Practical Issues Machine Learning - A.Y. 2022/23 - Padova



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Underfitting/Overfitting and learning parameters

• Suppose we have some data (60 points) that we want to fit a curve to



• Let fit a polynomial, of the form

$$y = w_0 + w_1 x + w_2 x^2 + \dots + w_p x^p$$



Underfitting/Overfitting and learning parameters

- How to choose *p* ? (Hypothesis Space)
- For various *p*, we can find and plot the best polynomial, in terms of minimizing the Empirical Error (Mean Squared Error in this case)
- Here are the solutions for different values of p

















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Practical Issues



Underfitting/Overfitting and learning parameters

• Here is a summary of the training error ... and the error on some new TEST data (100,000 extra points) from the same distribution, as a function of *p*:



- The BIAS measures the distortion of an estimate
- The VARIANCE measures the *dispersion* of an estimate





Bias and Variance



Let $y = f(\mathbf{x}) + \epsilon$ the target function, where ϵ has zero mean, variance σ^2 .

We want to find a function $\hat{f}(\mathbf{x})$ that approximates the true function $f(\mathbf{x})$ as well as possible, by means of some learning algorithm.

Given any pair (\mathbf{x}, y) , the following holds:

$$\mathbb{E}[(y - \hat{f}(\mathbf{x}))^2] = \left(\mathsf{Bias}[\hat{f}(\mathbf{x})]\right)^2 + \mathsf{Var}[\hat{f}(\mathbf{x})] + \sigma^2$$

where,

$$\begin{split} \mathsf{Bias}[\hat{f}(\mathbf{x})] &= \mathbb{E}[\hat{f}(\mathbf{x})] - f(\mathbf{x}) \\ \mathsf{Var}[\hat{f}(\mathbf{x})] &= \mathbb{E}[\hat{f}(\mathbf{x}) - \mathbb{E}[\hat{f}(\mathbf{x})]]^2 \end{split}$$

and the expectations refer to different sets of training data.



Underfitting/Overfitting and learning parameters

- For very low *p*, the model is very simple, and so cannot capture the full complexity of the data (Underfitting! also called **bias error**)
- For very high *p*, the model is complex, and so tends to overfit to spurious properties of the data (Overfitting! also called **variance error**)

Unfortunately we cannot use the test set to pick up the right value of p!

PRACTICAL PROBLEM: how can we use the training set to set the value of p?

Model Selection and Hold-out



We can keep some of our training data out **Hold-out procedure**

- A small subset of *Tr*, called the validation set (or hold-out set), denoted *Va*, is kept apart;
- **2** A classifier/regressor is trained using examples in Tr Va;
- Step 2 is performed with different values of the hyperparameter(s) (in our example, the value *p*), and tested against the hold-out sample.

It is possible to show (Hoeffding's Inequality) that the evaluation performed in step 2 gives an unbiased estimate of the error made by a classifier trained with the same values of the hyperparameter(s) and with training set of cardinality |Tr| - |Va| < |Tr|.

In an operational setting, after parameter optimization, one typically re-trains the classifier on the entire training corpus, in order to boost effectiveness (debatable step!).

K-fold Cross Validation



An alternative approach for model selection (and evaluation) is the K-fold cross-validation method

K-fold CV procedure

- K different classifiers/regressors h₁, h₂,..., h_k are built by partitioning the initial corpus Tr into k disjoint sets Va₁,..., Va_k and then iteratively applying the Hold-out approach on the k-pairs (Tr_i = Tr Va_i, Va_i)
- **②** Final error is obtained by individually computing the errors of h_1, \ldots, h_k , and then averaging the results

The above procedure is repeated for different values of the hyperparameter(s) and the setting (model) with the smallest final error is selected

The special case k = |Tr| of k-fold cross-validation is called **leave-one-out** cross-validation

Back to our example



• Let's apply 5-fold CV



• Minimum error reached for p = 3, rather than the optimal p = 12

• Clearly, cross validation is no substitute for a large test set. However, if we only have a limited training set, it is often the best option available.

Analysis of Cross Validation



- What happens varying k?
- For higher k's we have larger training sets, hence less bias! Smaller validation sets, hence more variance!
- For lower k's we have smaller training sets, hence more bias! Larger validation sets, hence less variance!



Almost all learning algorithms have (hyper)parameters!

- Support Vector Machines: *C*, type of kernel (polynomial, RBF, etc.), kernel parameter (degree of polynomial, width of RBF, etc.)
- Neural Networks: nonlinear/linear neurons, number of hidden units, η , other learning parameters we have not discussed (e.g., momentum μ)

Hold-out or Cross-Validation can be used to select the "optimal" values for the (hyper)parameters (i.e., select the "optimal" model).



