-NET data set. Hence, we have now estimated the prediction error of the model and have an idea of how well our model performs in solving such a problem. It is important to note that cross-validation is *just* an estimation method and when using our model in real-life applications we do not apply CV but rather train our model with all the data we have.



As concluded by Varma and Simon (2006), cross-validation is well implemented when everything is taken place within every CV iteration (including preprocessing, feature-selection, learning new algorithm parameter values, etc.), and the least bias can be achieved when using nested CV methods.

Figure 24: Stratified 10-fold cross-validation on 1000 labelled images of 8 different classes

Dimensionality Reduction

As the volume of data collected increased, several machine learning approaches have been formulated to identify patterns and perform predictions from this data. Apart from their large volume, these datasets constitute of a large number of variables. This high dimensionality within the data presented challenges such as excessive computational costs and increased memory requirements (Bolón-Canedo et al., 2015).

For these reasons, prior to applying any of these data mining techniques, pre-processing mechanisms are normally applied on the data. These methods include dimensionality reduction which is the process in which the large number of input variables are reduced to a smaller set of features (Sorzano et al., 2014).

Dua and Saini (2009) mention *feature selection* and *feature extraction* as two possible methods which reduce the dimensionality of the data. Feature selection involves selecting only the relevant features of the dataset, whilst ignoring the remaining features (Dua and Saini, 2009).

On the other hand, feature extraction techniques are applied to construct a feature vector with lower dimensionality. Linear Discriminant Analysis (LDA) and Principal Component Analysis (PCA) are two common techniques applied for feature extraction, in which a linear transformation matrix is used to project the original features to a new feature space (Wang and Paliwal, 2003).

Random Forests

Breiman (2001a) introduced Random Forests (RF) as an algorithm for *classification* and *regression* problems. However, this method is also useful in determining feature importance or irrelevance, thus widely used for variable selection (Genuer et al., 2010).

The concept behind RFs involves the iterative construction of a number of binary *decision trees*, which have a low degree of correlation (Genuer et al., 2010). In each iteration, a random sample is extracted from the training sample on which the decision tree will be trained. During training, each split involves randomly selecting k features from the original feature vector P , where $k < P$. Data is split using the best feature out of the selected k features. Once the separate decision trees, also referred to as random forests, are constructed, each tree can be used to predict outcomes for new samples. In classification problems, these outcomes are then averaged to obtain the final prediction (Kuhn

and Johnson, 2013).

Díaz-Uriarte and Alvarez de Andrés (2006) perform feature selection using a RF approach to select a smaller set of features that can still provide effective predictions in classification or regression problems. In their strategy, variable importances are initially calculated within each iteration. The variables with the smallest importances are eliminated and a new forest is then constructed using the remaining high importance variables. Finally, the prediction error of each forest, also referred to as the out-of-bag (OOB) error rate, is calculated. The features belonging to forests with the smallest error in prediction, are selected as the final variables (Díaz-Uriarte and Alvarez de Andrés, 2006).

Strobl et al. (2007) discuss that RFs are not ideal for feature selection when the features to be used for prediction constitute of a mixture of categorical and continuous values. Moreover, the variable importances measures are unreliable in cases when the number of categories for categorical predictors vary in the sample set.

Genetic Algorithms

Genetic algorithms (GA) are a commonly used technique in feature selection, which aims at minimizing the feature space by eliminating unimportant variables and keeping those which maximize accuracy in predictions. Introduced by Holland (1962), GA works using a procedure which mimics the theory of evolution, in which a population is manipulated to generate offsprings which form the new generation.

In GA's, a population is made up of a set of individuals (samples from a sample space), which are represented using binary strings, and referred to as *chromosomes*. Each chromosome is composed of a set of features, which are called *genes* (Siedlecki and Sklansky, 1989). A *fitness function* is applied to each individual to determine the fittest individuals within the population. GA then applies selection to obtain the fittest individuals. Pairs of individuals, referred to as parents, are then constructed from the selected individuals, and crossover is performed. Crossover involves generating offsprings from the identified parents by exchanging information amongst the parent pairs. Mutation, which is applied to increase the diversity within the population, entails that one or more bits are randomly flipped to generate a different offspring. GA repeatedly perform this process, also referred to as a generation, until a stopping condition is met (Chaikla and Yulu Qi, 1999).

In feature selection, the individual considered as the fittest from the new population obtained is used to determine the features with the highest importance. Genes with value '1' in the obtained chromosome are considered as the features to be used for predictions (Sivanandam and Deepa, 2007).

Principle Component Analysis

Jolliffe (2002) introduces PCA as the technique which aims "to reduce the dimensionality of a data set consisting of a large number of interrelated variables, while retaining as much as possible of the variation present in the data set." PCA aims to transform a set of correlated variables P into a set of uncorrelated k variables, where $k < P$ and the k variables maintain the majority of the information in P .

Prior to applying PCA, the dataset is adjusted such that the means of the original variables are subtracted from the observations, to produce a dataset with mean zero (Jackson, 2005). PCA works by initially constructing the *covariance matrix* of the data (Guan and Dy, 2009). A high covariance value indicates that the variables are correlated, whereas a zero value indicates that no correlation exists between the variables. Moreover, the covariance between x and y is equal to the variance of x , when $x = y$.

The variance between features is indicated by the diagonal values of a covariance matrix (Figure 25), whilst the off-diagonal values refer to the covariance between the features. High values in the off-diagonal terms indicate that high redundancy exists, and thus the covariance matrix must be diagonalized to minimize the correlations between the features.

The resultant diagonalized covariance matrix is used to calculate the *eigenvalues* and *eigenvectors*. The eigenvectors refer to the direction in which the data varies, and these are ordered such that the ones with the highest eigenvalues are chosen as the principal components of the data. Eventually, the selected vectors are kept to form a *feature vector*, which will be used to construct the new dataset with lower dimensionality, as in Equation 42

$$\text{NewDataset} = \text{FeatureVector}^T \times \text{AdjustedData}^T \quad (42)$$

Equation 42 multiplies the transpose of the feature vector by the transpose of the mean adjusted data, to obtain the original data with smaller dimensionality (Jackson, 2005).

Multi-dimensional scaling

Multi-dimensional scaling (MDS) is another dimensionality reduction technique which similar to PCA seeks to obtain a data set with lower dimensions. MDS aims at finding a configuration in which the distances between points on a dimensional space are as close as possible to the dissimilarities between the points. Different techniques of MDS exist which differ in the way that points are matched to dissimilarities. In *Metric MDS*, "dissimilarities between objects will be in numerical and distance". Classical scaling is one such technique which aims at finding a configuration in which the dissimilarities are equal to Euclidean distances. Another technique is least squares scaling in which the dissimilarities are transformed into distances using a function,

$$\begin{pmatrix} \text{cov}(x, x) & \text{cov}(x, y) & \text{cov}(x, z) \\ \text{cov}(y, x) & \text{cov}(y, y) & \text{cov}(y, z) \\ \text{cov}(z, x) & \text{cov}(z, y) & \text{cov}(z, z) \end{pmatrix}$$

Figure 25: Covariance Matrix.

and the configuration space is obtained by minimising this function. On the other hand, *Non-Metric MDS* seeks a transformation which preserves the rank order of the dissimilarities (Cox and Cox, 2000).

Linear Discriminant Analysis

LDA is a supervised dimensionality reduction technique which produces a lower dimensionality feature space in which a feature vector belonging to a class is distinguishable from that of the other classes (Sharma and Paliwal, 2015). Figure 26 depicts the projection of a two-dimensional feature vector to a one-dimensional feature space. C_1 , C_2 and C_3 denote the three different classes being used within the feature vectors. By projecting the feature vectors onto line \hat{W} , which represents one dimension, it is noted that a strong relationship exists between the feature vectors of the three classes. LDA seeks an orientation which strongly separates the feature vectors of one class from those belonging to other classes. This orientation can be obtained by rotating \hat{W} , resulting in projection W , where the feature vectors of the classes are highly separated. Algorithmically, Fisher's criterion is maximized to obtain the required one-dimensional feature vector W .

Feature extraction using LDA (Sharma and Paliwal, 2015; Huang et al., 2002) works by initially calculating the separability between the mean values of each class.

$$S_B = \sum_{i=1}^c n_i(\mu_i - \mu)(\mu_i - \mu)^T \quad (43)$$

This is referred to as the *between-class matrix* and is represented by Equation 43, where c is the number of classes, n_i is the number of samples for the i^{th} class, μ_i is the mean for the i^{th} class and μ is the total mean of all samples.

$$S_W = \sum_{j=1}^c \sum_{i=1}^{n_j} (x_{ij} - \mu_j)(x_{ij} - \mu_j)^T \quad (44)$$

Following this, the *within-class matrix* is calculated using Equation 44, where c is the number of classes, n_j is the number of samples for the j^{th} class, x_{ij} is the i^{th} sample in the j^{th} class, and μ_j is the mean for the j^{th} class. This matrix indicates the distance between the mean and sample values for each class. Finally, a *transformation matrix* W which maximizes the between-class variance and minimizes the within-class variance should be constructed.

Fisher's criterion, shown in Equation 45 is used to obtain the maximum ratio of the between-class matrix (denoted by S_B) and within class matrix (represented by S_W). By transforming the latter into Equation 46, where λ denotes the eigenvalues, eigenvectors and the corresponding eigenvalues of W may be calculated. A lower dimensional space is constructed by selecting the eigenvectors having the highest eigenvalues.

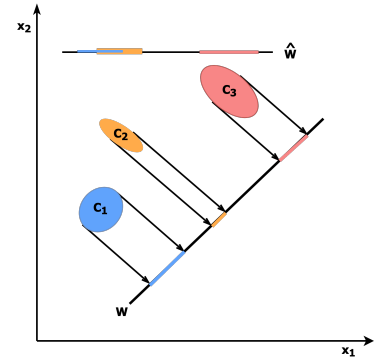


Figure 26: Projection of a two-dimensional feature vector to a one-dimensional feature space. Adapted from Sharma and Paliwal (2015).

$$\arg \max_w \frac{W^T S_B W}{W^T S_W W} \quad (45)$$

$$S_W W = \lambda S_B W \quad (46)$$

Empirical Risk Minimisation

Introduction

Empirical Risk Minimisation (ERM) is a principle of statistical theory that is used in supervised learning. Simply put, it states that for a function to make the most accurate predictions possible on an unknown true distribution, prediction errors made on the data belonging to the distribution must be minimised. To give a proper explanation of the principle some prior concepts need to be explained. The next section will discuss loss functions.

Loss Functions

A simple supervised learning scenario comprises of the following. Typically there would be one or more inputs or features denoted as x and the target output or labels denoted as y . Using the combination of features and output (x, y) , a prediction function $f(x)$ is built where, given a new input x' gives the best estimate for the value of y' , assuming that the data follows an underlying distribution that is not known. Once a set of predictions is generated, a way of measuring how good these predictions are is required. For a single data point this is simply the distance between y and y' . Repeating this process for multiple points will give us a loss function ([Wehenkel, 2018](#)):

$$L(f(x), y) \quad (47)$$

The loss function gives us the cost our prediction function ($f(x)$) which gives a clear indication of how efficient the function is at predicting the expected output. Common loss functions are squared-error loss and mean square error. This calculation of "cost" of the function can give us an indication of the risk of the prediction function. Therefore, the goal of finding the most optimal prediction function turns out to be the goal of finding the function which minimises the true risk.

True Risk

True Risk is the prediction error on the entire population of data. Unfortunately in most cases, it is impossible to collect the entire population dataset. For example, in the case of predicting the gender of a person based on their height and weight, every single person

across the globe would need to have their height and weight collected into one set of data. Since the entire population of data is unavailable, there needs to be a way of minimising the true risk, without the true distribution being known ([Vapnik, 2000](#))

Empirical Risk Minimisation

This is where Empirical Risk Minimisation comes in. When constructing the supervised learning model, the predictive function chosen is the one with the least discrepancy between predicted values and actual values. In other words, the one with the least risk. This risk is Empirical Risk and the process of finding the function that minimises this risk is called Empirical Risk Minimisation. Note the risk is empirical risk and not the true risk as the data is only subset of the entire population. In an ideal scenario the end goal would be to minimise the true risk as much as possible. Due to the fact that the data for calculating the true risk is unavailable this cannot be achieved, however it is said that the empirical risk is almost identical to true risk. Therefore by minimising the empirical risk the true risk will be minimised also.

Equation of Empirical Risk Minimisation

The following is the equation for the Empirical Risk Minimisation as proposed by [Vapnik \(1991\)](#):

$$R_{emp}(f) = \frac{1}{m} \sum_{i=1}^m L(f(x_i), y_i). \quad (48)$$

Here R_{emp} denotes the empirical risk or empirical error for a particular prediction function $f(x)$. The number of input and output pairs is denoted as m . As can be seen in the equation above, empirical risk equates to the sum average of the loss function on each pair of x and y points generated. The induction principle behind empirical risk minimisation makes the assumption that the prediction function $f(x)$, which minimises the risk R_{emp} , will result in a risk R_{emp*} which is close to its minimum.

Uniform Convergence

The convergence of the empirical risk to the true risk can only be achieved under specific conditions and bounds ([Clémençon et al., 2017](#)). These bounds are independent of the prediction function and are based on the VC-dimension of the dataset. This dimension basically measures the complexity of the prediction function. The VC-Dimension set also measures ability of the function to perform correct predictions on unseen data, using the least training data possible. A low VC Dimension indicates a high bias in the function, which will lead the function to perform poorly (especially on training data) as it will cause underfitting. On the other hand, a high VC-Dimension

which indicates that the function has a high complexity. The complexity could be high to the point that the function would memorise the data rather than identify the underlying pattern which causes overfitting. Based on this, prediction function with high complexity should be avoided. When comparing two functions which have roughly the same empirical error, the least complex one should be chosen in order to avoid the scenario of overfitting.

Problem of Empirical Risk Minimisation

The most common problem of ERM as mentioned above is overfitting. As the goal is to reduce the empirical error as much as possible, the result could be a prediction function that fits the training data perfectly. However when given unseen data this will perform poorly causing overfitting ([Shalev-Shwartz, 2014](#)).



Figure 27: Example of Overfitting

Figure 27 above is a perfect example of overfitting as a result of Empirical Risk Minimisation. The green line shows the line of best fit of the predictive function which indicates how the function would best represent the data. The orange line is the same predictive function after attempting to minimise the empirical risk to its lowest point. For the training data, the function will perform extremely well with almost no error however this is due to overfitting, resulting in the function to perform extremely poorly on the testing or real world data. For this particular case the Empirical Risk Minimisation would be giving a false representation of the true risk as it would appear to be a low empirical error when in reality it would be extremely high on unseen data.

Structural Risk Minimisation

The principle of Structural Risk Minimisation (SRM) was introduced as an effective method of tackling the problem of overfitting in Empirical Risk Minimisation. This does so by making use of the VC-Dimension mentioned previously to analyse a prediction function's complexity and by using Empirical Risk Minimisation. The aim of this minimisation principle is to find the most optimal predictive function, which can be defined as the function with a low value of empirical

risk and a low VC-Dimension (low complexity). This balance between these two bounds is the essence of Structural Risk Minimisation.

Factors affecting Empirical Risk Minimisation

From what can be seen there are several factors that indicate how good approximation of empirical risk will be. Firstly, the size of the dataset. A small dataset will most likely lead to high bias and overfitting meaning a large dataset is preferred as it will provide a better approximation of the true distribution. The underlying distribution of the data will also affect the approximation as complex and irregular distributions will cause the function to overfit as previously mentioned. Another factor is the loss function used. The loss function should be consistent with the data and not show irregular spikes in loss values at certain data points. Lastly, another factor that affects empirical risk minimisation is the prediction function. If the function considered is large and complex, it will be a much more difficult task in finding the empirical risk and the true risk.

Conclusion

In conclusion, the principle of Empirical Risk Minimisation is an effective way of minimising the empirical error of a prediction function. Under the appropriate conditions, by reducing this risk on a subset of the population data then the true risk will be reduced to the same extent. However, caution needs to be taken as to prevent the case of overfitting.

Ensemble Methods

An ensemble is a method used in machine learning to combine predictions made by multiple learning algorithms to achieve better predictive performance (Clemen, 1989; Perrone, 1993). This technique works with classification problems, for example predicting whether a company will go bankrupt (Zięba et al., 2016), and it also works with regression problems, for example predicting crude oil prices (Yu et al., 2008). The simplest ensemble technique used in classification problems is to take the maximum vote of all outputs made by each classifier. On the other hand, averaging the predictions made by each predictor is the most basic technique used for regression problems.

The general idea behind ensembles is to merge each learner's hypothesis into one with the intention of obtaining better predictions. There are various ensemble methods, some of which use a single base learner to produce homogeneous learners while others use individual learners to produce heterogeneous learners. Generally, ensemble methods are applied to supervised learning algorithms. The methods described below are the most common used ensemble techniques in machine learning.

Bagging

Bagging, also referred to as *Bootstrap Aggregating*, is a technique introduced by Breiman (1996a). In his work it was shown that classification and regression trees (Breiman et al., 1993) can have a significant increase in accuracy when combined with *Bagging*.

This method works by sampling the training dataset into multiple subsets or bags of equal parts. These subsets are sampled using replacement, meaning that an instance can be found in multiple subsets. The size of each subset is defined as a percentage of the total size of the training dataset. Once the training set is sampled, each subset is utilized to fit a different learner. A prediction \hat{y} on an unforeseen instance x is made by taking a maximum vote made by each learner as the predicted classification as shown in Equation 49. If it is a regression problem an average of the predicted values is taken as the final output as shown in Equation 50. This procedure can be better visualized in Figure 28. The idea behind this technique is to

avoid overfitting by reducing variance.

$$f(x) = \operatorname{argmax} \sum_{b=1}^B f_b(y|x) \quad (49)$$

$$f(x) = \frac{1}{B} \sum_{b=1}^B f_b(x) \quad (50)$$

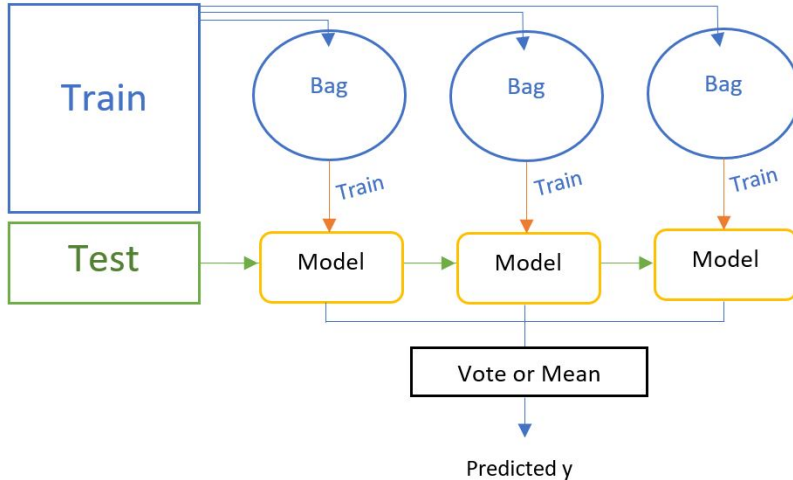


Figure 28: Dataset Bagging using 3 base learners. Each bag is a subset of the training set. Instances are selected randomly and with replacement. Each base learner is fitted with the respective bag. Predictions are then made by each base learner and the final output \hat{y} is the maximum vote or an averaged value of all predictions.

