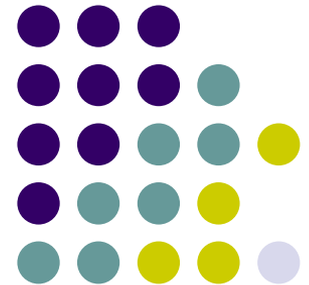


# *Big Data Analytics*

## Lecture 4/A Cross-Validation



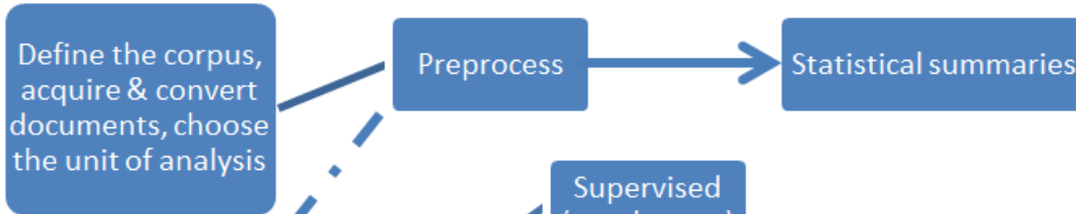
UNIVERSITÄT  
LUZERN



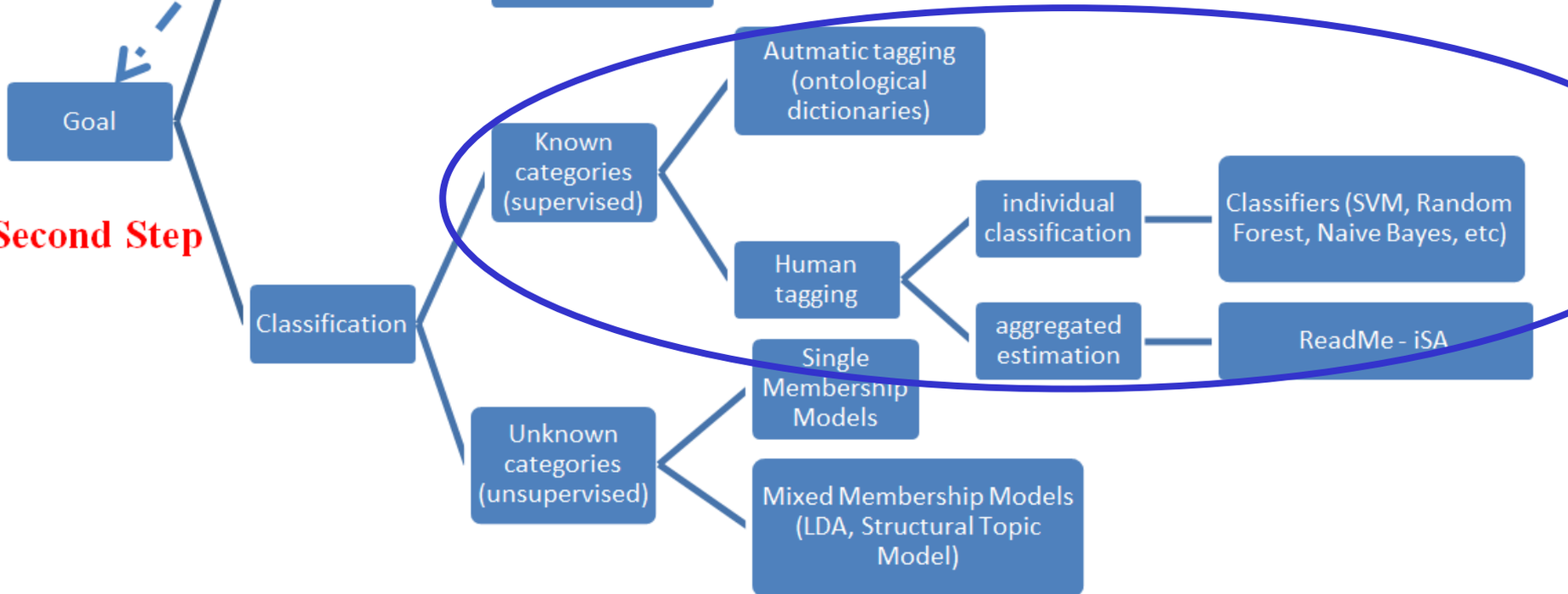
# Our Course Map



## First Step



## Second Step

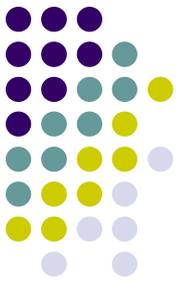




# Reference

- ✓ Grimmer, Justin, and Stewart, Brandon M. (2013). Text as Data: The Promise and Pitfalls of Automatic Content Analysis Methods for Political Texts. *Political Analysis*, 21(3): 267-297
- ✓ Curini, Luigi, and Robert Fahey (2020). Sentiment Analysis and Social Media. In Luigi Curini and Robert Franzese (eds.), *SAGE Handbook of Research Methods in Political Science & International Relations*, London, Sage, chapter 29
- ✓ Cranmer, Skyler J. and Desmarais, Bruce A. (2017) What Can We Learn from Predictive Modeling?, *Political Analysis*, 25: 145-166

# Validation

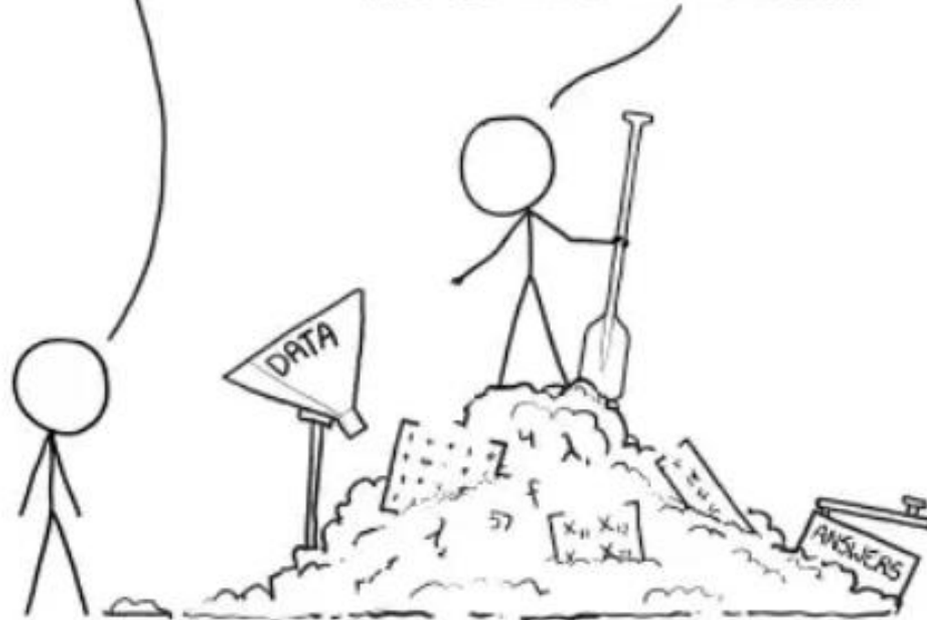


THIS IS YOUR MACHINE LEARNING SYSTEM?

YUP! YOU POUR THE DATA INTO THIS BIG PILE OF LINEAR ALGEBRA, THEN COLLECT THE ANSWERS ON THE OTHER SIDE.

WHAT IF THE ANSWERS ARE WRONG?

JUST STIR THE PILE UNTIL THEY START LOOKING RIGHT.

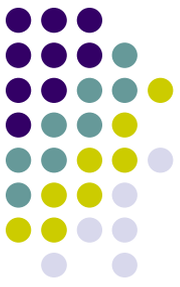




# Validation

Since the ultimate goal of supervised learning is to find generalizable patterns of association, models are typically subject to some form of regularization - typically in the form of a constraint that pushes the model toward parsimony - and are **selected based on their ability to generate good out-of-samples predictions**

Clearly, it is impossible to evaluate a model's performance on the universe of unsampled test instances (i.e., the test set), so an approximate measure of performance must be devised



# Validation

Supervised methods are designed to automate the hand coding of documents into categories or measuring the proportion of documents in categories as we have already noticed

If a method is performing well, it will **directly replicate the hand coding**. If it performs poorly, it will fail to replicate the coding – instead introducing serious errors

This clear objective implies a **clear standard for evaluation**: comparing the output of machine coding to the output of hand coding. From here the idea of **validation**!