Classification accuracy:

- Very common in ML,
- Proportion of correct decisions,
- Not appropriate when the number of positive examples is much lower than the number of negative examples (or viceversa)

Precision, Recall and F_1 are better in these cases!



	Relevant	Not Relevant
Retrieved	True Positive (TP)	False Positive (FP)
Not Retrieved	False Negative (FN)	True Negative (TN)

why not using the accuracy
$$\alpha = \frac{TP+TN}{TP+TN+FP+FN}$$
 ?

Alternative measures (precision and recall):

$$\pi = \frac{\mathsf{TP}}{\mathsf{TP} + \mathsf{FP}} \qquad \rho = \frac{\mathsf{TP}}{\mathsf{TP} + \mathsf{FN}}$$

Effectiveness for Unbalanced Classification



If relevance is assumed to be binary-valued, effectiveness is typically measured as a combination of

- Precision is the "degree of soundness" of the system: *P*(RELEVANT|RETRIEVED)



How can one trade-off between precision and recall? F-measure (weighted harmonic mean of the precision and the recall)

$$F_{\beta} = \frac{(1+\beta^2)\pi\rho}{\beta^2\pi + \rho}$$

 $\beta < 1$ emphasizes precision

$$\beta = 1 \Rightarrow F_1 = 2 \frac{\pi \rho}{\pi + \rho}$$



Multiclass classification consists of a classification task with more than two classes; e.g., classify a set of images of fruits which may be oranges, apples, or pears. Multiclass classification makes the assumption that each sample is assigned to one and only one label: a fruit can be either an apple or a pear but not both at the same time.

How can the multiclass problem be reduced to a set of binary problems?



The one-vs-rest strategy, is implemented in OneVsRestClassifier in sklearn. The strategy consists in fitting one classifier per class.

For each classifier, the class is fitted against all the other classes. In addition to its computational efficiency (only n_classes classifiers are needed), one advantage of this approach is its interpretability. Since each class is represented by one and only one classifier, it is possible to gain knowledge about the class by inspecting its corresponding classifier. This is the most commonly used strategy and is a fair default choice.



The one-vs-one strategy, is implemented in OneVsOneClassifier in sklearn. The strategy consists in fitting one classifier for each pair of classes.

At prediction time, the class which received the most votes is selected.

Since it requires to fit $n_classes * (n_classes - 1) / 2$ classifiers, this method is usually slower than one-vs-the-rest. However, this method may be advantageous for algorithms such as kernel algorithms which don't scale well with n_samples. This is because each individual learning problem only involves a small subset of the data whereas, with one-vs-the-rest, the complete dataset is used n_classes times.

Evaluation in Multiclass classification



A very intuitive way to show the results of a multiclass predictor is using the confusion matrix.

Example: Prediction : A B A B C C C A B True labels: A A A A C C C B B





Precision can be calculated separately for each class. For each row, we take the number on the diagonal, and divide it by the sum of all the elements in the column.

prec_A: 2/3 = 0.67 prec_B: 2/4 = 0.50 prec_C: 3/3 = 1.00

Recall can be calculated separately for each class. It is the value on the diagonal, divided by the sum of the values in the row.

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recall_A: 2/4 = 0.50
recall_B: 2/3 = 0.67
recall_C: 3/3 = 1.00
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Micro and Macro Averaging



To extract a single number from the class precision, recall or F1-score, it is common to adopt two different kinds of average.

The first, is to compute the score separately for each class and then taking the average value — this is called macro averaging. This kind of averaging is useful when the dataset is unbalanced and we want to put the same emphasis on all the classes.

$$\frac{r_A + r_B + r_C}{3} = 0.7233$$

The second is to calculate the measure from the grand total of the numerator and denominator, this is called micro averaging and is useful when you want to bias your classes towards the most populated class. For our 'recall' example:

$$\frac{2+2+3}{4+3+3} = 0.636$$



Recap

Notions

- Bias and Variance
- Under-fitting and Over-fitting
- Model Selection (Hold-out and Cross Validation)
- Unbalanced data
- Multi-class classification and evaluation

Exercises

- Bias and Variance evaluation in a controlled regression task
- Create a large regression dataset and demonstrate that by taking the mean of the models obtained with large *p* and relatively small training sets, we are able to break down the variance and thus significantly improve performance. Why can't this methodology actually be used? how to reduce the variance of the models even with small training sets?