Bagging can also be applied to the feature set where subsets of the features is selected at random and used to fit the base learners. This technique is referred to as *Feature Bagging*. One popular model that utilizes *Feature Bagging* is *Random Forests* introduced by [Ho \(1995\)](#). It uses multiple *Decision Trees* which are trained using subsets of the total feature set selected at random. It was shown that when applying this method with multiple *Decision Trees*, the model was able to generalize more when classifying handwritten digits. An extension to *Random Forests* was later developed which uses both *Bagging* and *Feature Bagging* to further control variance ([Breiman, 2001b](#)).

Boosting

This technique is a variation on *Bagging* which aims to improve the learner's predictive performance by correcting the errors made by the previous learners in the ensemble, hence the name *Boosting*. The idea of having multiple *weak learners* that adapt by correcting errors was introduced by [Freund and Schapire \(1997\)](#) and this technique is called *Adaptive Boosting* or *AdaBoost*, which is the most common *Boosting* method. It was shown that this technique can reduce both bias and variance ([Breiman, 1996b](#)). *AdaBoost* does not work well when having insufficient data or very weak hypotheses ([Freund et al., 1999](#)). It also suffers when having a large number of outliers in the dataset and can significantly reduce the predictive performance ([Dietterich, 1998](#)). Variants of *AdaBoost* were developed to combat large amount of outliers in the dataset such as '*Gentle AdaBoost*' which gives less

importance to outliers (Friedman et al., 2000) and '*BrownBoost*' which uses a combination of Brownian motion, *Boosting* and repeated games to handle such cases (Freund, 2001).

Boosting works by taking a subset of the training set to train the first learner. Once the learner is trained, it is validated using the whole training set. When validation is done, each instance in the original dataset is given a weight based on the error made by this learner. A new learner is now trained on a new subset, but this time instances with higher weights are more likely to be selected. Again, this learner is validated using the whole training set but the weights for each instance is now recalibrated using a combined prediction of the two learners as shown in Figure 29. This process goes on for the number of learners specified to be used in the ensemble and the idea behind it is that each learner corrects the previous learner's errors. Once this procedure is finished, a prediction is made on the weighted mean of each learner if it is a regression problem as shown in Equation 51. If it is a classification problem, it takes a weighted vote to come up with the predicted classification as shown in Equation 52.

Initial weights set to 1

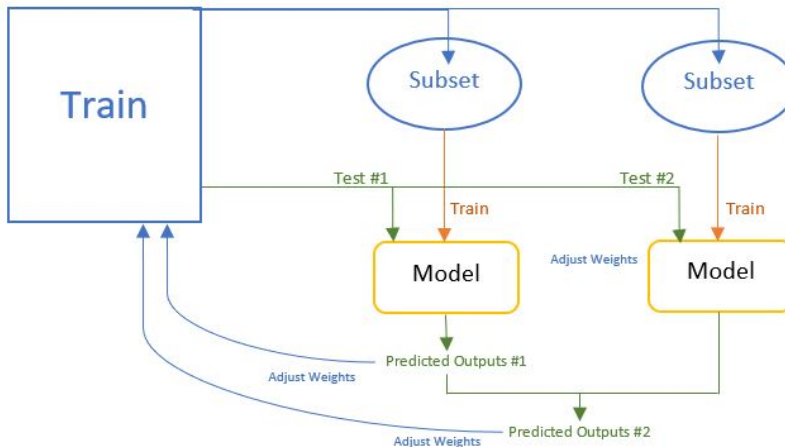


Figure 29: Boosting using 2 base learners. Initially weights are set to 1, so all instances have the same probability to be selected in the subset. The first learner is trained using the first subset and then validated using the whole training set. Weights are recalibrated depending on the errors of the predictions. On the second run, instances which had significant errors are more likely to be selected. The second learner is now trained and tested using the whole training set. This time weight are adjusted using the combined predictions of the two learners.

$$f(x) = \frac{1}{B} \sum_{b=1}^B \alpha_b f_b(x) \quad (51)$$

$$f(x) = \text{sign}(\sum_{b=1}^B \alpha_b f_b(y|x)) \quad (52)$$

This technique assumes that the learners are 'weak' and homogeneous. *Boosting* aims to build a stronger learner by combining a collection of *weak learners*. *Boosting* works in sequence and cannot be parallelized since the learners are dependent on each other. There are various other variations of boosting but one other popular implementation worth mentioning is *Gradient Boosting* 'GBM' which focuses on optimizing the loss function using boosting (Breiman, 1997).

Stacking

Stacking or *Stacked Generalization* is an ensemble method introduced by Wolpert (1992), it aims to reduce the generalization error of the combined model. Unlike the previous techniques it can be used to combine heterogeneous learners. In fact, this technique works better when the learners are different from each other and are known to be strong learners.

In *Stacking*, the dataset is first split into two; the training set and the test set. The training set is further split into K-folds like the K-fold cross validation technique. Each learner is then trained on $K - 1$ folds and validation is done on the holdout fold. For prediction made by each learner at the holdout fold is recorded and stacked together to form a new dataset. The actual value/label is also stacked at the end. This procedure continues for K times. Once all the learners have been trained using this method, a new dataset would be created with the predictions made by each learner which will be used as features. This dataset is then used to train the *combiner learner*, which is also referred to as the *meta-model* or the *aggregator model* as shown in Figure 30. After this model is trained using this newly generated dataset it will be validated using the original test set.

When using *Stacking* for classification it is advised to use class probabilities instead of the predicted outcomes as it gives a better measure of confidence for each model, thus enhancing the meta-dataset (Ting and Witten, 1999).

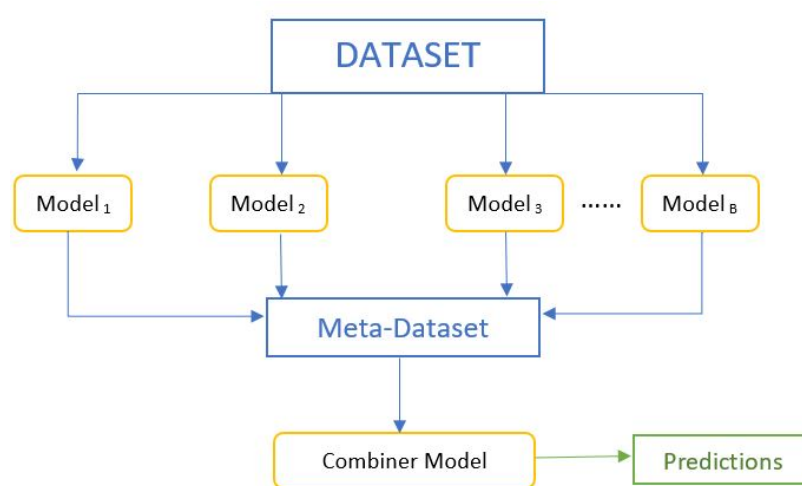


Figure 30: A visual representation of the steps required when Stacking multiple machine learning models.

The general idea behind this ensemble technique is to combine multiple different learners together, but unlike the previous methods, *Stacking* combines the models using another intermediate model which is trained on the predictions made by individual learners. *Stacking* technique was also studied to identify Higgs bosons at the Large Hadron Collider where *Stacking* outperformed *Boosted Decision Trees* (Alves, 2017).

Evaluation Methods

Although modern machine learning is exceptionally diverse and multidisciplinary, what unifies the field is standard experimental evaluation methodologies that are adopted based on the task at hand (Kibler and Langley, 1988). Evaluation is involved both in model selection, i.e. during hyperparameter tuning, and in comparing multiple models against each other or a random baseline. Ideally, evaluation is aimed at interpreting a model’s behaviour and analysing its errors (Doshi-Velez and Kim, 2017). Japkowicz and Shah (2011) outline four essentials of evaluation methodology: a test set, performance measures, error estimation, and statistical significance tests.

When evaluating machine learning models, it is common to rely on some annotated *gold standard* dataset that serves as a proxy of real-world phenomena (Manning et al., 2008). Examples of such datasets can be found in centralised collections such as the UCI Machine Learning repository (Dheeru and Karra Taniskidou, 2017) or online communities like Kaggle⁴. However, one needs to be aware that a dataset may not represent the whole domain (Japkowicz and Shah, 2011), contain annotation artefacts, or might have been constructed with vague class definitions (Hand, 2006).

While there is an assumption that training and testing sets come from the same distribution in order for a model to work as expected (Japkowicz and Shah, 2011), it is essential that the model is tested on previously unseen data. Otherwise, it yields overoptimistic results (Kibler and Langley, 1988). Error estimation techniques such as a held-out set or cross-validation are detailed on page 31, so the following is focussed on evaluation metrics and statistical tests.

Performance Measures

Stapor (2017) notes that according to the *no free lunch theorem* (Wolpert, 1996), it is unreasonable to expect that one classifier is generally better than another since there will always be problems where one outperforms the other and vice versa. Evaluation is also highly task-specific and selecting a performance measure should be done with the end goal in mind (Goodfellow et al., 2016). For example, natural language generation models used in image captioning are penalised for “hallucinating” extra information, i.e. such models are considered worse, but this aspect can be encouraged when applied to poetry generation.

⁴ <https://www.kaggle.com>

Automatic Evaluation Metrics

For supervised classification, the simplest measure is *accuracy* which is the ratio of correctly predicted examples, and its reverse metric *error rate*, i.e. the fraction of errors. More fine-grained classification metrics are described on page 19.

For machine-learned ranking algorithms that infer an ordered list of multiple outputs, standard metrics can be reported *at N* (Herlocker et al., 2004), e.g. accuracy at 5 (Acc@5) illustrating a relaxed accuracy that looks for a match among the top five outputs for each example.

For cluster analysis, despite it being unsupervised, typical evaluation metrics still require some labelled gold standard dataset. The simplest measure is *purity* that indicates the degree a model is able to cluster together instances of the same ground truth class (Equation 53):

$$p(\Omega, C) = \frac{1}{N} \sum_k \max_j |\omega_k \cap c_j|, \quad (53)$$

where $\Omega = \{\omega_1, \dots, \omega_k\}$ is the set of predicted clusters, C is the set of reference classes, N is the size of the test set (Manning et al., 2008). For other metrics used in clustering see, e.g. Friedman et al. (2001).

For regression methods, a common metric to report is the *root mean squared error* (RMSE) that characterises the discrepancy between predicted and true values (Equation 54):

$$RMSE(f) = \sqrt{\frac{1}{N} \sum_i^N (f(x_i) - y_i)^2}, \quad (54)$$

where $f(x_i)$ is a prediction for input x_i , y_i is the target value, and N is the test set size. Harrell (2015) overviews other regression metrics.

Manual Evaluation Metrics

It is not always possible to use the aforementioned automatic metrics to assess a machine learning model in a task-specific context, e.g. when there are no ground truth data available or there are yet no reliable metrics correlating with human judgement or other real-world phenomena. Therefore, *manual* evaluation is sometimes required and is typically done either by experts or via crowdsourcing when raters are performing small paid evaluation tasks (Alonso et al., 2008) using specialised platforms such as Amazon Mechanical Turk⁵. For example, assessors can be provided with Likert scales (Likert, 1932) to rate various facets of a model's output, e.g. fluency of machine translation on a scale from 0 to 5; or they can be asked to rank outputs according to some aspect, allowing comparing multiple models. There have been attempts to replicate such judgements automatically, e.g. using *adversarial* evaluation (Jurafsky and Martin, 2018) that involves training Generative Adversarial Networks (Goodfellow et al., 2014) to distinguish between human and machine-generated outputs.

⁵ <https://www.mturk.com>

Another way to overcome the lack of metrics and data is to shift the task on which a model is evaluated, i.e. conduct *extrinsic* evaluation (Belz and Gatt, 2008) that commonly involves assessment of a

model's performance in a "downstream" application rather than the training task, as opposed to *intrinsic* evaluation. For instance, if the goal is to evaluate a feature reduction mechanism, one can measure its effect on the accuracy of sentiment classification (Schnabel et al., 2015). An example of manual extrinsic evaluation is A/B studies (Hofmann et al., 2016) aimed to quantitatively measure a change in user experience before and after deploying a new machine learning model. A/B testing consists of dividing users into two groups A and B and comparing them against each other according to some metrics. A/B tests can be employed to investigate the average time it takes users to find the desired webpage depending on the search algorithm (Manning et al., 2008) or the change in time users spend on a website after introducing a new recommendation system (Davidson et al., 2010).

Statistical Significance Testing

Whenever a new algorithm is proposed, it needs to be compared against either a random baseline or previously published results for the task and dataset at hand. However, often improvement is rather incremental (Hand, 2006), so it might not be evident if it is of any importance. To decide whether a change is significant, *statistical significance tests* are encouraged, at least in case of scarce testing data (Drummond and Japkowicz, 2010).

Details about general statistical tests can be found in (Urdan, 2011), whereas Japkowicz and Shah (2011) provide recommendations on the correct use of such tests for classifier comparison. In evaluating machine learning techniques, one needs to be cautious when using standard parametric tests such as the paired *t*-test (Urdan, 2011) because of the assumptions expected to be met, e.g. equal variance of obtained metrics (Japkowicz and Shah, 2011). In contrast, *non-parametric* tests do not impose such restrictions on the population distribution which makes them more applicable but less powerful (Colquhoun, 1971). What follows is an illustration of applying a non-parametric test.

An Illustration of Statistical Significance Testing

Given two trained classifiers *A* and *B* and a set of ground truth labels of size *N* against which predictions can be compared, it is possible to construct a 2×2 contingency table such as Table 7. Assuming that accuracy is selected as a performance metric, it can be seen for model *A* it is $(CC + CI)/N = 1210/1243 = 0.973$, while for model *B* it is 0.986. Based on the accuracy alone, the improvement of classifier *B* appears rather small, so one would want to report statistical significance. A *null hypothesis* H_0 is formulated that there is no difference in accuracy between classifiers, whereas the alternative hypothesis H_1 states that these are significantly different. A *significance level* α representing the probability of wrongly rejecting H_0 is specified, e.g. 0.05. Supposing that it has been found that assumptions for parametric tests do not hold, a non-parametric alternative is used, e.g. *McNemar's test* (McNe-

		B	
		Correct	Incorrect
A	Correct	CC = 1200	CI = 10
	Incorrect	IC = 25	II = 8

Table 7: Contingency table breaking down predictions of two classifiers *A* and *B* into different cases based on comparison against the ground truth labels (correct or incorrect).

mar, 1947) with the following statistic:

$$\chi_{MC}^2 = \frac{(|IC - CI| - 1)^2}{IC + CI}, \quad (55)$$

where IC and CI correspond to the cases described in Table 7. The χ_{MC}^2 statistic from Equation 55 is compared against the χ^2 distribution table value (or a binomial distribution in case of $CI + IC < 20$) (Japkowicz and Shah, 2011) that allows to find out the probability of obtaining such χ_{MC}^2 under the assumption that H_0 is true. If this probability, referred to as p -value, is less than α , H_0 is rejected.

For the example in Table 7, χ_{MC}^2 is 5.6 and the p -value is 0.017, which is less the selected $\alpha = 0.05$. Therefore, it is said that the improvement in accuracy is significant.

Conclusions

In conclusion, each component of a machine learning evaluation framework, i.e. selecting a testing dataset and performance measures, conducting error estimation and statistical significance tests, requires careful consideration with respect to the end goal; however, rigorous evaluation can pave the way to interpretable and more reliable machine learning systems (Doshi-Velez and Kim, 2017).

Feature Selection

Feature Selection (FS) is a process that chooses an optimal subset of features according to certain criterion (Liu and Motoda, 2012). According to Sammut and Webb (2017), in many real-world applications such as: data mining, machine learning, computer vision, and bioinformatics we need to deal with high-dimensional data. Lately, the dimensionality of the data involved in these areas has increased exponentially. The trend of this growth is also reflected in the *UCI machine learning repository*⁶ as shown in Figure 31.

Guyon and Elisseeff (2003) state that the objective of FS is three-fold: improving the prediction performance of the predictors, providing faster and more cost-effective predictors, and providing a better understanding of the underlying process that generated the data. Furthermore, the aim of the FS is to select features that are *independent* of each other but *highly dependent* on the value to be predicted.

FS keeps the original feature, that is, it maintains the physical meanings of the original feature without any transformation. Furthermore, Masaeli et al. (2010) found that this property has its significance in many practical applications such as finding relevant genes to a specific disease and building a sentiment lexicon for sentiment analysis.

According to Zhao et al. (2010), FS is an optimization problem that can be split into two perspectives:

1. searching for the best subset of features,
2. principle model for: selecting, adding, removing or changing new features during the search using a certain evaluation measure.

Mainly there are three models used for FS: filter, wrapper and embedded.

Filter models

Filter models operate independently of the learning algorithm. A filter algorithm usually consists of two steps. First, each feature is given a weight then, the features are ranked and the top-ranking features are chosen whilst the others are discarded. *Selection criteria* section describes some popular measurements used to calculate feature weight.

⁶ <https://archive.ics.uci.edu/ml/index.php>

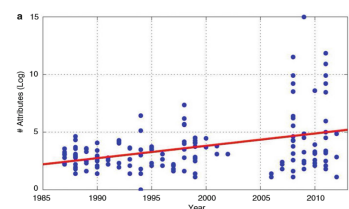


Figure 31: Growth of the number of features in the UCI ML repository. Reproduced from Sammut and Webb (2017).

Selection criteria

- *Correlation*: this measures the ability to predict the value of one variable from the value of another. In FS for classification, we look for how strongly a feature is associated with the class. A feature x_1 is preferred to another feature x_2 if the association between feature x_1 and class Y is higher than the association between x_2 and Y . In FS for clustering, the association between two random features measures the similarity between the two (Liu and Yu, 2005). According to Guyon and Elisseeff (2003), one of the simplest dependency measure function is *Pearson's correlation coefficient* defined in Eq. 56, where x_i is the i_{th} feature, Y is the output (class labels), cov is the covariance and var is the variance.

$$\rho(i) = \frac{cov(x_i, Y)}{\sqrt{var(x_i) * var(Y)}} \quad (56)$$

- *Information gain*: this is one of the most popular measures in FS due to its computational efficiency and simple interpretation (Tang et al., 2014). It is used to measure the dependence between features and labels. The information contained in label Y can be defined by *Shannon's entropy* $H(Y)$. This is defined in Eq. 57 where y_j are the labels and $P(y_j)$ is their probability. If we observe feature X then the conditional entropy $H(Y|X)$, can be defined by Eq. 58 where $P(y_j, x_i)$ is the joint probability. Finally, *mutual information* $MI(Y, X)$ between label Y and feature X can be defined by Eq. 59.

$$H(Y) = - \sum_j P(y_j) \log_2 P(y_j) \quad (57)$$

$$H(Y, X) = - \sum_i \sum_j P(y_j, x_i) \log_2 P(y_j, x_i) \quad (58)$$

$$MI(Y, X) = H(Y) - H(Y|X) \quad (59)$$

Filter models normally are very fast because usually the evaluation measurement complexity is simple and cheap, both in terms of performance and memory. Having said this, García et al. (2015) found that due to it's simplicity and low time complexity it can handle large data. On the other hand, Guyon and Elisseeff (2003) found that due to the fact that filter models score features independently of each other, the best features are not always chosen. This is because, a feature that is completely useless by itself thus ranked very low with this model, can provide a significant performance improvement when taken with others. *Wrapper models* address this limitation by evaluating subsets of features rather than individual features.

Wrapper models

The aim of the *Wrapper model* is to achieve the highest predictive accuracy possible by selecting the features that accomplish this for a fixed learning algorithm. It uses a learning algorithm as a black-box

to select the best feature subset, agreeing on a predictive measure (Kohavi, 1995). Figure 32 shows a conceptual view of a wrapper model. The *Feature Selector* component takes on the original features and uses a search strategy to generate subsets of features for evaluation. The selected feature subset uses the learning algorithm to *measure* and evaluate its performance. These steps are repeated until a *stopping criterion* is met. Typically, the stopping criterion is that refinements to the subset do not yield any performance improvements.