# Validation

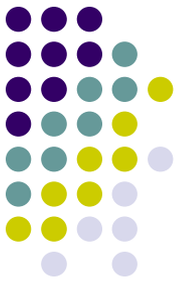


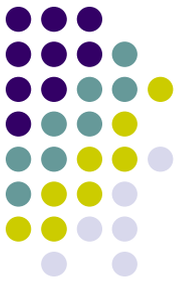
The *ideal validation* procedure would divide the data into **three subsets**

1. Initial model fitting would be performed on the training-set
2. Once a final model is chosen, *a second set of hand-coded documents - the validation set* - would be used to assess the performance of the model
3. The final model would then be applied to the test to complete the classification

# Validation

This approach to validation is difficult to apply in most settings. But **cross-validation** (also called: **K-fold validation**) can be used to replicate this ideal procedure





# Validation

In K-fold cross-validation, the training set is randomly partitioned into some groups (say two: K1 and K2)

For each group, the first model is trained on K1, then applied to the K2 to assess performance; similarly a model is trained on the K2 and then applied to K1 to assess performance

Then you take the average across the results you get in the two scenarios

# Validation



And if you want to run a K-fold cross-validation with K larger than 2?

The algorithm is as follow:

1. Randomly split the data set into k-subsets (or k-fold) (for example 5 subsets)
2. Reserve one subset and train the model on all other subsets (4 in this case)
3. Test the model on the reserved subset and record the prediction error
4. Repeat this process until each of the k subsets has served as the test set
5. Compute the average of the k recorded errors. This is called the **cross-validation error** serving as the performance metric for the model



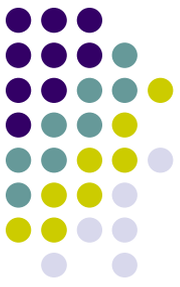
# Validation



Typical question, is how to choose right value of  $k$ ?

Lower value of  $k$  is more biased and hence undesirable. On the other hand, higher value of  $k$  is less biased, but can suffer from large variability

In practice, one typically performs  $k$ -fold cross-validation using  $k = 5$  or  $k = 10$ , as these values have been shown empirically to yield test error rate estimates that suffer neither from excessively high bias nor from very high variance

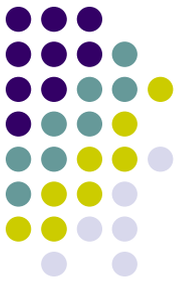


# Validation

When you run a ML algorithm on the test-set, there are no available statistics to control for the goodness-of-fit of your prediction (by definition the “true” values of the test-set are unknown!)

That is why **cross-validation** is so important! This is the **only way** to control if the ML algorithm you are using is doing a good job or not (unless you are ready to believe in that by fiat)!

Moreover, cross-validation **avoids overfitting** by focusing on **out-of-sample prediction** and **selects the best model** for the underlying data from a set of candidate models

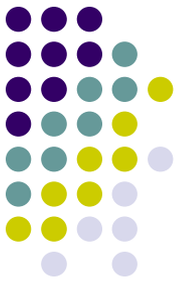


# Validation

Which statistics (or **performance metrics**) should we use to assess model performance?

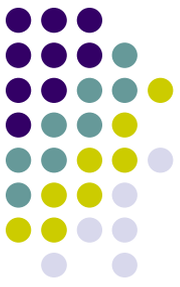
There are several of them, but we are going to focus on three metrics for individual classifiers with text-analysis

# Validation



**Accuracy:** proportion of correctly classified documents

While of course we want this score to be as high as possible, it can also be important to look at the two components which make up that score, known as **recall** and **precision**



# Validation

**Recall** or **Sensitivity** (for a category  $k$ ) is a measure of what proportion of instances of a given category the algorithm correctly identified; so for example, if there were 10 instances of the category “positive” in the data set, and the algorithm correctly identified 8 of them, we would say that this algorithm has “recall of 0.8 for the category *positive*”

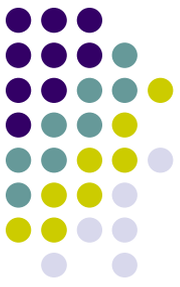
- ✓ given that a human coder labels a document as belonging to category  $k$ , what is the chance the machine identifies the document?

# Validation



**Precision or Positive Predictive Value** (for a category  $k$ ) on the other hand is a measure of how many of the times the algorithm identified a category were actually correct, as against how many times were false positives. In the above example, where the algorithm correctly identified 8 of the 10 instances of *positive*, perhaps the algorithm also mis-identified 4 other documents as *positive* - so 8 out of its 12 *positive* classifications were correct, allowing us to say that it has a “precision of 0.667 for the category *positive*”

- ✓ given that the machine guesses category  $k$ , what is the probability that the machine made the right guess?

