A Machine Learning Study Guide



Machine Learning Handbook



ICS5110, class of 2018/9





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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

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). Each of these approaches are further divided into different cross-validation methods, which are explained below.

Exhaustive cross-validation

• Leave-*p*-out (L*p*O)

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. 2a. 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 nchoose1 = n times, however might still be inapplicable for large values of n (Arlot et al., 2010).

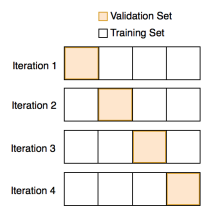
Non-exhaustive cross-validation

· 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. 3. 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).



• 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

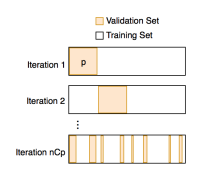


Figure 1: Exhaustive cross-validation methods: Leave-p-Out

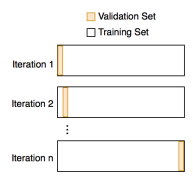


Figure 2: Exhaustive cross-validation methods: Leave-One-Out

Figure 3: k-Fold Cross Validation where k=4

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. 1, 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.

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

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. 5a. 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. 5b, 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, lets take an example using the stratified k-fold crossvalidation 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 o 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 class². As depicted in Fig. 6, we can now start forming our 10 folds by taking 10% of each age-group bucket, randomly without replacement. Hence, we will 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

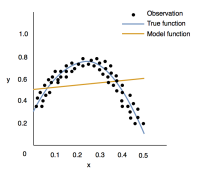


Figure 4: Underfitting

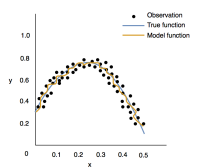


Figure 5: Overfitting

² Down-sampling or up-sampling are common techniques used when there is an unequal amount of samples for the different classes.

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. 3). 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.

Labelled Data Age Group 6 Age Group 7 Age Group 2 Age Group 3 Age Group 1 Age Group 4 Age Group 5 Age Group 8 125 125 125 125 125 125 125 125 10% 10% 10% 10% 10% 100 100 100 100 100 100 100 100 100 100 10-folds

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 6: Stratified 10-fold crossvalidation on 1000 labelled images of 8 different classes

Confusion Matrix

A *Confusion Matrix*³ (CM) is a cross-tabulation illustrating 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).

³ Also called a Contingency Table.

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

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

Even though CMs are commonly used to evaluate binary classifiers, they aren't restricted to 2-class models (Martin and Jurafsky, 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).

M&M'sSkittlesSmartiesM&M's3438Skittles1285Smarties2422

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.

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

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

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.

In binary classification, the CM consists of 2 specially designated classes called the *positive* class and the *negative* class (Saito and Rehmsmeier, 2015). 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 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 (Eq. 2-8) 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 (e.g. a rare diseases) is underrepresented (Raeder et al., 2012).

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 or not. Even though both misclassifications are undesirable, in medical applications it's better to err on the side of caution since FN could be fatal.

Accuracy (ACC) is the proportion of correct predictions (Eq. 2). 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're more class-sensitive and give better indicators when the dataset is imbalanced.

$$ACC = \frac{|TP \cup TN|}{|TP \cup FP \cup TN \cup FN|} \tag{2}$$

Negative Predictive Value (NPV) is the ratio of the correct negative predictions from the total negative predictions (Eq. 3).

$$NPV = \frac{|TN|}{|TN \cup FN|} \tag{3}$$

True Negative Rate (TNR), or *Specificity*, is the ratio of the correct negative predictions from the total true negatives (Eq. 4).

$$TNR = \frac{|TN|}{|TN \cup FP|} \tag{4}$$

True Positive Rate (TPR), also called *Sensitivity* or *Recall*, is the ratio of the correct positive predictions from the total true positives (Eq.5).

$$TPR = \frac{|TP|}{|TP \cup FN|} \tag{5}$$

Sensitivity and Specificity can be combined into a single metric (Eq. 6). These metrics are often used in domains in which minority classes

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).

are important (Kuhn and Johnson, 2013). For example, a medical classifier (El-Dahshan et al., 2010) could be evaluated by measuring Sensitivity to find the number of patients with a condition who tested positive, and Specificity to measure the number of patients who don't have the condition and tested negative.

$$Sensitivity \times Specificity = \frac{|TP| \times |TN|}{|TP \cup FN| \times |TN \cup FP|}$$
 (6)

Positive Predictive Value (PPV), or Precision, is the ratio of the correct positive predictions from the total positive predictions (Eq. 7).

$$PPV = \frac{|TP|}{|TP \cup FP|} \tag{7}$$

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 (Eq. 8) can be derived by finding their harmonic mean (Kelleher et al., 2015).

$$F\text{-}score = 2 \times \frac{PPV \times TPR}{PPV + TPR} \tag{8}$$

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.

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 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.

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	$Skittles \cup M\&M's$

Taking into account the above, CMs are suitable for visualising, evaluating, and comparing the performance of binary or multi-class

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.

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 Fig. 7 were trained on 70% of the wines dataset included with Scikit-Learn, whilst the remaining 30% was kept as a test set to evaluate the model. The dataset has 3 classes and 13 features, but only the results for class 1 and 2; and the alcohol and malic acid features; will be presented. It was scaled by standardising the features using $\frac{x_i - \mu}{\sigma}$, for all $x_i \in S$ using mean μ and standard deviation σ .

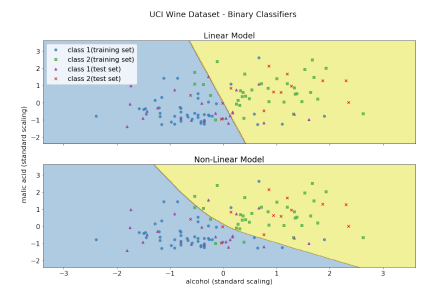


Figure 7: Decision boundary learned by a linear and non-linear binary classifier.

Linear	Non-Linear
0.72	0.78
0.77	0.77
0.70	0.78
0.84	0.86
0.76	0.82
	0.72 0.77 0.70 0.84

Table 6: Statistics derived from the CMs in Fig. 8.

As it can be intuitively deduced from Fig. 7, the decision boundary of the non-linear model is a better fit than the linear model. The CMs in Fig. 8 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.

UCI Wine Dataset - Confusion Matrix Linear Model Non-Linear Model 16 16 18 class 1 14 12 10 8 10 10 class 1 class 2 class 1 class 2

Figure 8: The linear classifier has 16 TP, 10 TN, 7 FN and 3 FP, whilst the nonlinear classifier has 18 TP, 10 TN, 5 FN and 3 FP.

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