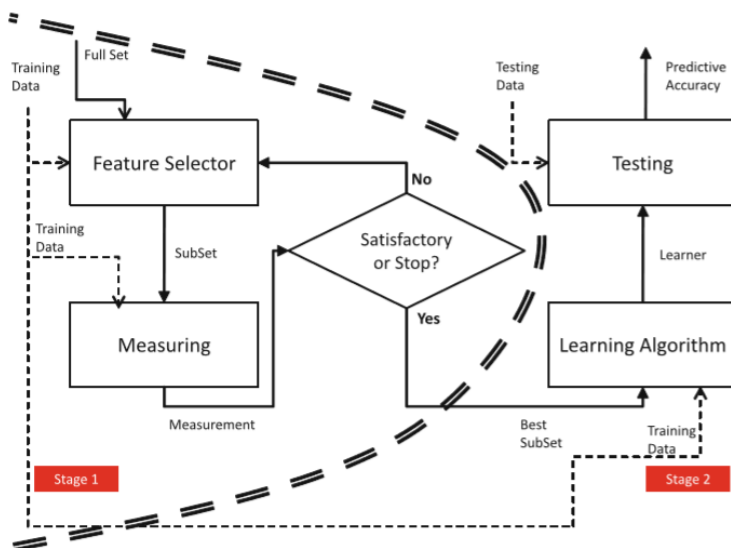


Figure 32: A general framework for Wrapper Methods of FS. Reproduced from Tang et al. (2014)

Feature Selector plays an important role in wrapper models because the order of the feature search space is $O(2^N)$ (Figure 33) making exhaustive search methods computationally unfeasible for large datasets. Therefore, search algorithms such as *Sequential selection algorithms* and *Heuristic selection algorithms*, described in section [Search algorithms](#), need to be used (Chandrashekhar and Sahin, 2014).

Search algorithms

- **Sequential selection algorithms:** These algorithms are called sequential due to the iterative nature of the algorithms. Sequential forward feature selection (SFFS), sequential backward feature elimination (SBFE) and sequential bidirectional selection are greedy search algorithms that add or remove features one at a time (Liu and Yu, 2005). SFFS initiates with an empty set and SBFE initiates with a full set where as a bidirectional search initiates a SFFS and SBFE simultaneously ensuring that the features selected by SFFS are never eliminated by SBFE. The mentioned algorithms suffer from producing nested subsets since the forward inclusion is always unconditional which means that two highly correlated variables might be included if it gave the highest performance in the SFFS evaluation (Chandrashekhar and Sahin, 2014). Sun et al. (2006) developed the Adaptive Sequential Forward Floating Selec-

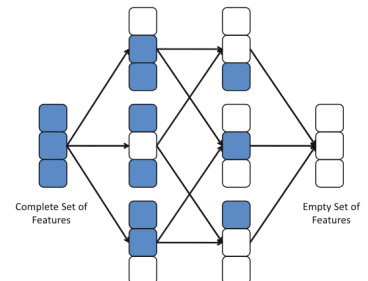


Figure 33: Search space for FS. Reproduced from Liu and Motoda (2012)

tion (ASFFS) algorithm, an adaptive version of the SFFS, to avoid the nesting effect and produce a less redundant subset than the SFFS algorithm.

- **Heuristic algorithms:** One of the widely used *Heuristic algorithm* in FS is *Genetic algorithm* (GA); a general adaptive optimization search methodology that mimic Darwinian forces of natural selection to find optimal values of some function (Chandrashekhar and Sahin, 2014). The algorithm starts by creating a set of candidate solutions referred to as chromosomes which form the first population. For each chromosome the fitness value is calculated. The fitness value is randomly impacted mainly by the *crossover* and *mutation* functions. Then based on the Darwinian principle of 'survival of the fittest' referred to *offspring*, the chromosomes with the best fitness values are combined randomly to produce the next population. This process is repeated again until acceptable results are obtained. For FS, a simple GA approach is to encode features in the chromosome (e.g. the chromosome 01001000 could mean that the 2nd and 5th features are selected), and the fitness value can be some measurement of model performance (De Silva and Leong, 2015). Huang and Wang (2006) used a chromosome with parameters of a *Support Vector Machines* SVM and the resultant solution provided the best features and the appropriate SVM parameters.

According to Sammut and Webb (2017), wrapper models generally achieve better recognition rates than filter models since the learning algorithm is involved in the selection of the subset, thus any bias introduced by the chosen algorithm is removed. The drawback of this model is that, wrapper models must train a classifier for each feature subset thus the method might become unfeasible.

Embedded models

Some learning algorithms have in-built mechanisms to deal with feature selection (Guyon and Elisseeff, 2003; Sammut and Webb, 2017). This makes the selection of the features more efficient than the wrapper method and is specifically designed for the algorithm of choice. Another improvement on the other methods is that whilst the filter and wrapper methods select the features as part of the pre-processing of the data, the embedded models cater for feature selection as part of the learning process. One such example is the use of *L₁-norm* regularisation in (SVM) (Guyon and Elisseeff, 2003; De Silva and Leong, 2015). Here the coefficients of less important features are reduced to zeros thus eliminating them. *L₂-norm* regularisation can also be used for feature selection but was found to be less effective than *l₁-norm* regularisation (De Silva and Leong, 2015). *Decision trees* also have a feature selection strategy embedded within the algorithm (García et al., 2015).

Long Short-Term Memory

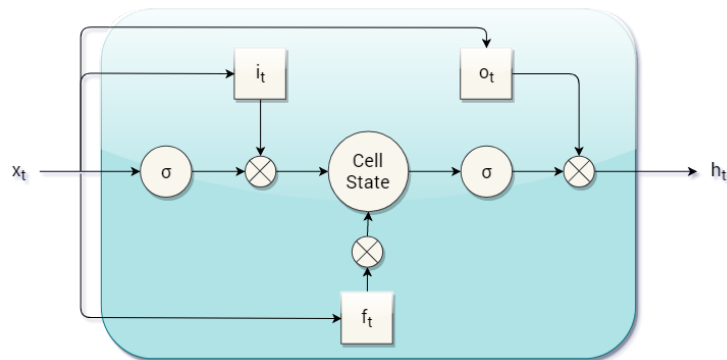
Some problems in Machine Learning, especially NLP-focused tasks, generally require some sort of **context** from previous iterations to form a more comprehensive understanding of the problem it is being given.

When facing context-heavy sentences such as, “I am from Malta, and I speak *Maltese*”, most models became unable to forge enough contextual clues in order to determine the nationality despite such clues, or **long-term dependencies**, being given earlier in the sentence (Bengio et al., 1994).

In order to solve this problem, Hochreiter and Schmidhuber (1997) proposed the Long Short-Term Memory network, or LSTM.

LSTM Architecture

This module is a type of Recurrent Neural Network (RNN), that is, a repeated chained module, with a gradient-based unit applied. The main difference is that the repeated module itself is different, with the LSTM module being composed of a series of gates named the **input gate**, **hidden gate**, **forget gate**, and **output gate**, as described in Figure 34.



The traversal from one gate to another is handled through a number of **activation functions**, which have been discussed in a previous chapter. The LSTM framework uses three main functions:

σ_g - denoting the Sigmoid Function (Rumelhart et al., 1986). σ_c - denoting the Hyperbolic Tangent Function. σ_h - also denotes the Hyperbolic Tangent Function in the case of the hidden gate, and implies that $\sigma_h(x) = x$.

Figure 34: Architecture of standard LSTM module. Adapted from: Olah (2015)

In order for the LSTM module to learn, backpropagation is implemented in the form of the Constant Error Carousel (CEC) function, which updates the hidden gate h_t through the following equations:

- h_t - the hidden state, denoted by

$$h_t = o_t \circ \tanh(c_t) \quad (60)$$

where \circ implies the Hadamard Product (element-wise multiplication).

- o_t - the output gate, denoted by

$$\sigma(W_{xo}x_t + W_{ho}h_{t-1} + W_{co} \circ c_t + b_o) \quad (61)$$

- c_t - the cell transfer state, denoted by

$$f_t \circ c_{t-1} + i_t \circ \tanh(W_{xc}x_t + W_{hc}h_{t-1} + b_c) \quad (62)$$

- f_t - the forget gate, denoted by

$$\sigma(W_{xf}x_t + W_{hf}h_{t-1} + W_{cf} \circ c_{t-1} + b_f) \quad (63)$$

- i_t - the input gate, denoted by

$$\sigma(W_{xi}x_t + W_{hi}h_{t-1} + W_{ci} \circ c_{t-1} + b_i) \quad (64)$$

These functions are not only the foundation of the LSTM module, but it is processed in such a way that it also reduces the Vanishing or Exploding Gradient Problem - the problem in which gradient descent begins to converge to zero or infinity, leaving the results almost unchanging in value (Hochreiter et al., 2001). They are typically trained by Connectionist temporal classification (CTC) Score functions (Graves et al., 2006), similar to how training and testing work in normal Neural Networks with the additional function of handling time-variable problems. It works by taking the input as the recorded observations, sequential labels as an output, and proceeds to estimate a probability distribution for the sequence of inputs against outputs for time. These processes take place for every LSTM module present in the global network, sending their output to the next LSTM module which can be seen in Figure 35.

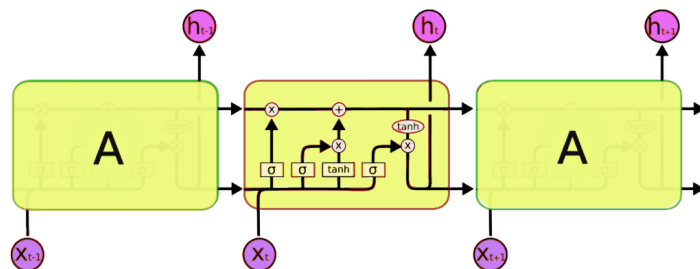


Figure 35: Typical LSTM Network. Retrieved from: [Strivastava \(2017\)](#)

Structural Variants

Since their introduction in 1997, many researchers have refined and improved this structure to suit different applications, with the following variants being the most notable amendments.

Peephole LSTM

First introduced by Gers et al. (2003), the Peephole LSTM variation is very similar to the typical LSTM structure, with the only core difference being that each cell is able to look into the current CEC values for each gate, allowing for more control into the network as shown in Figure 36.

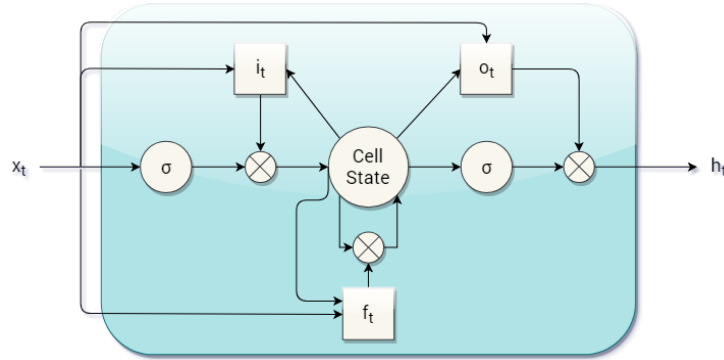


Figure 36: Peephole LSTM module variation, Adapted from Graves et al. (2013)

Convolutional LSTM

Another variation of LSTM first proposed by Shi et al. (2015), which used the LSTM's long-term dependency property in conjunction with a Convolutional Neural Network in order to process multiple subsequent image frames and retain contextual knowledge of different scenes. Figure 37 demonstrates the operation of Convolutional LSTMs.

The operations which take place are the following:

- h_t - The hidden gate, denoted by $o_t \circ \sigma_h(c_t)$
- c_t - The current cell state, denoted by $f_t \circ c_{t-1} + i_t \circ \sigma_c(W_c * x_t + U_c * h_{t-1} + b_c)$
- o_t - The output gate, denoted by $\sigma_o(W_o * x_t + U_o * h_{t-1} + V_o \circ c_{t-1} + b_o)$
- i_t - The input gate, denoted by $\sigma_i(W_i * x_t + U_i * h_{t-1} + V_i \circ c_{t-1} + b_i)$
- f_t - The forget gate, denoted by $\sigma_f(W_f * x_t + U_f * h_{t-1} + V_f \circ c_{t-1} + b_f)$

Gated Recurrent Units (GRU)

This variation, first proposed by Cho et al. (2014), is very similar to the LSTM architecture in that it also a gated network, depicted in Figure 38. It had been proposed as an alternative to LSTM's to handle problems within the same domain (i.e. Speech Synthesis, for example). The main architecture of a GRU consists of a fully gated unit, where for time $t = 0$ and output $y_t = 0$, the output vector h_t is defined as:

$$h_t = z_t \circ h_{t-1} + (1 - z_t) \circ (W_h x_t + U_h(r_t \circ h_{t-1}) + b_h) \quad (65)$$

where

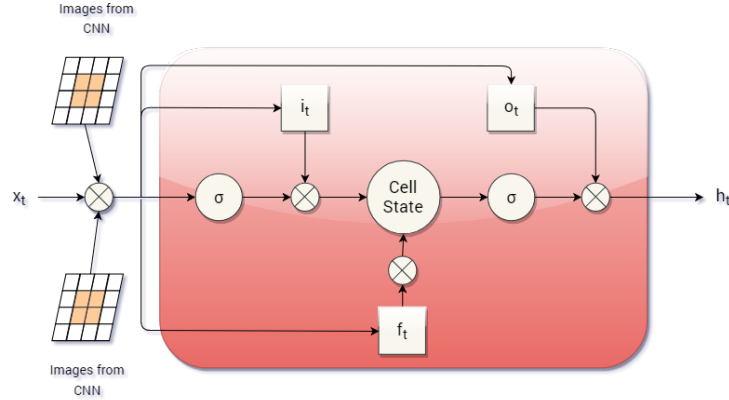


Figure 37: Peephole Convolutional LSTM module variation. Adapted from [Gers et al. \(2003\)](#)

- x_t is the input gate
- z_t is the update gate, denoted by the function $\sigma_g(W_z x_t + U_z h_{t-1} + b_z)$
- r_t is the reset gate, denoted by the function $\sigma_g(W_r x_t + U_r h_{t-1} + b_r)$

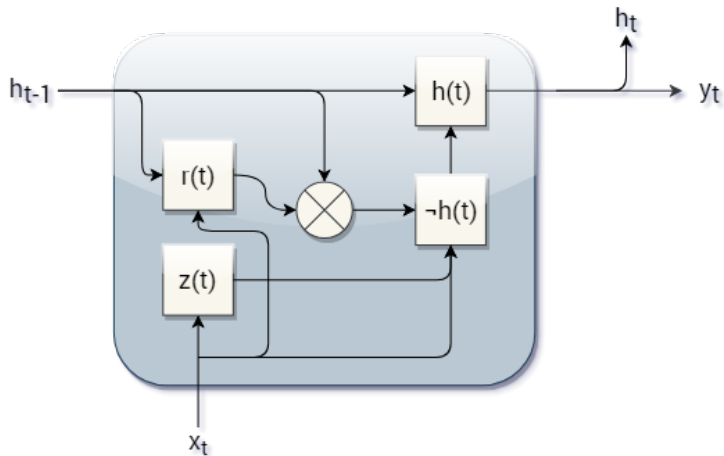


Figure 38: Structure of GRU, based on figure by [Le et al. \(2016\)](#).

In conclusion, LSTMs are an excellent deep learning tool for time-series problems such as Natural Language Processing, where sentence fragments require memory ([Graves et al., 2006](#); [Gers and Schmidhuber, 2001](#); [Schmidhuber et al., 2005](#); [Schäfer et al., 2006](#)). However, those are not the only applications of LSTMs. Other applications can involve speech recognition ([Saon and Picheny, 2017](#)), handwriting recognition ([Graves et al., 2009](#)), patient subtyping ([Baytas et al., 2017](#)), and many more applications. Handling context is very important in many domains, and as such, LSTMs have been and will continue to be used and improved on to support the ever-growing problems in Machine Learning.

Neural Networks

Neural Networks (NNs) are the product of various academic facets, the most prominent one being neuroscience as the structure of a neural network mimics the computation done by the human brain, which makes use of neurons to perform complex calculations such as object recognition from visual input of the eyes. This process is very inefficient to be performed on a computer due to its computational difficulty, which is remedied by the neural network's parallel nature. Neural Network learns with a supervised approach, supplying the model with training datasets in order to adjust the synapses to match the output. The trained model can then be used on unseen data to produce predictions (Guresen and Kayakutlu, 2011).

The way the brain trains its neurons is through experience and thus creating a learning process, and neural networks are designed to work the same way by creating synthetic nodes which will behave like organic neurons. Since this structure makes use of learnt experiences, the progress is stored in the form of weighted values at the synapse of each node (Haykin, 1994).

Types of Neural Networks

Artificial Neural Network

An Artificial Neural Network (ANN) (see Figure 41) can be defined as a collection of Machine Learning techniques that together are able to process complex inputs into meaningful spaces or classifications without any further input from the user (such as rules). The ANN is a network of layers, where every node is connected to each node of the previous and next layers. A common setup for an ANN will have an input layer, hidden layer and output layer. Neural Networks are capable of solving both regression (continuous label) and classification (discrete classes) problems, although the main purpose of using these models are for classification purposes, which is highly influenced by the number of nodes in the output layer (Devulapalli, 2015).

With regards to hyperparameters, the neural network contains parameters that require many trial and error to discover their optimal value. One of which is the number of hidden layers and nodes, as a small value can result in underfitting and the model would not have enough complexity to learn the dataset, and a large number will result in overfitting, where the model would be able to learn the dataset

by rote. The latter is a very common issue with neural networks and can be mitigated with dropout procedure. This method assigns a probability to each node during training in order to periodically deactivate it, thus eliminating the possibility of the next node to base its calculations on the deactivated node, thus eliminating overfitting (Srivastava et al., 2014a).

Another hyperparameter is the learning rate, which is used as a scale to which the model updates its weights. The learning rate should not be very large, as this could overshoot the local minimum and never converge. The ideal way is to use a decaying learning rate in order to alleviate the problems of having a large learning rate as well as small, while still keeping their benefits Jacobs (1988). One method to find out hyperparameters is to perform Bayesian Optimization, which builds a probability model on the dataset and selects the hyperparameters with the best score Eggensperger et al. (2013).

As the ANN is traversed, an activation function is fired at each step. Activation functions form the output behaviour of each of the neurons, by applying a specific function to the input nodes and producing the desired output. The activation function may contain a numerical bias, which will transform its output upwards or downwards (Jain et al., 1996). Examples of activation functions are as follows:

- **Sigmoid function:**

The Sigmoid function is an improvement over the step function as it smoothens out the steep change between the two states. This function maps the input to a value between 0 and 1, both ends being an asymptote. The Sigmoid function shown in Figure 39 is as follows (Haykin, 1994):

$$f(x) = \frac{1}{1 + e^{-x}}$$

- **Hyperbolic Tangent function:**

The Hyperbolic Tangent function (Tanh) is very similar to the Sigmoid function, with a more stable gradient and an asymptotic range between -1 and 1, thus including negative values. The function shown in Figure 40 is the following (Abdelsalam et al., 2017):

$$\tanh(x) = \frac{2}{1 + e^{-2x}} - 1$$

The traversal goes from input nodes to the output nodes, and the output will be a mapped function of the input, firing activation functions at each step. Before performing any predictions, the ANN needs to be trained with labelled data in order to adjust the synapses to their optimal values to form the mapping (Guresen and Kayakutlu, 2011).

The goal of the learning process is to minimize the output of a cost function, taking into consideration the error value. Each time the input is fed forward through the network, the error of the predicted

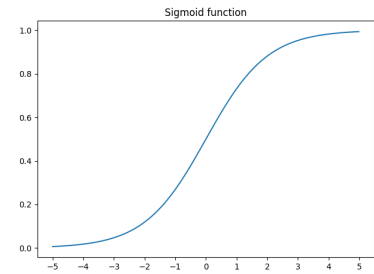


Figure 39: The sigmoid function.

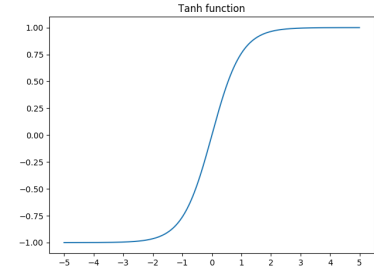


Figure 40: The Tanh function.

output is compared to the actual output and thus produce an error value. The cost function $C(n)$, in terms of error $e_k(n)$ is defined as $C(n) = \frac{1}{2}e_k^2(n)$. The error delta in terms of weight w_{kj} is calculated with the following formula: $\Delta w_{kj}(n) = \alpha e_k(n)x_j(n)$ where α is the learning rate, $e_k(n)$ is the error and $x_j(n)$ is the input. (Dawson and Wilby, 1998).

Back-propagation is the process where the biases are updated by distributing the error from the layer closest to the output inwards, applying the derivative of the activation function in the process. The synapses are updated by adding the calculated delta to the weight, formally, $w_{kj}(n+1) = w_{kj}(n) + \Delta w_{kj}(n)$ where $w_{kj}(n)$ is the previous weight and $\Delta w_{kj}(n)$ is the calculated weight difference (Rosen and Goodwin, 1994). Each layer will calculate the error based on the previous error calculations and this will cause a diminishing effect on the error value, leading towards the vanishing gradient problem. Hochreiter (1998) points out that this will cause the calculated error delta to be so small that the improvement will be nearly negligible, thus causing the model to be under-trained. One of the remedies to the above phenomenon is to implement Long short-term memory to the model. As the ANN performs epochs, the error of the synapses will converge towards the optimal value, which once reached, the model will be trained. Once the model is trained, one can perform predictions by passing an input through the network and observing the output (Taşdemir et al., 2008).

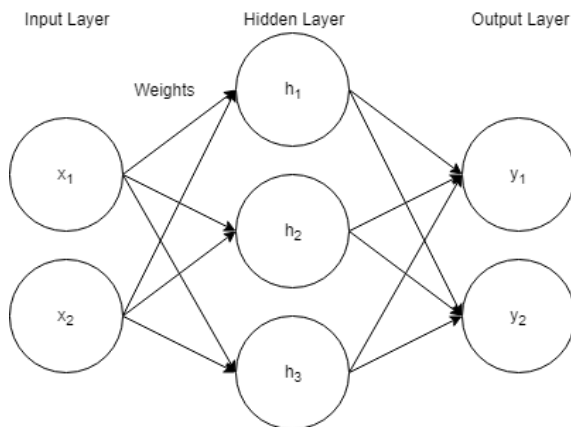


Figure 41: This diagram is a visual representation of an ANN.

Convolutional Neural Network

A Convolutional Neural Network (CNN)(see Figure 42) is a variant of ANN and are commonly used for image processing and recognition. A CNN usually is a multi-layered network, where each layer will apply a specific function to an image (Krizhevsky et al., 2012b). A CNN can use different kinds of layers, each with their own function and output, for example:

- **Features:** The images are reduced to similar patterns, where each pattern is a small subset of the image and is present in both images

(Rohrer, 2016).

- **Convolutions:** A filter is applied across every pixel to calculate the similarity between the filter (also referred to as a kernel) and the sample with a convolution function. This will output a value that represents the likeness of the feature to the image; the higher the number, the higher the similarity (Alwani et al., 2016).
- **Average/Max Pooling:** This process shrinks the images while still keeping its important features. A small window which is typically 2x2 iterates over pixels, stepping a specific number of pixels each time to possibly avoid overlapping calculations. The output of this filter is the maximum or average value inside the window. Pooling greatly helps to reduce the computational overhead of processing a large number of pixels and thus converge in less time (Fei et al., 2018; Cireşan et al., 2011).
- **Rectified Linear Units:** A Rectified Linear Unit (ReLU) is considered as an activation function, in that the output is filtered to contain only positive values. This function sets all negative values to 0 and ignores positive ones. This is used to prevent learned values from lingering close to 0 or becoming such a large negative number that it will inhibit the performance (Han et al., 2017).
- **Fully-Connected Layer:** A fully-connected layer (FC) treats the input as a singular list of nodes where each node is connected to all the previous nodes. Each node contains a weighted value which determines the contribution to a specific category when the node is fired. When the layer is activated, the nodes go through a voting process, where each node will vote to which category the input image falls under. The weights will determine which nodes will have more impact in the voting process, and the category with the bigger value of votes will classify the image (Xie et al., 2017).

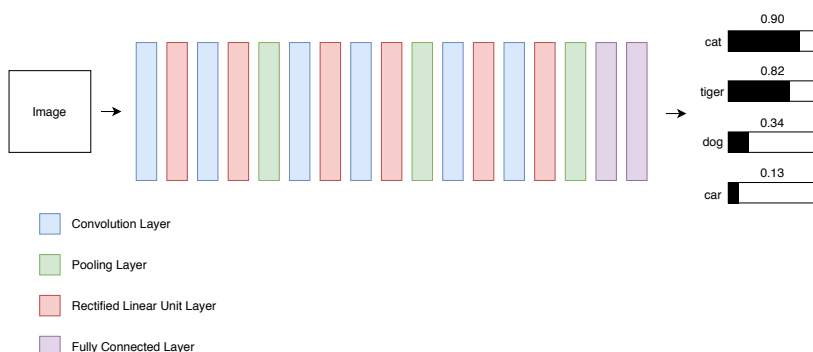


Figure 42: A diagram of a CNN with deep learning.

A CNN can have multiple repetitions of a set of layers, which will form a deep learning model with a multitude of layers that will eventually produce a final classification of the image. Training of a CNN occurs by feeding the network labelled data and adjusting the weights and features during a backpropagation step, distributing the error and converging to a set error threshold, similar to how ANNs perform their training (Iizuka et al., 2016).

Noise in datasets

As defined by Hickey (1996), noise is anything that distorts the relationship between the describing attributes and the class. Broadly speaking there are two types of noise: attribute (or feature) noise and class (or label) noise (García et al., 2013; Frénay and Verleysen, 2014). In attribute noise, errors in one or more of the attributes that describe the class distort the true representation of the data. Class noise on the other hand, is the mislabelling of instances in a dataset. Missing observations can exist in both attributes and class, and are also considered as noise (Zhu and Wu, 2004).

The example dataset in Table 8 shows the various forms of noise. Instances 5 and 6 exhibit attribute noise, with instance 5 having a missing value for attribute x_1 , and instance 6 having attribute x_2 erroneously marked as c . Instances 1, 2, 4 and 7 exhibit class noise. Instance 1 was mislabelled as o , which should read 1 . Instances 2 and 4 are contradicting instances, implying that either one was mislabelled or the attribute readings were interpreted differently. Instance 7 has a missing label.

Attribute noise is normally introduced through errors in the data collection or data processing stages, but also by corruption whilst the data are stored or transported (García et al., 2013). Class noise on the other hand can be introduced through (Frénay and Verleysen, 2014):

- insufficient information when labelling the instances;
- expert labelling errors;
- subjectivity of the classes (for example in the case of medical diagnosis where experts can classify differently);
- encoding and communication problems; and
- using a cheap labelling method (such as non-expert or automated approaches).

Unfortunately, noise is very common since reliable, noise free data are expensive and time consuming to obtain (Frénay and Verleysen, 2014). The typical error rate in a dataset is 5%. (Zhu and Wu, 2004).

Effects of noise

When learning a concept, algorithms assume and expect a correct and perfectly labelled dataset (Frénay et al., 2014). Primarily, the noise in the training set reduces the predictive ability of the inferred model (García et al., 2013; Frénay et al., 2014; Frénay and Verleysen, 2014).

#	x_1	x_2	Y
1	a	d	0
2	a	c	1
3	b	c	1
4	a	c	0
5	-	d	0
6	b	c	1
7	b	d	-

Table 8: A sample from a noisy dataset.
Note: Noise is marked in red for ease of demonstration.

This can be seen in Figure 43 that shows the effect of noise on the classification accuracy of various classifiers.

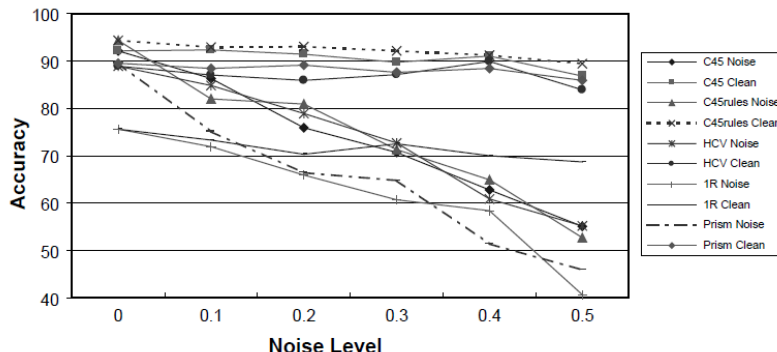


Figure 43: A graph showing the effect of noise on the classification accuracy of various classifiers. Each model was trained on a noisy dataset and on a manually cleaned dataset. These are represented as *XXX Noise* and *XXX Clean* respectively, where *XXX* denotes the classifier in question (Zhu and Wu, 2004).

The extent of the damage caused by noise depends on various factors. Class noise has been found to be more harmful than attribute noise (Zhu and Wu, 2004). The reason for this is that whilst an attribute shares the predictive capability with other attributes, there is only one label and errors confuse the learning algorithm (Zhu and Wu, 2004).

Zhu and Wu (2004) also found that not all attributes are equal in predicting the class and noise in the more important features are more damaging than others. The authors found that the higher the correlation between attribute and class the greater the impact of noise in the attribute.

Learning from a noisy dataset requires a larger training set (Frénay et al., 2014; Frénay and Verleysen, 2014) and is usually lengthier (García et al., 2013). Another cost of noise is the higher complexity of the inferred model (García et al., 2013; Frénay et al., 2014; Frénay and Verleysen, 2014).

Dealing with noise

To improve the performance of the inferred model, the effect of noise must be minimised (Zhu and Wu, 2004). There are various ways addressed in literature by which one can minimise such effects, which can be broadly classified into two categories: (i) improving the quality of the dataset (i.e. cleansing noise); and (ii) building models that are robust to noise.

Cleansing noise

Noise is usually identified by a domain expert since automatic noise identification is difficult (García et al., 2013). However, a domain expert is not always available and manual identification of noise is a lengthy process, therefore automated noise identification techniques are needed that either correct the noisy data or eliminate it.

One cannot identify noise without making assumptions (Frénay

and Verleysen, 2014). Techniques that filter out noise remove instances that appear mislabelled or that disproportionately increase the model complexity (Frénay et al., 2014). In some cases a classifier (noise filter) is trained to identify noisy instances. Other techniques aim at correcting errors or imputing missing values (Zhu and Wu, 2004).

For class noise, filtering out instances that appear noisy was found to improve results, but in the case of attribute noise, removing an instance for a noisy (including missing) attribute would not make sense, especially since other attributes may contain information that is useful for learning (Zhu and Wu, 2004). Instead, correcting or imputing the values was found to achieve better results. Techniques that deal with class noise include *ensemble filters*, *cross-validated committees filters* and *iterative-partitioning filters* (García et al., 2015).

Ensemble filters aim to identify and remove mislabelled instances in the pre-processing phase (García et al., 2015; Brodley and Friedl, 1999). The filter consists of an ensemble of m different classifiers (for example a decision tree, a 1-NN and an SVM) that are trained on the training data to act as noise filters. The training data is split into n parts and for each of the m classifiers, n different algorithms are trained. Each algorithm classifies one of the n subsets after being trained on the remaining $n-1$ subsets. If the predicted class does not match the true class, the element is marked as potentially noisy. The results of each of the m classifiers are then compared and a consensus whether an element is noisy is obtained through a voting scheme. Elements deemed noisy are removed from the training data.

Cross-validated committees filters use an approach very similar to that of ensemble filters except that the ensemble is made up only of decision trees (García et al., 2015; Verbaeten and Van Assche, 2003). It uses k-fold cross-validation to split the training data and train the base classifiers. Once again the noisy elements are identified through a voting scheme and eliminated.

Iterative-partitioning filters are used for cleaning large datasets (García et al., 2015). The training data is partitioned into manageable parts and cleansed in iterations until a stopping criterion is met. The stopping criterion is normally the percentage of noise that is tolerated.

Figure 44 depicts the approach taken by Zhu and Wu (2004) to correct attribute noise, where for each attribute a strong correlation with other attributes is sought upon which noisy instances can be predicted (through a learning model) and corrected. If a correlation is not found, corrections are based on other methods such as *clustering* or *k-nearest neighbour*.

The choice of noise filtering method should be based on the task at hand (Frénay and Verleysen, 2014). The effectiveness of the noise filter should be evaluated on a dataset that is degraded with artificial noise. The filtering precision can then be calculated through the number of correct instances that are filtered (*Type I errors* - Equation 66) and the number of incorrect instances that are not filtered (*Type II errors* -

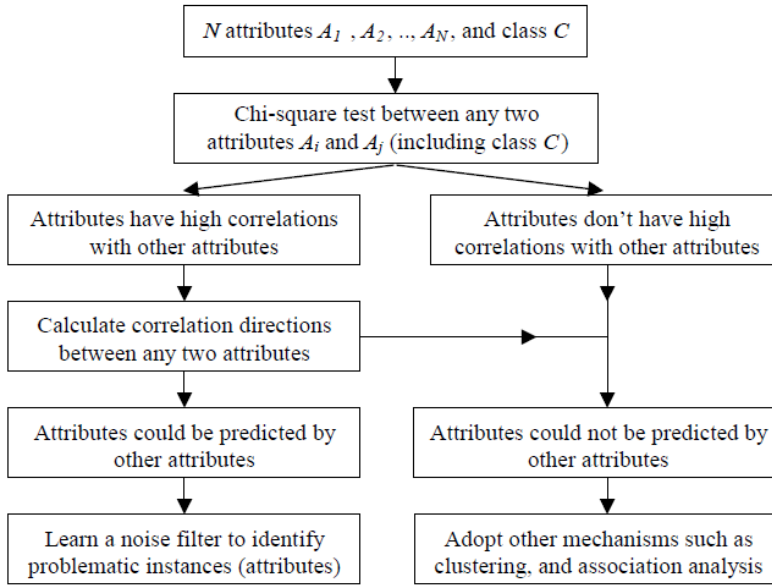


Figure 44: The approach adopted by [Zhu and Wu \(2004\)](#) to filter and correct attribute noise.

Equation 67). These are calculated as follows:

$$ER_1 = \frac{\# \text{ of correctly labelled instances which are removed}}{\# \text{ of correctly labelled instances}} \quad (66)$$

$$ER_2 = \frac{\# \text{ of mislabelled instances which are not removed}}{\# \text{ of mislabelled instances}} \quad (67)$$

Consequently, the *Noise Elimination Precision* (NER) can be calculated through Equation 68 ([Frénay and Verleysen, 2014](#)).

$$NER = \frac{\# \text{ of mislabelled instances which are removed}}{\# \text{ of removed instances}} \quad (68)$$

Noise robust models

No learning algorithm is immune to noise but some algorithms perform better than others in the presence of noise ([Frénay et al., 2014](#)). [Kalapanidas et al. \(2003\)](#) studied the noise sensitivity of ten different machine learning algorithms against various levels of artificially induced noise. The results show that classifiers are much more noise tolerant than regressors. The *linear regression* algorithm proved to be the regressor least sensitive to noise, whilst the *decision table classifier* proved to be the best performer overall. A similar study ([Nettleton et al., 2010](#)) found the *naïve bayes* algorithm to be the most robust to noise.

Online Learning Algorithms

Introduction

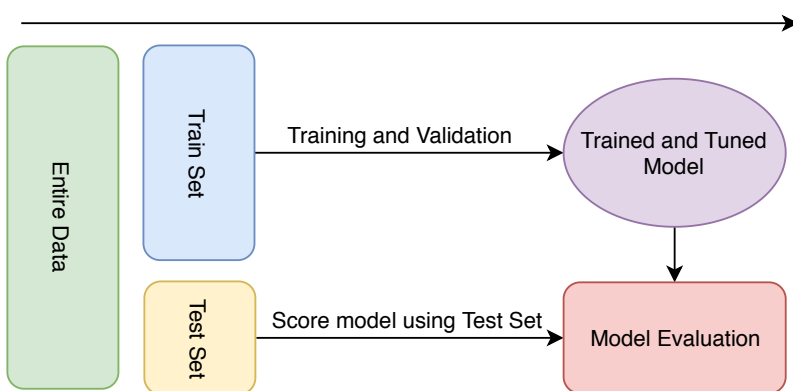


Figure 45: The traditional batch train-test machine learning approach workflow.

In the traditional machine learning approach depicted in Figure 45, we usually have some historical data to train an algorithm on for predicting some future events (Oza, 2005). However, since most data environments are dynamic and will change, the trained model eventually becomes outdated. To tackle this, we usually automate model re-training based on a timeframe (i.e. weekly or daily basis). Although this helps with keeping the model up-to-date, this is still not enough. Even if we consider model re-training on a daily basis, the model would still be at least one day late.

Furthermore, as Pagels (2018) argued, no matter how good a specific model is, it would always be an imperfect representation of the problem. Moreover, to have the best prediction for today, we cannot rely on a model with knowledge about yesterday only. Enter online learning algorithms, a family of techniques that are modelled to consume as much data available (one sample at a time), as fast as possible, while continuously learning and adapting different learning parameters. In the following sections, we will dive deeper into the subject of online learning, and its particular usage.

Why use Online Learning?

While being highly adaptable to dynamic underlying data structures since they make no statistical assumptions on the distribution of the

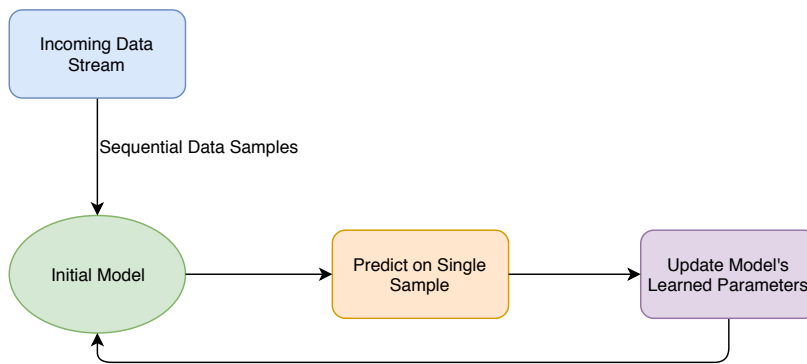


Figure 46: Basic workflow architecture of online learning.

data (Hoi et al., 2014), online learning techniques are also highly data efficient. Since online learning algorithms are only updated using the most recent data samples in the stream (as illustrated in figure 46), such data samples are no longer stored or needed once the algorithm has passed over them, maintaining a much smaller data storage (Oza, 2005). Such algorithms are also very fast since only a single pass on a smaller data set is made, in contrast to the standard approach where the optimisation function needs multiple iterations over the entire dataset. Thus, as argued by Hoi et al. (2018), online learning algorithms scale much better than the traditional approach.

As aforementioned, in offline machine learning, we load an entire dataset in memory, process it, then train a specific model, and then deploy the model into production. However, as more and more data is being generated, especially with the bright spotlight on Big Data, this methodology is proving to be more and more tedious. Some data sets are too large to fit into memory, even with distributed computing measures in place. Thus, online learning can drastically help in this scenario due to its small data storage property, especially when considered as online distributed computing and out-of-core computation (Zhang et al., 2017), which is a huge plus.

To further extract the important usability of online learning, let us, as an analogy, consider the case of an online news portal where news articles are custom and shown to the users based on which categories that respective user usually tends to click. Pretending that a terrible disaster is happening or has happened on one specific day, and the government issues a 24-hour emergency evacuation; therefore, the majority of the user-base would start clicking on this news more and more. With the traditional batch approach, even with a re-training time of 24 hours, the system would fail to push this article to users who typically do not click on domestic affairs articles (i.e. users only interested in sports or entertainment). As a result, the same data content structure will be assumed by the algorithm even though there was a drastic change of events.

In addition to this, given the same batch algorithm, after re-training in the following day, it would now start to suggest this article to a high percentage of the user-base, which by this time, such news might

no longer be relevant or applicable.

Another small application resides in the online advertising domain. With different events and occasions happening every day, especially unscheduled or unforeseeable events which go viral, ads must stay relevant all the time to ensure the highest click-rate probability, and thus, must always synchronise to the affairs of the physical world. Ads must be intelligent enough to be aware of the hidden data distribution to adapt to data morphism.

In both examples, a traditional static model will fail due to being too slow to react to the dynamic underlying relationships present in the data. This problem is more formally known as concept drift (Schlimmer and Granger, 1986). The following section will further explain the notion of Concept Drift.

Concept Drift

Concept Drift occurs when the hidden context of the data changes. For instance, weather predictions are highly dependant on the season (the context), and as the seasons change so does the weather (Widmer and Kubat, 1996). As highlighted by Krawczyk and Cano (2018); Gama et al. (2014), based on the distribution drift speed and severity, concept drift can be of four types as depicted in Figure 47:

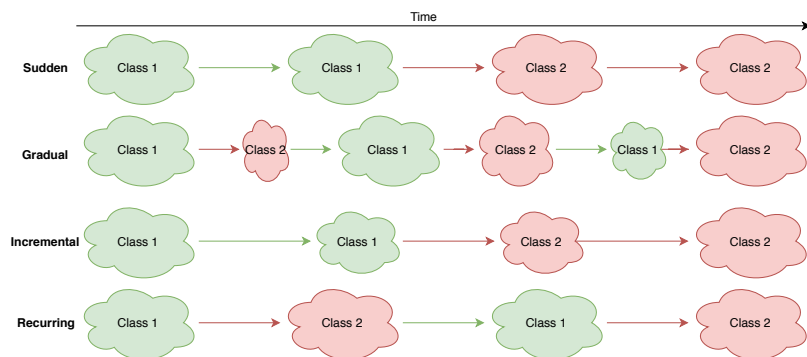


Figure 47: The four types of Concept Drift.

1. **Sudden:** *The data distribution is immediately changed to a different class.*
2. **Gradual:** *The data distribution gradually transitions by having varying proportions of the different classes mixing together over time, until it completely changes to the new class.*
3. **Incremental:** *The data distribution slowly morphs from one class to another.*
4. **Recurring:** *The data distribution periodically transitions between previous classes.*

Dealing with Concept Drift

Thus, to combat this, as the concept drifts, so must the model's transition function that maps the inputs to the outputs. Due to the constant model updates performed through online learning (sample by sample), the transition function would be dynamic and adapts to the changing distribution (Gama et al., 2014; Hoi et al., 2018; Lane and Brodley, 1998). In addition to this, another approach is to have a sliding window that shifts with the data stream. The purpose of this window, as discussed by Wozniak (2011), is to keep a set of instances that offer the best representation of the present data distribution. As newer data samples arrive in the stream, the window slides towards more recent instances, resulting in the exclusion of the oldest samples from the window. Online learning achieves this window technique through the 'forgetting rate' which sets how fast older data is discarded to make room for newer instances.

Forgetting Rate

Even though the design for most online learning algorithms is for fast execution speeds and thus adapted from less complex algorithms, implementation challenges are also present. As argued by Gepperth and Hammer (2016), this leads us to one of the most significant problems in online learning, Catastrophic Interference. The latter happens when the model abruptly forgets knowledge learnt for previous data. Most online learning algorithms have a forgetting rate parameter. This parameter allows the user to decide the speed at which the learning algorithm forgets old data; thus, how much data to retain. Moreover, the correct calibration of this rate is essential and challenging to perfect since a high value would result in catastrophic interference, while a lower value would result in the algorithm not adapting to the incoming samples in the stream. In addition to this, good initialisations are critical in this approach to steer away from slow convergence.

Conclusion

In this chapter, we introduced and discussed the sub-field of online learning algorithms concerning machine learning. Online learning is a highly useful tool that allows us to take machine learning to a whole other level by solving problems that otherwise would seem to be out of our technical ability. With the exponential importance for Big Data analytics, online learning arms us with the capabilities to process high-velocity data while also being fast to adapt to frequent changes in the data due to the ever-increasing data velocity.

Regularisation of Models

Introduction

Regularisation of Models deals with the widespread problem of overfitting in machine learning (ML) models. When a ML model is overfitting it implies that the model has been trained in such a way to perform well on the particular training data but performs badly when using test or unseen data. The overfitting model learns the pattern as well as the noise in the training data. This can be caused when the model has a high variance and when the model is highly complex with respect to the underlying data. The other extremity is underfitting where the model has a high bias. These are illustrated in Figure 48 where the underfitting model is represented by a low degree polynomial ($y = x$) with respect to the underlying data, and the overfitting model is represented by a high degree polynomial ($y = x^n$) (Taschka, 2015). High variance refers to the variability of the model output prediction for test inputs that were not used during training and represent the overfitting case, whereas high bias refers to the errors due to the ‘assumptions’ of the model that differ from the actual, since the model does not reflect the underlying relationship of the data. The challenge is to find a good bias-variance trade-off.

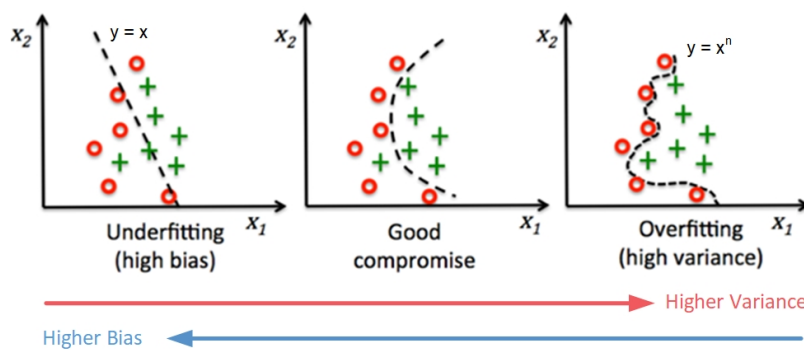


Figure 48: Overfitting and Underfitting: Challenge for a good bias-variance trade-off (Adapted from Taschka (2015)).

The *compromise* or *generalisation* challenge can be tackled using regularisation techniques. A number of regularisation techniques will be discussed in this chapter including i) L₁ regularization, ii) L₂ Regularization, iii) Dropout, and iv) Early Stopping.

L₂ and L₁ regularization

Regularization is a technique that alters the learning algorithm to add noise during the learning process in such a way that the model can generalise better. This generalisation is achieved by introducing a bias so that an overfitting model achieves bias-variance compromise as illustrated in Figure 48. The learning algorithms such as regression and deep learning typically learn using a cost or error function which is minimized such that the model with minimum error is determined. Regularisation uses a *regularisation* parameter λ to add the bias or penalty to the error function as the model complexity increases. Function 69 defines a simplified loss/cost function with regularisation.

$$\text{minimize}(\text{cost function}, c(x) + \lambda(\text{penalty function})) \quad (69)$$

Function 69 includes two terms; i) the cost function, and ii) the penalty (regularization) function, where the penalty function is constrained to be less than or equal to a constant, t (Hastie et al., 2001).

To train the model, minimisation is done on both the cost and penalty functions. The cost function, $c(x)$ depends on the training data whereas the regularisation function is independent of the data variable (x_n). Parameter λ is determined empirically or through cross validation, and it is used to control how to balance out the two terms in function 69 by balancing how much the model should learn the training set, and how much bias to add. There are two methods for regularisation that apply a bias which are termed L₂ and L₁, known as Ridge and Lasso Regression respectively. L₂ and L₁ penalize weights differently.

L₂ Regularisation

L₂ regularisation penalizes the *weight*². Therefore, considering the cost function for linear regression as an example, function 69 can be re-written as:

$$\text{minimize}\left(\sum_{i=1}^n(y_i - w_0 - \sum_{j=1}^p(x_{ij}.w_j))^2 + \lambda \sum_{j=1}^p(w_j^2)\right) \quad (70)$$

Where y_i is the predicted value from which the actual value is subtracted. The weight, (w_0) (intercept in linear regression) is left of out the penalty function. In L₂ regularisation λ is a complexity parameter that controls the amount of shrinkage (Hastie et al., 2001). The idea of penalizing by the *weight*² is also used in neural networks where it is known as weight decay (Krogh and Hertz, 1992; Moody, 1992). Krogh and Hertz (1992), claim that the “generalisation ability of a Neural Networks depend on a balance between the information in the training example and the complexity of the network”, where the complexity is related to the number of weights in the model. L₂ regularisation tries to minimize the number of weights, and therefore

make the model less complex (decrease the polynomial degree), whilst still minimizing the error.

L₁ Regularisation

On the other hand, L₁ penalizes on the |weight| (Hastie et al., 2001). Therefore the function is rewritten as:

$$\text{minimize} \left(\sum_{i=1}^n (y_i - w_0 - \sum_{j=1}^p (x_{ij} \cdot w_j))^2 + \lambda \sum_{j=1}^p (w_j) \right) \quad (71)$$

The derivative of L₂ and L₁ regularisation term would result in $2w$ and k (a constant) respectively (considering penalty function only), when computing partial derivatives with respect to the weights, (w_n). Therefore, L₂ regularisation removes a percentage from the weight, whereas L₁ subtracts a constant from the weight. This creates a significant difference from the Ridge function as it will cause some of the coefficients to be exactly zero for an appropriate value of λ (Hastie et al., 2001). L₂ regularization pushes less important weights towards zero however it does not force them to be exactly zero.

Ng (2004) considered supervised learning problems where the feature space is made up of many irrelevant features (noisy), and studied L₁ and L₂ regularization applied on logistic regression methods for preventing overfitting. L₁ regularisation cause the weights of some features to go to zero, making it highly suitable for models where many of the features should be ignored. He has found L₁ regularisation to be effective in these scenarios and concluded that L₁ regularized logistic regression can be effective even if there are exponentially many irrelevant features as there are training examples.

Since L₁ regularisation encourages the weight for meaningless features to drop to exactly zero, consequently being removed from the model, it makes this technique suitable for sparse datasets where there could be potentially considerable meaningless features or dimensions. On the other hand L₁ regularisation can have the negative affect that it zeros the weight for weakly informative dimensions.

Dropout

Dropout is another regularisation technique that is targeted for neural networks (NN) models and was proposed by Srivastava et al. (2014b). The dropout techniques adds bias or noise to the NN model in order to prevent overfitting similar to the L₁ and L₂ regularisation techniques described earlier. In order to add this bias, the dropout technique removes nodes together with their input and output connections randomly during training. The Dropout technique is illustrated in Figure 49,50.

The random dropout action is performed during the training phase only, and it creates a different thinned NN (Figure 50) for each epoch. Therefore the minimisation, through back propagation, of the NN loss function is only applied to the thinned network, and the inactive

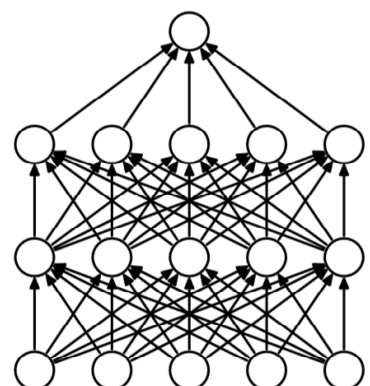


Figure 49: Standard Neural Net (Reproduced from Srivastava et al. (2014b)).

neurons do not participate in the training of that epoch. The intensity of the dropout is regulated by hyperparameter p , describing the probability of retaining a unit, where $p = 1$ implies no dropout. Dropout can be applied both for the input layer, and also on each of the hidden layers. Srivastava et al. (2014b) have determined that the typical values of p for hidden layers is between $0.5 - 0.8$, whilst for real-valued inputs 0.8 is used. Choosing an incorrect value of p can induce too much bias and may lead to underfitting.

During each training epoch a *masking* vector is created using the probability p applied on a *Bernoulli Distribution*, where its output is either 1 or 0 to determine which neurons are activated. Therefore considering a layer of n neurons, $(1 - p) \times n$ neurons would be masked at each epoch.

At the end of the training each node would have been trained a different number of times, however each epoch contributes to the same sets of weights. On the other hand, during the testing phases dropout is not utilised and all the neurons are again active. However, before testing, the weights obtained from the different thinned networks are further penalized by multiplying each outgoing weight with the probability p that was used during testing.

Srivastava et al. (2014b) et al have determined that dropout was successful in various domains including speech recognition and document classification to mention a few. The authors have concluded that the dropout method is a general technique that can be applied across different domains, however it has the drawback of extending the training time by typically $2 - 3$ times.

Early Stopping

Early Stopping is a method applied to NN where the training model is stopped before the training error is minimized (Sarle, 1995). Figure 51 illustrates a typical plot of the accuracy of model versus the epochs for the ‘Training Data’ and ‘Test Data’. At each iteration the test data is used to evaluate out the model being trained. As can be noticed although the training accuracy continues to increase as cost function is minimized, the ‘Test Data’ accuracy start dropping after a certain epoch.

Early Stopping determines the number of epochs a model is allowed to run by evaluating the *training* model after each epoch (or n epochs). The model is stopped if subsequent training results in lower performance. This marks the ‘Early Stopping Epoch’ as shown in Figure 51. Zur et al. (2009) showed that early stopping reduces the effect of overfitting but is it is not as effective as weight decay using L1 and L2 regularisation. Early Stopping is considered a form of regularisation method since it helps the model from overfitting.

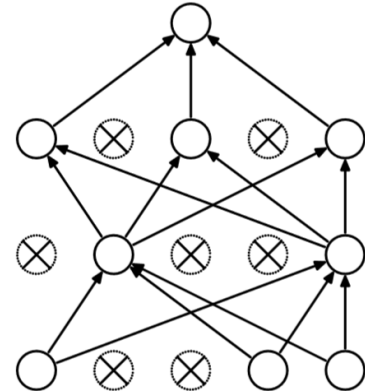


Figure 50: Thinned Neural Net after applying dropout (produced from Srivastava et al. (2014b)).

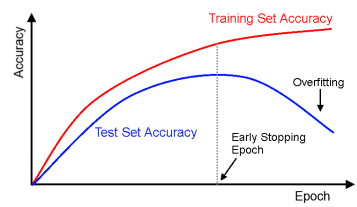


Figure 51: Training Data Accuracy Vs Test Data Accuracy

Sample Selection Bias

In traditional statistics, the algorithms assume that the data samples are being drawn in line with the same distribution, and different classes and values of data should appear with roughly the same frequency that they actually occur in the real world. However, this is rarely the case and results in the data becoming biased, meaning that the method of sample collection favours a particular type of data, skewing the distribution of the values (Cuddeback et al., 2004).

There could be a number of reasons as to why this bias occurs (Tommasi et al., 2017), such as:

1. It might be costly or unpractical to collect certain data.
 - For example, measuring spending habits of large groups of people would be cumbersome, and these people may not necessarily give accurate information.
2. The entity collecting the data only has access to data belonging to a particular class.
 - For example, if a doctor only has access to patients that are sick, a large portion of the sample would represent sick patients, whereas in relation to the total population the number of sick people would be relatively much smaller.
3. Confirmation bias, whereby people tend to recall only examples that confirm their existing beliefs.
4. Incorrect sampling techniques (Marshall, 1996), such as sampling from the top of a list instead of randomly
5. Sampling using results generated from another process.
 - For example, if a system is being trained on detecting fraudulent transactions, and some of these transactions were classified incorrectly by another process, then the model will be trained on false data.

An interesting example that tends to happen within Maltese populations is related with politics. When a political party passes an online poll regarding a particular issue, the results always tend to be in its favour. This stems from the fact that even though the poll is open to the general public, the outreach of the poll would be much more inclined to reach people within the party - they would

have subscribed to social media and news pages, and would regularly follow or check up on their articles and news.

False Information created by Selection Bias

Having selection bias within a dataset can create false information which does not exist in the actual population, and can lead to inaccurate estimates of the relationships between variables ([Cuddeback et al., 2004](#)).

For example, assume a college accepts students that either have high math skills or high science skills. Therefore if a student in this college does not have high math skills, then he must instead have high science skills. This means that a negative correlation between maths and science skills has been created, which does not represent the actual population.

How To Prevent Selection Bias

When collecting a sample of data, one should be attentive in order to prevent selection bias occurring in the dataset. This step of preventing bias is important as not only could it skew the results, but might end up rendering the analysis and conclusions gained from the dataset as useless.

Stratified Sampling

This process involves splitting the population into sub-populations before selecting, in order to ensure that an adequate number from each group is selected. For example, splitting a population into age categories and selecting a number from each category ([Krautbacher et al., 2017](#))

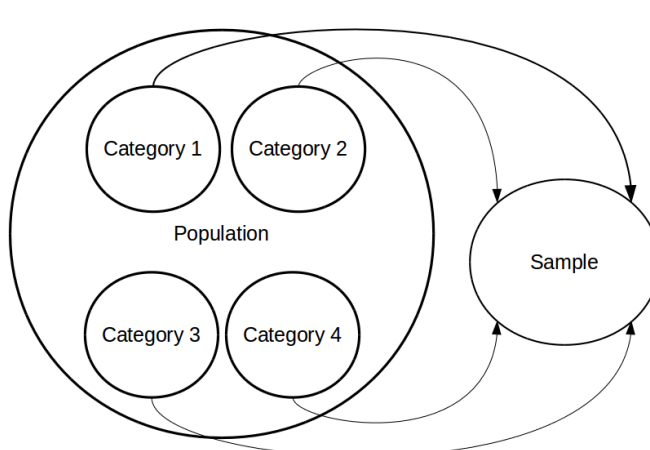


Figure 52: Stratified sampling: splitting the population into categories and selecting accordingly

Not Self Selecting

Draw from a sample that is not self selecting. If the sample is self-selected, people who are willing to volunteer may share similar characteristics (such as being more outgoing and extroverted), which will create a bias.

Analyze Dropouts

Determine factors affecting drop outs and establish that no differences exist between participants and non-participants. If those refusing to participate all belong to a certain category, this will create a bias and results need to be corrected to account for these differences ([Alonso et al., 2006](#)).

Sufficient Sample Size

Ensure a large enough sample size. If the sample size is too small, it will make it more difficult to create an accurate representation.

Detecting Selection Bias within a Dataset

Once data has been collected, or when working on a dataset gained from a different source, it is important to determine whether any sample selection bias exists within the dataset before performing analysis or drawing any conclusions. The processes mentioned below are some methods which can be useful in the detecting such bias.

Comparing Distributions

If it is possible to capture information regarding distribution about the whole population, one may then compare this with the sample population. If the distribution is drastically different, it will indicate that a sample selection bias is present.

As a simple example, if the total population contains 50% males and females, whilst your sample contains 75% males and 25% females, it will indicate a selection bias which will skew the results.

Techniques exist to allow the comparison of distributions and measuring the difference, such as Bayesian Analysis, Kolmogorov-Smirnov test or Chi-Squared test. ([Griffin et al., 2013](#))

Two Step Estimator

This method, as defined by [Heckman \(1979\)](#), comprises the use of two multiple regression models:

- One model is used to examine the interest of the study.
- The other regression model is used to detect selection bias and to statistically correct the substantive model. The independent variable could be set to represent participation. The dependent variable could represent different related statistics.

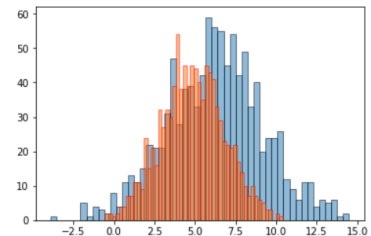


Figure 53: Comparing Distributions

Two Step Estimator Example

Imagine a survey aiming to collect information about European citizens quality of life. This would include factors such as job status, salary, level of education, country GDP, amongst others.

The first model would simply use these factors to create a model determining each citizens overall quality of life.

The second model would then measure relationships between participation and other variables. If for example a positive correlation exists between level of education and participation, then a bias in the study has occurred whereby those with a lower level of education are being underrepresented.

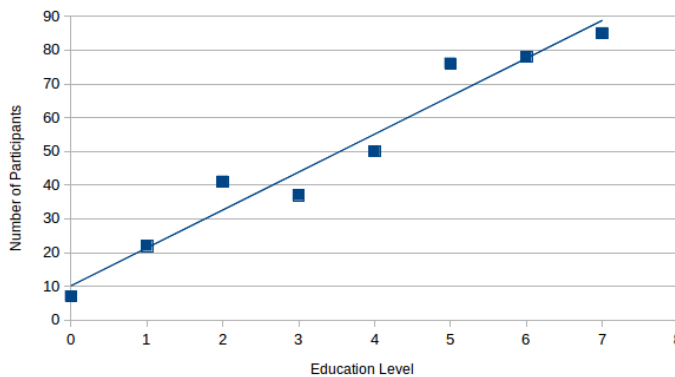


Figure 54: Two Step Estimator: Using regression models to determine correlations between participants and non-participants

How to Deal With Selection Bias

If presented with a dataset which contains selection bias but which cannot be changed or modified due to a number of reasons, then adequate measures should be taken to either mitigate the bias or account for it in the results.

Post-stratification

In an attempt to make the results more representative of the total population, higher weightings are given to the lower class. If say a sample contains 25% women, but the general population has 50%, then you could adjust the estimates by assigning women in the sample higher weights ([Holt and Smith, 1979](#)).

As explained by [Cortes et al. \(2008\)](#), the generalization error R on the newly weighted sample is defined as follows:

$$R(h) = \sum_{i=1}^m w_i c(h, z_i) \quad (72)$$

Where m is the number of samples, h is the error value, w is the weight, c is the cost function and z is the sample value.

Synthetic Minority Oversampling TECnique (SMOTE)

Another method of dealing with imbalanced classes is that of generating synthetic data that closely resembles the underrepresented class. In cases where the majority class greatly outweigh the minority class, having a larger dataset that more closely resembles the minority class will balance the results generated.

The way this process works is that new samples are created by finding the midpoint between the line segments joining the existing samples within the minority class.

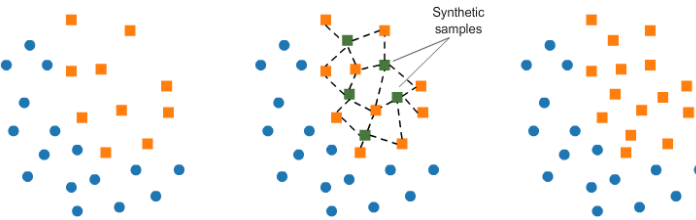


Figure 55: SMOTE: generating synthetic samples of the minority class

SMOTE may also create certain drawbacks. Since the synthetic results are generated on a small set of existing features, it may create an overfitting scenario which might not be able to correctly classify the true outlier nature of the minority class. SMOTE might also skew the distribution of the dataset which could affect certain analysis. The large addition of additional samples could also slow down learning speed. (Weiss et al., 2007)

Studies seem to have shown that using SMOTE techniques to overrepresent the minority class, as opposed to under-representing the majority class by selecting fewer, tends to perform better (Chawla et al., 2002).

Applicability Disclosure

Giving full disclosure: If it is not possible to eliminate the selection bias from the dataset, then it is important to fully disclose who exactly the results and knowledge gained from the data are applicable to.

Semi-Supervised Learning

SUPERVISED LEARNING involves a labelled dataset on which a classifier is trained. Once trained, it is then used to predict the labels of similar unlabelled data. Unsupervised Learning deals with the identification of patterns in a given unlabelled dataset.

Semi-supervised learning lies somewhere in between these two techniques, blending them together. It makes use of both labelled and unlabelled data and is often used in applications where labelled data is difficult to come by. The goal of the semi-supervised classification method is to train a classifier on both labelled and unlabelled data, with the aim of getting a better result than that provided by a supervised classifier ([Zhu and Goldberg, 2009](#)).⁷

Semi-Supervised Classification and Clustering

⁷It is also worth noting that semi-supervised methods do not exist only for classification, but also for regression problems.

SIMILAR TO SUPERVISED LEARNING, Semi-Supervised Learning can be split into two categories; Semi-Supervised Classification and Semi-Supervised Clustering. We now take a closer look at each of these two areas.

Semi-Supervised Classification

SEMI-SUPERVISED CLASSIFICATION is a classification problem which makes use of both labelled and unlabelled data in the same dataset. As is typical of such a problem, we assume that the volume of unlabelled data in the dataset is larger than that of labelled data.

In semi-supervised learning, we make use of Pseudo-Labelling ([Lee, 2013](#)) to increase the amount of labelled data upon which the classifier is being trained. The process is as follows:

1. We first use the smaller portion of labelled data to begin training our model.
2. We then use this model to predict the labels for the unlabelled data in the dataset.
3. The model is re-trained on all of the labelled data, including the original labelled entries as well as the new pseudo-labelled entries.

4. Steps 2 and 3 are then repeated for any unlabelled data left (if any).

Hence in this way, semi-supervised classification offers the same performance as a Supervised Classification, with the added benefit that the unlabelled data is automatically labelled by the classifier itself, reducing the effort needed for manual labelling in the dataset.

Semi-Supervised Clustering

SEMI-SUPERVISED CLUSTERING, also known as Constrained Clustering, can be considered as a supervised extension added to Unsupervised Clustering ([Zhu and Goldberg, 2009](#); [Bradley et al., 2000](#)).

In such a case, the dataset in question consists of unlabelled data, the same as a typical Unsupervised Clustering problem. Distinctively however, in Constrained Clustering one also finds a degree of supervised information about the data clusters inside of the dataset. Such information may contain constraints such as *must-link* and *cannot-link*, where in the former, two data elements x_i and x_j must be in the same cluster, while in the latter they must not ([Zhu and Goldberg, 2009](#)). Using Constrained Clustering we aim at clustering better than a typical unsupervised clustering technique.

Method

INITIALLY, IT MAY SEEM ILLOGICAL that a semi-supervised process making use of unlabelled data can perform as good as or better than a supervised labelled solution. Unlabelled data is incapable of providing a relationship between an element x and a label y , which is what a Supervised Model is trained upon. What gives Semi-Supervised Learning its strength is the assumptions made between the unlabelled data and the target labels.

To examine the discussed process, let us take an example proposed by [Zhu and Goldberg \(2009\)](#). First we represent each data instance by a one-dimensional feature $x \in \mathbb{R}$. x can be in one of two classes; positive or negative.

- In a case of Supervised Learning, we are given the labelled training instances $(x_1, y_1) = (-1, -)$ and $(x_2, y_2) = (1, +)$. In such a case the best Decision Boundary would be $x = 0$, where all instances having $x < 0$ are classified as $y = -$, and those having $x \geq 0$ as $y = +$.
- Now let us consider also a large number of unlabelled instances with unknown correct class labels. We observe however, that they form two groups. Taking the assumption that instances in each class form a coherent group, we know that instances tend to center around a central mean in a Gaussian Distribution.

- Through this assumption, the unlabelled data is capable of providing us with more information. We now notice that the two labelled instances detailed in the first point are not the best examples for each of the classes in our dataset.
- Hence we take a semi-supervised estimate, which shows us that the Decision Boundary should be between these two latter groups instead, at $x \approx 0.4$.

If the assumption holds, then using both the labelled and unlabelled data gives us a more reliable Decision Boundary than the one initially proposed. As we can see from this example (displayed also in Figure 56), the distribution of the unlabelled data helps us in identifying the regions having the same label, with the smaller amount of labelled instances providing us with the actual labels.

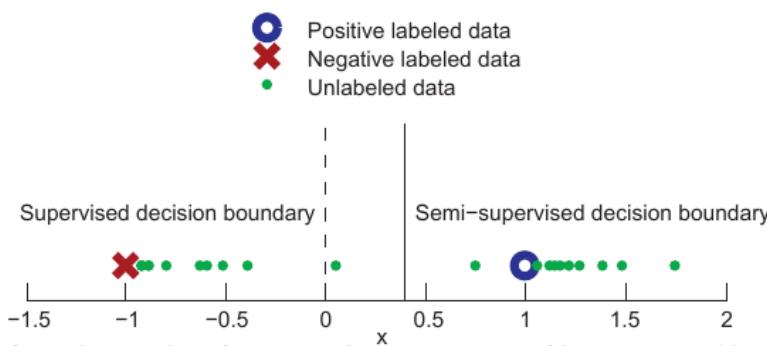


Figure 56: An example of the decision boundaries determined by a Supervised system and a Semi-Supervised system. Source: Zhu & Goldberg (2009)

Inductive vs. Transductive Semi-Supervised Learning

IN SEMI-SUPERVISED LEARNING, on finds two modes. Since in our dataset we have both labelled and unlabelled data, we have two goals to achieve, namely:

1. To predict the labels for the upcoming data instances in the test set. This is known as **Inductive Semi-Supervised Learning**.
2. To predict the labels of the unlabelled data instances inside of the training set. This is known as **Transductive Semi-Supervised Learning**.

Formally, in Inductive Semi-Supervised Learning (Zhu and Goldberg, 2009), given a training sample;

$$\{(x_i, y_i)\}_{i=1}^l, \{x_j\}_{j=l+1}^{l+u} \quad (73)$$

...our model learns a function $f : X \mapsto Y$, such that f is expected to be a good predictor on future data, beyond $\{x_j\}_{j=l+1}^{l+u}$.

In Transductive Learning (Zhu and Goldberg, 2009), given a training sample;

$$\{(x_i, y_i)\}_{i=1}^l, \{x_j\}_{j=l+1}^{l+u} \quad (74)$$

...our model learns a function $f : X^{l+u} \mapsto Y^{l+u}$, such that f is expected to be a good predictor on the unlabelled data $\{x_j\}_{j=l+1}^{l+u}$.

Limitations of Semi-Supervised Learning

SEMI-SUPERVISED LEARNING may seem as a revolutionary step on Supervised Learning; providing the same, if not better, performance, while using incomplete datasets with unlabelled data.

In reality however it is not that simple. Blindly opting for Semi-Supervised methods for any specific task can often lead to worse results than a Supervised solution ([Zhu and Goldberg, 2009](#)).

This is due to the assumption we make when handling the unlabelled data in our dataset. Since our Semi-Supervised model relies heavily on this assumption, a wrong one can lead to a significant decrease in performance and accuracy. Careful evaluation of the data is a must before committing to which type of learning algorithms to use.

Semi-Supervised Learning in Algorithms & Applications

SEMI-SUPERVISED LEARNING can be found in various practical applications, including i) Image Searching, ii) Genomics, iii)Natural Language Processing and iv) Speech Analysis. It can be integrated inside of well-known algorithms, each of which having its own advantages and disadvantages, and must be used dependently on the application in question.

A simple yet effective algorithm making use of Semi-Supervised Learning is known as the Self-Training Model ([McClosky et al., 2006; Zhu and Goldberg, 2009](#)). It is a Wrapper Method, capable of wrapping itself around other algorithms without altering their inner workings. It also comes with a crucial limitation; small errors occurring in the initial training iterations can get reinforced throughout the rest of the training.

To combat this limitation, among other improvements, one also finds other, more complex algorithms. Transductive SVMs ([Bennett and Demiriz, 1999](#)) are Support Vector Machines embedded with Semi-Supervised Learning. These methods however have difficulty scaling to large amounts of data. Graph-Based Methods ([Goldberg and Zhu, 2006; Zhu and Goldberg, 2009](#)) are some of the most used techniques. Labelled information is spread through the graph from labelled to unlabelled nodes, connecting similar observations. One also finds Neural Network solutions such as Generative Models and Deep Generative Models ([Zhu and Goldberg, 2009; Kingma et al., 2014](#)), which are capable of allowing a more robust set of features to be used than Linear Embedding used by other solutions.

Structural Risk Minimization

Structural Risk Minimization (SRM) is a technique first developed in 1974 by Vapnik and Chervonenkis (1974), used to identify the function $f(x)$ that solves a machine learning problem $y = f(x)$. SRM attempts to find a balance between training the model as accurately as possible, while not having an overly complex solution.

Consider the equation below (Guyon et al., 1992):

$$\min_f [\hat{R}(f) + \epsilon(f)] \quad (75)$$

The first part of this equation is the training loss, or what we can refer to as the expected risk $\hat{R}(f)$. The expected risk is found based on the training data, and represents the frequency of errors on our training set. Parameters are adjusted to minimize $\hat{R}(f)$ so as to obtain the best possible model for our problem. This technique is referred to as Empirical Risk Minimization (ERM). The second part of the equation $\epsilon(f)$ refers to the capacity measure of the function. This is used to quantify the complexity of our function f .

VC-Dimension

A vital topic for one to be able to understand SRM is the VC-Dimension. The theory about uniform convergence of empirical risk to actual risk describes various conditions, as well as the bounds for the rate of convergence. These bounds are based upon a measure of the capacity of the set of functions implemented by the model, also known as the VC-Dimension (Vapnik, 1992). Before looking at the definition of the VC-Dimension, it is important to understand the following definition:

Definition 1 Given a collection F of subsets of a set S , we say that the finite subset A of S is shattered provided that every subset B contained in A can be written as intersection of A with an element of F .

The VC-Dimension is defined(Vapnik, 1992) as:

Definition 2 The VC-dimension of a set of indicator functions is the maximum number h of vectors that can be shattered in 2^h ways by using functions within that set.

It should be noted that a VC-Dimension of h does not mean that all sets of h points can be shattered by a given set of functions, but there is at least one example of it occurring.

Structural Risk Minimization

Suppose m is the size of our sample, and h is our VC dimension. When $\frac{m}{h}$ is large enough, our VC confidence is negligible and hence, it can be ignored and ERM can take place. However, when $\frac{m}{h}$ is small, the VC confidence can no longer be ignored. When this is the case, we need to be able adjust the VC-dimension h of our model (Vapnik, 1992). This is where Structural Risk Minimization comes in. It is a technique that allows a trade off in accurate model fitting and avoiding overcomplexity. One may think that we would want our VC-dimension h to be a high value, as our risk $R(f)$ drops. However, while this is true, this would increase the chances of overfitting. This is why we have our complexity term ϵ , as it increases when this occurs. On the other hand, if h is too low, then we would have an issue of our $R(f)$ being too high. So the user must have the right balance of accuracy and complexity.

In mathematical terms, SRM is carried out by first selecting a family of classifiers $\{F(x, w)\}$, and then defining a structure of nested subsets of the elements of the family: $S_1 \subset S_2 \subset \dots \subset S_{n-1} \subset S_n$, where $S_i = \{f(x, w), w \in W_i\}$. Due to the structure of the subsets, their respective VC-dimension satisfies: $h_1 < h_2 < \dots < h_{n-1} < h_n$. Our goal is to find the optimal structure for our problem, denoted S^* .

To find this optimal structure, we carry out the following steps, as described by Vapnik (1992) and Sewell (2008):

1. Find the ERM for each element of the structure $\hat{f}_i = \arg \min_{f \in S_i} \hat{R}(f)$
2. Next we calculate $i_* = \arg \min_{i \in \mathbb{N}} R(\hat{f}_i) + \epsilon_i$ and return f_{i_*}

A loss function $L(f(x), y)$ is used to represent our risk function $\hat{R}(f)$, of which there are many examples, such as the hinge loss function or the ϵ -insensitive loss function, used in support vector regression. This function calculates the gap between the function $f(x)$ used as our solution, and the output y .

The generalization error of our function, $\epsilon(f)$, is a more technical proposition, explained below using the theorems 1 and 2 described by Shawe-Taylor (1997). These theorems are used to provide an upper bound for $\epsilon(f)$ with confidence $1 - \delta$. This is a very important result for us, as while we want our loss function $L(f(x), y)$ to be as small as possible, it is important that $\epsilon(f)$ is not too high.

$Er_z(s)$ is the number of errors that s makes on z . $er_\mu(s)$ represents the expected error, based on probability measure μ , when x_1, \dots, x_m are drawn independently.

Theorem 1 Let $0 < \epsilon < 1$ and $0 < \gamma \leq 1$. Suppose S is a hypothesis space of functions mapping the input space X to $\{0, 1\}$, and let μ be any probability measure on $G = X \times \{0, 1\}$. The probability with respect to μ^m that for some $\mathbf{z} \in G^m$, $\exists s \in S$ s.t. $\text{er}_\mu(s) > \epsilon$ and $\text{Er}_{\mathbf{z}}(s) \leq m(1 - \gamma)\text{er}_\mu(s)$, that is at most:

$$4\Pi_S(2m)\exp\left(-\frac{\gamma^2\epsilon m}{4}\right) \quad (76)$$

Theorem 1 is used to prove theorem 2, shown below. p_d represents the probability that S_d is the smallest class containing a consistent hypothesis. On the other hand, q_{dk} represents the probability of errors in the training set. It influences the trade off between the selected function's complexity and accuracy.

Theorem 2 Let S_1, S_2, \dots be a sequence of hypothesis classes mapping X to $(0, 1)$ and having VC-dimension i . Let μ be any probability measure on $S = X \times (0, 1)$, and let p_d, q_{dk} be any sets of positive numbers satisfying

$$\sum_{d=1}^{\infty} p_d = 1 \quad (77)$$

and $\sum_{k=0}^m q_{dk} = 1$ for all d . Then with probability $1 - \delta$ over m independent identically distributed examples x , if the learner finds a hypothesis f in S_d with $\text{Er}_x(f) = k$, then the generalization error of f is bounded from above by:

$$\epsilon(m, d, k, \delta) = \frac{1}{m} \left(2k + 4 \ln \left(\frac{4}{p_d q_d k \delta} \right) + 4d \ln \left(\frac{2em}{d} \right) \right) \quad (78)$$

provided $d \leq m$.

The function $\epsilon(m, d, k, \delta)$ provides an upper bound on the generalization error of f with confidence $1 - \delta$. The above theorem also takes into consideration the possibility of errors occurring within the training set.

An algorithm, such as the support vector machine (SVM), is used to find d which contains a hypothesis $s \in S_d$ that is consistent with our dataset $\{\mathbf{x}, y\}$. The above theorems result in the inequality shown in equation 79 below.

$$R(f) \leq \hat{R}(f) + \epsilon(m, d, k, \delta) \quad (79)$$

Examples

Consider a simple machine learning problem, where we need to find the ideal function to represent our model. As mentioned above, to carry out SRM we have two parts to our procedure, the first of which is finding the function f_i that returns to us the minimum risk $R(f)$ for each structure S_i . The definition of our risk function depends on the type of problem we are solving. For a classification problem, one may want to use a zero-one loss function or a hinge-loss function. On the other hand, for a regression problem, a Huber loss function may

be favourable. Once we have selected a suitable loss function, our next step is to look at functions to represent our capacity measure. The l_1 -regularizer and l_2 -regularizer are two examples of functions that can be used to calculate the capacity measure.

Below is the formula that needs to be solved for a support vector machine, according to SRM:

$$\min_f \left[C \cdot \sum_{i=1}^n \max [|y_i - f(x_i)|, 0] + \|\mathbf{w}\|_2^2 \right] \quad (80)$$

with the solution $\hat{f}(x_i) = C_k \cdot K(x, x_k)$, where K represents the kernel function. Notice how SVMs use the hinge-loss function as their risk function and the l_2 -regularizer to quantify complexity.

If we are to carry this out for linear regression, our formula would look something like this:

$$\min_f \left[\sum_{i=1}^n (f(x_i) - y_i)^2 + \|\mathbf{w}\|_2^2 \right] \quad (81)$$

In reality, we can use whichever regularizer we prefer, however, the l_2 -regularizer has an advantage over the l_1 -regularizer as it is strictly convex and differentiable everywhere.

Figure 57 is a diagram illustrating the general idea of SRM:

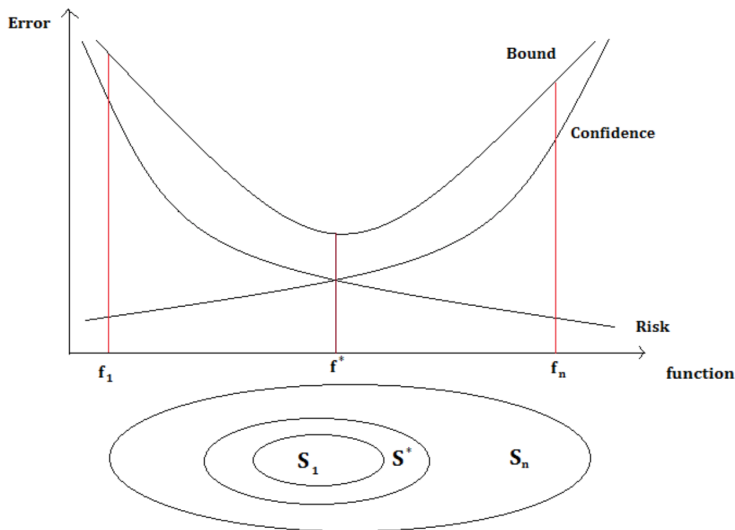


Figure 57: An illustration of the empirical risk and confidence for different functions. The optimal function is found at the minimum of the bound, listed as f^*

Synthetic Features

The problem

Training a machine learning algorithm requires inputting some form of training data. This training data comprises of all the features from which the algorithm learns from and builds a model. This input is often referred to as the training dataset.

Whilst in concept the above seems straightforward, it often transpires that the various data-points provided in the training dataset do not fit a structure that is easily understood by the algorithm. For this reason, an important pre-processing step is needed to:

1. Understand the original data well
2. Subsequently, if and where needed, generate synthetic features

If we look at the following example (Alberto et al., 2015): It contains a number of records used to train a spam / ham classifier for comments on a YouTube video.

Video	Comment ID	Author	Date	Content	Class
Psy	LZQPQhLyRh9MSZYnf8dyk0gEF9BHDPYrrK-qCc2lY8	Evgeny Murashkin	2013-11-08T17:34:21	just for test I have to say murdeco.com	Spam
Psy	z13bgdypuluhf11122rgxuhaluvalzz1os04	Zielmeek21	2013-11-28T21:49:00	I'm only checking the views	Ham
Psy	z13cx1pqgss0hlryd04cc1dxcyngljnjngk	Tasha Lucius	2014-01-19T13:25:56	2 billion....Coming soon	Ham
Psy	z12lg1v1zrmsgxnu3p23coij4aqrijxjdd1p	Holly	2014-11-06T13:41:30	Follow me on Twitter @mscalifornia95	Spam

Table 9: Sample of four rows from the Psy dataset from the YouTube comment training dataset.

Table 10 describes each feature in the original unmodified dataset.

As one can see, there is very little input the machine learning algorithm can reliably take just from using the four features described above. One could easily realise this by asking oneself the following question (in plain English):

How can I describe the components of a comment well enough to decide whether it is probably spam or ham?

One can therefore summarise this problem paradoxically as: *Having enough data to solve the problem, but very little meta-data to actually understand it and solve it.*

Ways of solving the problem

A way of solving this problem is to apply a synthetic features approach, sometimes referred to as feature engineering. This is the generation of features derived from other existing features, in a way

Feature	Description
Video	<i>The video this comment was written for. The relevance depends whether the classification model is being built generically for all videos, or a per-video specific model is also considered.</i>
Comment ID	<i>Random comment ID generated by the YouTube comment board system. This probably has no impact on the final class.</i>
Author	<i>The author / account that generated the comment. This has relevance only if this account has a lot of spam comments. If that is the case, two things should happen, none of which are directly related to the machine learning algorithm:</i> <ul style="list-style-type: none"> - <i>Maintain a blacklist of accounts that are probable spam (if a particular author often has flagged comments).</i> - <i>Block such accounts.</i>
Date	<i>The date does not directly have a huge relevance on the classification of a comment.</i>
Content	<i>The comment body definitely has a big relevance in the classification result, however, can the whole sentence be easily understood by the algorithm as it is?</i>

Table 10: Description of each feature in the original unmodified YouTube comment dataset.

that can be more easily captured or understood by a machine learning algorithm (Li et al., 2013). It is a way of generating meta-data for the existing features in the original dataset.

In essence, the idea is to look at every available feature and for each determine the following:

- Does the feature contain more than one feature within it? If so, try exploding it into sub-features and test.
- Does the feature contain too little information for any relevance, but could benefit from adding some context to it? If so, attempt at looking at other features that might be related, and produce new features as a result, and test.
- For each of the above, the original feature(s) might not be relevant anymore and be entirely replaced by the newly generated synthetic features instead.

What or how an explosion of features or a composite of features is generated depends on the very specific nature of the components involved and there is no generic formula behind it that works without some additional specificity. For example, two pairs of geo-coordinates probably qualify in giving a distance feature, however the formula applied here is specific to the geo-coordinates domain.

There is not a one-size-fits-all approach but rather it is more of an iterative approach with new synthetic features being outputted per iteration, following which one then assesses whether it is enough to generate a reliable machine learning model from the new features or not.

Following below is a practical example of this technique, using the dataset described at the introduction of this chapter.

Analysing each feature

- Video

- The video name / ID could be useful if a per-video classifier is also generated over and above the generic one. This together with other features could have some relevance.
- Comment ID
 - This feature does not have any relevance to the outcome whatsoever. It is a unique ID, built randomly, assigned to each comment. For this reason, it is out of scope for this discussion.
- Author and Date
 - As described earlier these two features independently do not have much of a direct impact on the outcome, however a synthetic feature could be generated which might have some form of effect on the outcome: A ratio of comment count over a time period for a particular author. The idea is to make it easier for the algorithm to detect a potential pattern related to volume over a typical short period, thus the definition of time period can be assigned via testing.
- Comment
 - The comment body is not an easy feature and it could grow into a number of features, however it is the most relevant input for this spam classifier. Quite a number of features could be exported from this comment, and most of them relate to natural language processing techniques (Cormack et al., 2007). For this reason, output quality could also vary based on the language in context. Some example features that could be extrapolated:

<i>Synthetic Feature</i>	<i>Scope / Description</i>
Language	<i>This depends on the availabilities of various NLP implementations for different languages, however one could have an indication of spam / non-spam probabilities based on the comment languages for each particular video.</i>
Readability score	<i>A readability score could be calculated per comment which gives an indication on the quality of such text. An example of such a score could be the Flesch Reading Ease score.</i>
Length (excl. stop words)	<i>Very short or very long comments might have a probabilistic impact on the outcome.</i>
Presence of account tags / URLs / emojis	<i>The presence of account tags (ex. a Twitter username), URLs or emojis could increase probability of the comment being spam.</i>

Table 11: Example of possible features that can be extracted from textual comments.

Updated feature / data set

Following the synthetic feature generation described above, the updated data set used as an example here would look as follows:

Looking at the output in table 12, the effect of synthetic features can immediately be appreciated, as with such new features more meaning is given to the original dataset.

Video	Author Comments in last minute	Language	Readability	Length excl. stop words	Presence of account tags	Presence of URLs	Presence of emojis	Class
<i>Psy</i>	1	EN	94.3	3	No	Yes	No	Spam
<i>Psy</i>	1	EN	103	3	No	No	No	Ham
<i>Psy</i>	1	EN	83.3	3	No	No	No	Ham
<i>Psy</i>	1	EN	32.6	3	Yes	No	No	Spam

Naturally the above contains just a sample, and one must experiment with:

- more or less synthetic features
- a further iteration of synthetic features from the generated features
- a much bigger data-set (the example above is too small to build a reliable classifier)
- perform feature selection (such as Principle Component Analysis) to identify the features that actually matter and remove extra noise

Therefore, employing a synthetic feature approach on your dataset as a pre-processing step, should in general give you positive results.

Table 12: Updated sample of the four rows from the Psy dataset from the YouTube comment training dataset now containing the synthetic features.

Transfer Learning

Machine learning algorithms typically operate within isolated problem domains - for example, a model trained to recognize motorcycles would not aim to recognize bicycles. However, this approach is not an accurate representation of human intelligence; in fact, we know that humans can relate knowledge of motorcycles to bicycles (Aytar and Zisserman, 2011).

Unlike classical machine learning approaches, human beings instinctively learn from different sources, drawing from varied past experiences and transferring knowledge across different contexts.

Transfer Learning is the study of extending classical machine learning approaches to apply knowledge acquired from a number of source tasks, to a different but related target task, similar to the way humans learn (Thrun and Pratt, 1998).

Source tasks and target tasks can be related in different ways. For example, if a source task is a '*Dog or Cat*' classifier; we can transfer knowledge to a similar domain but a different task, such as an '*Elephant or Tiger*' classifier; or we can transfer knowledge to the same task in a different domain, such as a '*Cartoon Dog or Cat*' classifier (Torrey and Shavlik, 2009).

Transfer Learning Approaches

When labeled data is in short supply or learning is computationally expensive and time consuming, one may select a surrogate task to train for, and transfer knowledge to the intended target task. This is especially beneficial in cases where we do not have enough data, or where training and test data do not possess the same characteristics; containing a different feature space, different distributions, different data sources or even different target labels.

In fact, Transfer Learning approaches can be generalised into three categories, depending on the availability of labeled data for the source and target tasks (Pan and Yang, 2010):

1. **Inductive Transfer Learning;** applicable when data is available for the target domain. Cases where there is no data for the source domain are referred to as self-taught learning; whereas cases where data is also available in the source domain is referred to as multi-task learning.
2. **Transductive Transfer Learning;** applicable when where labeled

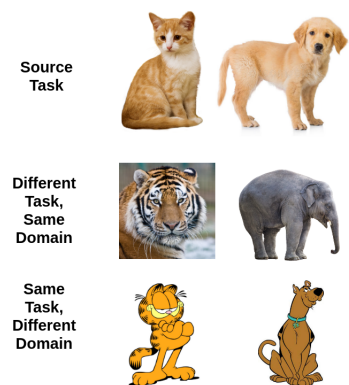


Figure 58: Transfer learning can occur across domains or across tasks. Images reproduced from commons.wikimedia.org (Tiger, Cartoon Cat), imdb.com (Cartoon Dog), pixabay.com (Cat), pnghunter.com (Elephant) and shutterstock.com (Dog).

data is only available in the source domain. Transductive learning where we assume the same task but different domains is referred to as domain adaptation, while learning a similar domain but a different task is referred to as Sample Bias Selection.

3. Unsupervised Transfer Learning; applicable when labelled data is not available neither for the source nor the target task.

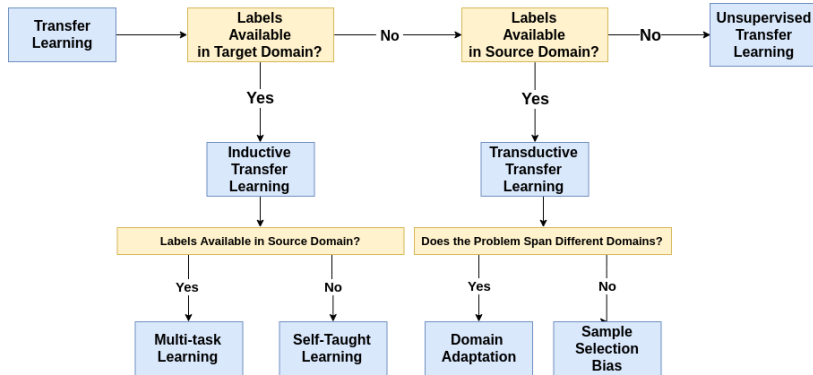


Figure 59: Transfer learning approaches vary depending on the availability of labeled data for the source and target tasks.

The above approaches vary on what data is available for the learning task - conversely, this also affects what knowledge can be transferred across tasks.

For example, it is sometimes impossible to apply a source task's data directly to a target task. Instead we can attribute selected instances or features from the source task, which are in turn re-weighted and re-distributed to fit the target task. These methods are referred to as Instance Transfer and Feature-representation Transfer approaches.

In cases where labels are available for either source tasks or target tasks (both inductive and transductive transfer learning), some forms of instance-transfer and feature-representation transfer are possible. Moreover, unsupervised transfer learning techniques are limited to feature-representation transfer (Pan and Yang, 2010).

Other examples of transfer learning include broader knowledge transfers such as parameter transfer (or model transfer), where full or parts of parameterized models are transferred from the source task to the target task; or relational-knowledge transfer, where relationships between data points within the source task are transferred to redistribute data within the target task (Cook et al., 2013). Such transfer is only possible where the target task's labels are available.

	Inductive Transfer Learning	Transductive Transfer Learning	Unsupervised Transfer Learning
Instance Transfer	X	X	
Feature-representation Transfer	X	X	X
Parameter Transfer	X		
Relational-knowledge Transfer	X		

Table 13: Transfer types by Approach

Deep Transfer Learning Techniques

The above approaches have manifested themselves in two major techniques of transfer learning within deep learning. Multi-layer neural networks are built in such a way that lower-level layers address generic features and higher-level layers address more complex and specific features. The initial layers for a group of related tasks are similar if not identical; this characteristic can be exploited to enable the transfer of knowledge between models (Yosinski et al., 2014).

Off-the shelf models, or pre-trained models, involve training one or more multi-layer networks and adapting parts of the model to another target task. The source models act as a form of pre-training, to learn shallow parts of the problem, while the target learning task is solely limited to the final layer of the model. In fact, one application of pre-trained models can be seen as a feature-selection process (Zhu et al., 2018).

Another approach is to use transfer learning to replace selective parts of the model, rather than just the final layer, this is referred to as model Fine-tuning.

With Fine-tuning techniques, we pre-train a model using a number of different source tasks, and then re-train the model for the target task; however the target training is done selectively, selecting which layers are to be frozen (and thus inherited from the source tasks) or fine-tuned (to be updated during the backpropagation process). We can define a metric to decide the learning rate for each layer, creating a variable degree of freezing and fine-tuning (Chu et al., 2016).

The success of transferability is highly dependent on source task's level of specialization, successful transfer learning projects work on generalized source tasks. As a matter of fact, transfer learning across tasks which are too dissimilar as a result of being too specific would result in a negative learning (Rosenstein et al., 2005).

Applications and Improvements

Notwithstanding savings on time and computational resources, the above techniques for transfer learning in multi-layer neural networks frequently render better results when compared to training the neural networks from scratch (Yosinski et al., 2014). This is especially the case in fields where data collection and annotation is notoriously difficult, such as computer vision and natural language processing.

The use of pre-trained models machine learning has achieved new state-of-the-art results in several tasks; within the field of computer vision, in object detection (He et al., 2017), semantic segmentation (Zhao et al., 2016), human pose estimation (Papandreou et al., 2017) and action recognition (Carreira and Zisserman, 2017); and within the field of natural language processing in text classification, question answering, language inference and conference resolution amongst others (Howard and Ruder, 2018) (Joshi et al., 2018).

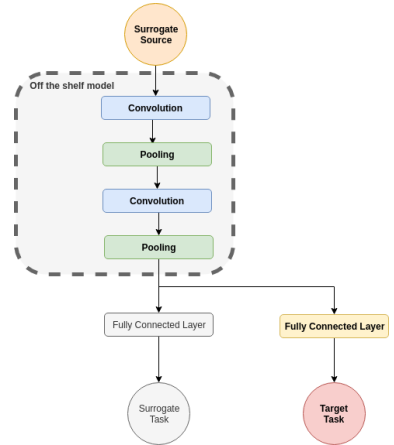


Figure 60: In off-the-shelf pre-trained models, we replace the final layer of the neural network.

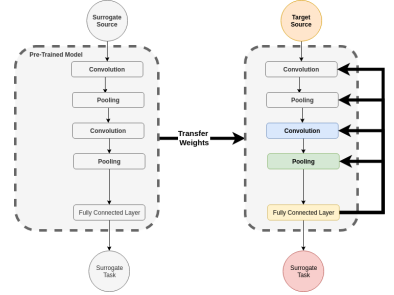


Figure 61: In Pre-trained model fine tuning, we use the source model to initialize the neural network, and fine-tune it using backpropagation.

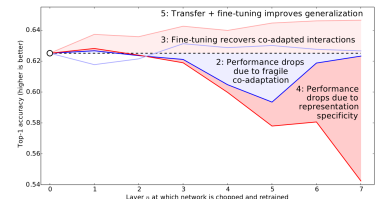


Figure 62: The performance gained from fine-tuning a neural network is only relative to the layer generalization and specificity of each distinct layer (Yosinski et al., 2014).

Receiver Operating Characteristic Curves

Introduction

A Receiver Operating Characteristic (ROC) curve is a graphical plot that presents the performance of a binary classifier when the discrimination cut-off, or threshold, is varied. ROC curves and the associated analysis area under the curve and finding optimal threshold, have proved to be very useful analysis tools in many fields of research. Numerous studies evidence that they have been found to be very effective techniques for evaluating diagnostic and predictive accuracy.

Essentially, for creating and plotting the ROC curve one needs to compute the true positive rate (TPR) and the false positive rate (FPR) at various threshold settings and plot them in a two-dimensional space. In machine learning the TPR is referred to as the sensitivity or recall which measures the proportion of actual positives that are correctly classified. The true negative rate (TNR), or specificity, measures the proportion of actual negatives that are correctly identified as such. These measures can be easily summarised in Table 14. ROC is an evaluation procedure based on the sensitivity-specificity pair of indices.

	True Class			
Predicted Class	Positive	Negative		
Yes / 1 / Positive	True Positives	False Positives	TPR=TP/P	<i>Sensitivity</i>
	False Negatives	True Negatives	FPR=FP/N	<i>1-Specificity</i>
Totals	P	N		

Table 14: Sensitivity and specificity

Consider a diagnostic test that seeks to determine whether a person has a certain disease or not. The two-class predictor yields an outcome that is labelled either positive or negative. From table 1 there are four possible outcomes from the discrete classifier: true positive or true negative if the outcome matches the actual class, false positive if the person is incorrectly classified as sick, and false negative if the person is incorrectly classified as healthy. If the test is extended to a larger sample then given a classifier and a set of instances, a 2x2 contingency table, or confusion matrix, can be constructed for the classifier with values for the four possible outcomes. Sensitivity and specificity are computed accordingly as shown in Table 14.

If one extends the test further to six classifiers, where the same metrics are extracted from the outcomes of these classifiers, one ends up with six sensitivity-specificity pairs. The latter are represented as points on an ROC plot as illustrated in figure 63 (adopted from Fawcett (2006)). A to F are sensitivity-specificity pairs representing the six classifiers used in the hypothetical test. TPR (sensitivity) is plotted on the y-axis and FPR (or $1 - \text{specificity}$) on x-axis. This plot depicts the relative trade-offs between true positives and false positives. The higher the sensitivity, the more beneficial and trustworthy the classifier is, so TPR is a measure of the classifier's benefits. A low specificity value represents a costly classifier, so FPR is synonymous with a classifier's cost. The best predictor is the one that maximises the benefits and minimises the costs.

Evaluating Classifiers

Classifier C lies on the diagonal line from $(0,0)$ to $(1,1)$ of the ROC plot. It represents a test that cannot distinguish between diseased and non-diseased states. Its TPR equals FPR. This means that classifier C is merely guessing the labels randomly. It is as accurate as predicting heads or tails in a coin toss. The diagonal line often serves as a benchmark for judging how good (or bad) a test performs. E and F lie on the right-hand side of the diagonal line with relatively low sensitivity and specificity metrics compared to the others. In general, points on this side of the ROC plot are considered worthless (or costly) classifiers.

Classifiers A, B and D are considered better predictors from the group since they lie on the left-hand side of the diagonal line with high TPR and low FPR. D is clearly the best predictor in this case because it is the closest to the top left-hand corner of the graph. The perfect predictor would yield 100% sensitivity and 100% specificity with zero false positives and zero false negatives, that is, zero cost. However this is realistically very difficult to achieve. In this way the ROC plot provides an easy comparison of classifiers and leads to choosing the one that is closest to $(0,1)$ and furthest from the diagonal line.

Consider now a continuous random variable X , denoting the width in millimetres of metal pipes from an automated production line, where X is normally distributed. An inspection test might measure the diameter of the pipes and classify a pipe as defective if the width is above a certain value. Thus given a threshold width w , the instance is labelled defective if the continuous random variable X is greater than w and non-defective otherwise. As shown in figure 64, one can idealise curves for both the number of positive and the number of negative results of the test. Since the test does not distinguish good pipes from defective pipes with 100% accuracy, the distributions overlap. The overlap area indicates where the test cannot distinguish between the two classes. In practice, one chooses a cut-off (or threshold) above which the test is considered normal. Different cut-off points are used

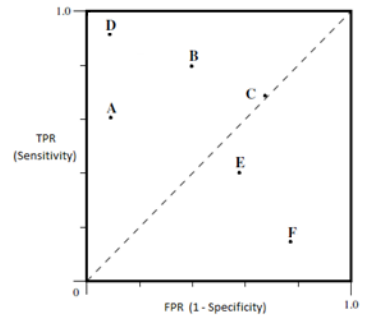


Figure 63: Basic ROC plot with five classifiers.

in different situations in order to minimise one of the erroneous types (false positives or false negatives) of the test result. The sensitivity and specificity of a classifier system depends on the particular confidence threshold that the system is using and changes inversely as the confidence threshold is changed. If the confidence threshold is made less strict, then sensitivity will increase due to more instances being classified as true positives. However, specificity will decrease because more false positives will appear. TPR and FPR increase or decrease simultaneously as the confidence interval threshold is changed. By repeating the experiment with adjusted thresholds, different value pairs are obtained leading to different positioning in the ROC space. ROC curves allow the visualisation of the trade-off between sensitivity and specificity for all possible thresholds rather than just the one that was chosen by the modelling technique.

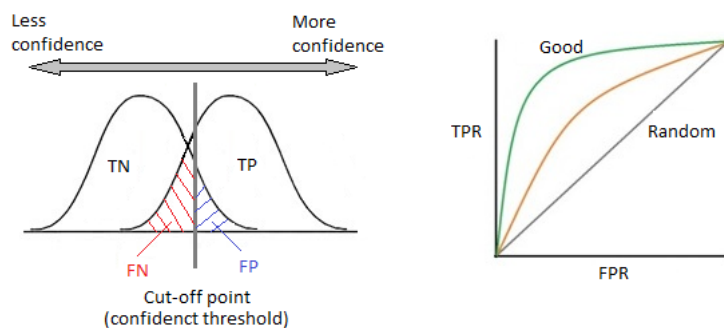


Figure 64: Left: Overlapping distributions with threshold. Right: ROC curves resulting from different thresholds.

The classifier that is able to increase the rate of detection while keeping the false alarm rate low is considered a good classifier. Figure 64 shows the ROC curves with different thresholds. As the ROC curve gets closer to the top left hand of the graph the more optimal the threshold and hence the better is the classifier's ability to discriminate between the classes.

Area Under Curve

The most common condition for choosing the optimal parameterisation is to maximise the area under the curve, AUC for short. The AUC is a widely used measure of performance of supervised classification rules. It is a measure of the usefulness, or rather the discriminatory ability, of a test in general where the greater the area the more discriminative the test is in a given situation. A model or test with perfect discriminatory ability will have an AUC of, or very close to, 1.0, while a model unable to distinguish between classes will have an AUC value of 0.5 or less.

A precise mathematical computation of the AUC is $\int_0^1 ROC(t)dt$, however since the ROC curve is normally plotted by joining small points close to each other, it can be very closely approximated by adding the area of many small rectangles next to each other covering

the whole section under the curve as shown in figure 65.

However, the true value of AUC comes when comparing two or more models or tests by reducing the ROC to a single scalar value representing the expected performance of the models. Figure 66 (adopted from Linden (2006)) provides a comparison between two predictive models. Clearly Model A yields a better predictive ability than Model B since its AUC is bigger. Therefore, if a choice was to be made between selecting one of the models, Model A would be the choice.

AUC is a desirable measure because (1) it is scale-invariant: it measures how well predictions are ranked rather than their absolute values; (2) it is classification-threshold-invariant: it measures the quality of the model's predictions irrespective of the classification threshold chosen.

The majority of studies on binary classifiers in conjunction with imbalanced datasets still use the ROC plot as their main performance evaluation method and this make it easier to compare results. However a recent study (Saito and Rehmsmeier, 2015a) shows that Precision-Recall plots provide a more accurate prediction of future classification performance although they are less frequently used. It is therefore appropriate to use the AUC metric together with other measures of evaluation depending on the nature of the dataset.

Partial AUC

AUC is closely related to another widespread statistic, the Wilcoxon statistic (Hanley and McNeil, 1982). Thanks to this relationship, AUC methods for AUC-based analyses are well developed and widely used. However one of the major practical drawbacks of the AUC as an index of diagnostic performance is that it summarises the entire ROC curve including the regions that are frequently not relevant to practical applications, such as regions with low level of specificity (Ma et al., 2013). To alleviate this deficiency while benefiting from some of the advantageous properties of the area under the ROC curve, one can use a partial area under the curve (pAUC for short), which summarizes a portion of the curve over the pre-specified range of interest. A number of approaches have been developed for pAUC-based analysis (Dodd and Pepe, 2003; He and Escobar, 2008). However, Ma et al. (2013) argue that the same features that increase the practical relevance of the pAUC introduce some difficulty to resolve issues related to arbitrariness of specifying the range of interest.

In the case of population screening, for instance, Dodd and Pepe (2003) claim that in order to avoid high monetary costs, the region of the curve corresponding to low false-positive rates is of primary interest whilst in diagnostic testing it is critical to maintain a high TPR in order not to miss detecting diseased subjects. They go on to consider a summary index for the ROC curve restricted to a clinically relevant range of false-positive rates.

Essentially the partial AUC is simply the area under the ROC curve

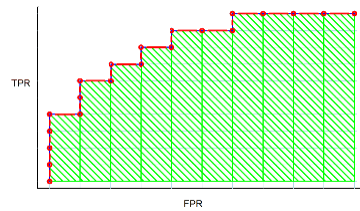


Figure 65: Area under the curve.

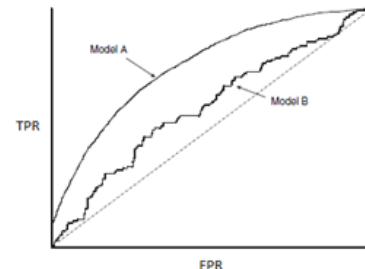


Figure 66: A comparison of two AUC curves.

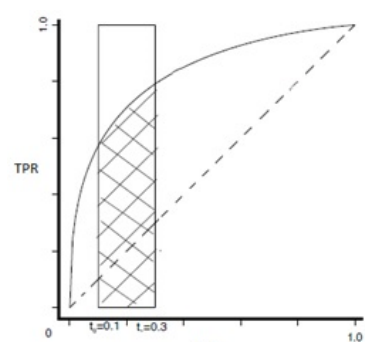


Figure 67: Illustration of an ROC curve and its partial AUC with $t_0 = 0.1$ and $t_1 = 0.3$.

between t_0 and t_1 (Figure 67) where the interval (t_0, t_1) denotes the false-positive rates of interest. The partial AUC is thus the integral between t_0 and t_1 of the ROC curve function as shown in equation 82 (Dodd and Pepe, 2003).

$$AUC(t_0, t_1) = \int_{t_0}^{t_1} ROC(t)dt \quad (82)$$

Two Specific Applications of ROC

The use of ROC curves and Area Under ROC curves is ubiquitous and the list of scientific research papers citing the use of AUC as an essential and effective evaluation technique is endless. They have been used by numerous authors to evaluate models, scoring systems, diagnostic tests, manufacturing defects, spatial images processing, psychiatric evaluations, behaviour analysis, and so on.

An interesting recent paper by Fung et al. (2018) evidences how ROC curve statistics are used to evaluate Convolutional Neural Network for breast cancer classification. It compares the ROC curves generated by the classifier with other reviewed studies. The analysis of AUC helped to evaluate the superiority of the neural network as compared to methods based on experimental results conducted on real patient subjects. Another recent paper (Ding et al., 2017) analyses a deep learning method that accurately recognises an identity from heavily noisy face images. The paper discusses ROC curves with optimal thresholds to show that noise-resistant Deep Neural Network is visibly superior to some state-of-the-art feature extraction algorithms.

Another field where ROC proves to be contributory is psychology. Correct identification of criminals by eyewitness can help to remove a dangerous criminal from society but a false identification can lead to the erroneous conviction of an innocent suspect. A paper by Gronlund et al. (2014) describes how by constructing ROC curves, researchers can trace out discriminability across levels of response bias for each lineup procedure. They illustrate the shortcomings of ratio-based measures and demonstrate why ROC analysis is required for evaluating the performance of a lineup procedure. Luby (2017) goes a step further and examines the application of ROC methodology to lineup data. He shows that by using a log-linear analysis in conjunction with an ROC approach to eyewitness identification, it is possible not only to visualise the trade-off between true positives and false positives, but also to identify which variables are interacting with one another and explain the trade-off.

History

A final note on the history of ROC. The first use of ROC curves is traced back to World War II where in conjunction with signal detection theory, it was developed for the analysis of radar images. Radar operators had to decide whether a blip on the radar screen

represented an enemy target, a friendly ship or just noise. ROC curves helped signal detection theory to measure the ability of radar receiver operators to make these important distinctions. Their ability to do so was called the Receiver Operating Characteristics.

Decades later in the 1970s, ROC analysis was introduced into the biomedical field via the radiological sciences. It has been used extensively to test the ability of an observer to discriminate between healthy and diseased subjects, using a given diagnostic test. As well as to compare the efficacy among the various tests available at that time. Since then, ROC analysis has been extended for use in (but not limited to) visualising and analysing the behaviour of a broad range of diagnostic systems across many fields of science.

Hyperparameters

Li et al. (2017) state that ‘performance of machine learning algorithms depends critically on identifying a good set of hyperparameters’. Hyperparameter tuning is a topic often discussed in literature describing any combination of machine learning algorithms. Adequate values set for hyperparameters will massively influence training and testing of a model, be it in terms of processing time, as well as the actual robustness of the model in question.

Introduction

Defining Hyperparameters

As explained by Bergstra and Bengio (2012), consider a learning model A attempts to map a set of training data X_{train} to a function f which minimizes generalization loss. The model has a set of parameters which may be optimized during training, which are referred to as *model parameters*. The learning algorithm also has a set of parameters which define its architecture and how the system will behave. These may not be changed at runtime, and are referred to as *hyperparameters*. The training and testing performance of a function in relation to minimizing validation error depends heavily on selecting appropriate hyperparameter configurations.

Approaches for Hyperparameter Optimization

Various methods exist to find optimal hyperparameter sets. A multitude of published works go into methods such as *grid search*, where one may refer to Duan and Keerthi (2005). *Random search* is another approach, explored in detail by Bergstra and Bengio (2012). Other approaches make use of *Bayesian Optimizations*, such as Snoek et al. (2012) and Snoek et al. (2015). *Gradient-Based* methods have also been used for hyperparameter tuning, in works such as Keerthi et al. (2006). An overview and comparison of a number of these approaches will be provided in the sections to follow.

Grid Search vs. Random Search

Two common approaches for tuning hyperparameters are grid search and random search. Grid search is an exhaustive method which will

go through every possible combination of a set of hyperparameters, similarly to a brute-force methodology. The hyperparameter set with the best results is then kept and used for the model. Figure 68 shows grid search being applied on a system consisting of two hyperparameters.

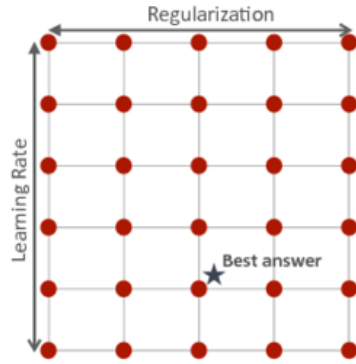


Figure 68: Grid search on a 2D hyperparameter space, adapted from <http://prog3.com/sbdm/blog/google19890102/article/details/50276693> and researchgate.net

By observing Figure 68, it is clear that grid search is guaranteed to find a correct answer due to its exhaustive nature. In order to compare this method with a different approach, Bergstra and Bengio (2012) conducted a study to compare randomly sampled hyperparameters with those obtained through a grid search. Here they refer to a set of model coefficients θ and hyperparameters λ . As stated previously, the objective for a learning algorithm is to find the λ which minimizes a loss function f . Figure 69 shows random search working within the same space as the previous image.

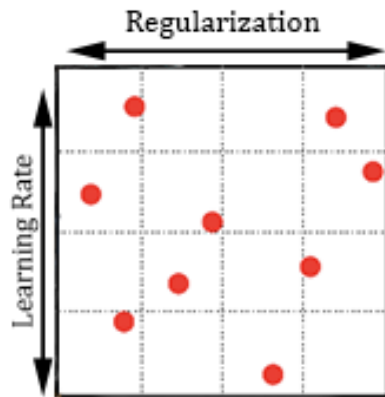


Figure 69: Random search on a 2D hyperparameter space, image adapted from Figure 68

Weighing both approaches together, random search could land on the best answer in the first trial, or the last. Consider the following example:

An SVM has several hyperparameters, one may consider γ , C , and the kernel type for the sake of this example. γ defines the influence of observations on the model, C affects the strictness of classification, and the kernel will affect the shape of the SVM's decision boundary.

These may be represented as follows:

$$\text{kernel} = [\text{linear}, \text{polynomial}, \text{RBF}]$$

$$\gamma = [0.0001, 0.001, 0.01, 0.1, 0.5, 1, 1.5]$$

$$C = [0.001, 0.01, 0.1, 1, 1.5]$$

For this very simple example, grid search would take $3 * 7 * 5 = 105$ runs of an SVM to check each and every hyperparameter combination. With random search, there is a possibility to discover a good configuration much faster, perhaps even instantly. This is particularly the case in real life examples where there may be many more hyperparameters than the ones presented. Since grid search makes use of every possible combination, the system suffers from the *curse of dimensionality*, described by [Chen \(2009\)](#) and [Bellman \(2015\)](#). This results in the algorithm complexity growing exponentially as the number of hyperparameters increase. This constrains grid search to being competitive only in lower dimensional spaces.

Thus, once a problem exceeds two hyperparameters, and sometimes even with only two, random search becomes vastly more efficient according to [Bergstra and Bengio \(2012\)](#). Nevertheless, studies such as [Duan and Keerthi \(2005\)](#) and machine learning libraries such as *libsvm* by [Chang and Lin \(2011\)](#), employed a combination of grid search and manual search. Knowing this, Bergstra and Bengio showed that random search actually generally outperforms grid search for hyperparameter optimization. This is true even when taking into account the element of luck required by random search, since hyperparameter trials are randomly sampled without any selection criteria. [Snoek et al. \(2012\)](#) states that this property provides random search yet another advantage grid search, since the algorithm can operate without caring about which hyperparameters are important to the model.

Bayesian Optimization of Hyperparameters

[Press \(2005\)](#) defined Bayesian techniques as methods making use of Bayes' theorem to combine observational data with subjective beliefs. Bayes' theorem states that considering a hypothesis H and evidence E , the probability of H happening given that E has occurred is calculated as follows:

$$P(H|E) = \frac{P(E|H)P(H)}{P(E)}$$

Bayesian optimizations provide a reasonable approach to modelling uncertainty. According to [Snoek et al. \(2015\)](#), they also provide a natural framework for optimizations of expensive black-box functions. Returning to the scope of hyperparameter tuning, one may

consider a hypothesis as a set of hyperparameters λ and the evidence as data D . Bayesian approaches will keep D static, and modify hyperparameter configurations λ instead to see how the output changes. Essentially, the resulting data obtained through a certain hyperparameter set is then used to predict an expectation for different hyperparameter values. This may be repeated until convergence to a local optimum is reached. Regarding performance, Snoek et al. (2012) compared Bayesian approaches to an exhaustive grid search for optimizing hyperparameters of M3E models with three hyperparameters. The conclusion of the study was that the Bayesian approach was reasonably more efficient than grid search. This also makes sense

Gradient-Based Hyperparameter Optimization

A particular project by Keerthi et al. (2006) investigates hyperparameter values in SVMs and Bayesian models. In this system, the variables C and γ are considered. C is the regularization coefficient, influencing how leniently the SVM hyperplane classifies observations. Figure 70 and Figure 71 show how different values of C influence a decision space for classification.

As previously stated, γ defines the influence of points on the variance of a model. As expected, experiments in this study showed that performance of the SVM varied considerably as a function of hyperparameter values. Gradient-based optimization techniques were used, computing the slope of a performance validation function with respect to the vector h of hyperparameters. The gradient-descent approach attempts to minimize the cost function of the hyperparameter problem by converging to local minima. To reach this goal, the gradient of the function decreases with each iteration of the algorithm. This will update the coefficients β of the system, which will then be applied to the feature vectors in the next iteration. This process is repeated until convergence towards a local minimum is reached. This system allowed for a fast method to determine optimal sets of hyperparameters, which also outperformed grid search even for models with a low dimensionality of two hyperparameters.

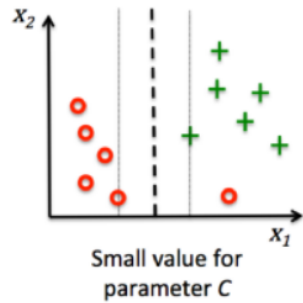


Figure 70: Small C

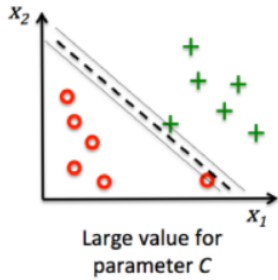


Figure 71: Big C , both images from https://www.bogotobogo.com/python/scikit-learn/scikit_machine_learning_Support_Vector_Machines_SVM.php

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