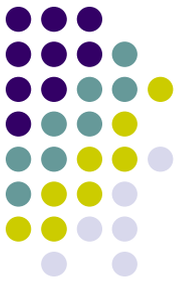


# Validation

If you find market differences between the **recall and precision** (for example, with a recall rate  $\gg$  precision) implies that your algorithm guesses too often that a document belongs to category  $k$

The result is that it labels a large portion of the human coder's as  $k$  correctly (and so you have a **high recall rate**). But it also includes several documents that humans label differently (and so you have a **low precision**)

This sometimes applies when the original  $k$  category, compared to the other  $k_{n-1}$  categories, is the most relevant category in the training-set



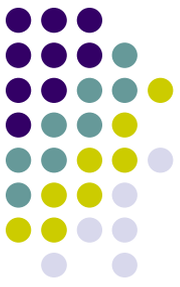
# Validation

The aggregate of the recall and precision scores for a category is known as the **f1 score**

More precisely, the traditional F-measure or balanced F-score (**f1**) is the harmonic mean of precision and recall:

$$\mathbf{f1} = (2 * \text{precision} * \text{recall}) / (\text{precision} + \text{recall}) \dots$$

...where the highest level of performance (of f1) is equal to 1 and the lowest 0



# Validation

The **average of the f1 scores for all the categories** is a reasonable rough measurement of the performance of the algorithm (more than accuracy alone!)

However, before using the algorithm for any serious analysis work, it is advisable also to take a look at the precision and recall scores for individual categories - you may find that a category you are planning to use in your analysis actually has very high rate of false-positive or false-negative identifications, which could cause serious problems for your results

Let's an example on how to compute the statistics we discussed up to now from the Confusion matrix

# Performance metrics



## Confusion matrix

Classification (algorithm)	Actual label	
	Black	White
Black	True Black	False Black
White	False White	True White

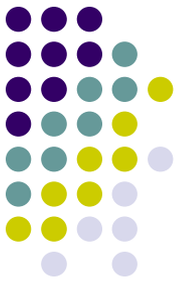
$$\text{Accuracy} = \frac{\text{TrueBlack} + \text{TrueWhite}}{\text{TrueBlack} + \text{TrueWhite} + \text{FalseBlack} + \text{FalseWhite}}$$

$$\text{Precision}_{\text{Black}} = \frac{\text{TrueB}}{\text{TrueB} + \text{FalseB}} \longrightarrow \text{Think horizontally}$$

$$\text{Recall}_{\text{Black}} = \frac{\text{TrueB}}{\text{TrueB} + \text{FalseW}} \longrightarrow \text{Think vertically}$$

$$f_{1\text{Black}} = \frac{2 * \text{precisionB} * \text{recallB}}{\text{precisionB} + \text{recallB}}$$

# Performance metrics: an example



Classification (algorithm)	Actual label	
	Black	White
Black	800	100
White	50	50

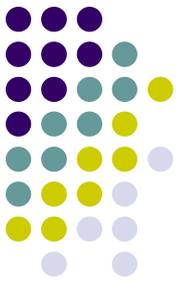
$$\text{Accuracy} = \frac{800 + 50}{800 + 50 + 100 + 50} = 0.85$$

$$\text{Precision}_{\text{Black}} = \frac{800}{800 + 100} = 0.88$$

$$\text{Recall}_{\text{Black}} = \frac{800}{800 + 50} = 0.94$$

$$f_1 \text{ Black} = \frac{2 * .88 * .94}{.88 + .94} = 0.91$$

# Performance metrics



## Confusion matrix

Classification (algorithm)	Actual label	
	Black	White
Black	True Black	False Black
White	False White	True White

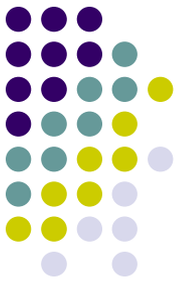
$$\text{Accuracy} = \frac{\text{TrueBlack} + \text{TrueWhite}}{\text{TrueBlack} + \text{TrueWhite} + \text{FalseBlack} + \text{FalseWhite}}$$

$$\text{Precision}_{\text{White}} = \frac{\text{TrueW}}{\text{TrueW} + \text{FalseW}} \longrightarrow \text{Think horizontally}$$

$$\text{Recall}_{\text{White}} = \frac{\text{TrueW}}{\text{TrueW} + \text{FalseB}} \longrightarrow \text{Think vertically}$$

$$f_{1\text{White}} = \frac{2 * \text{precisionW} * \text{recallW}}{\text{precisionW} + \text{recallW}}$$

# Performance metrics: an example



Classification (algorithm)	Actual label	
	Black	White
Black	800	100
White	50	50

$$\text{Accuracy} = \frac{800 + 50}{800 + 50 + 100 + 50} = 0.85$$

$$\text{Precision}_{\text{White}} = \frac{50}{50 + 50} = 0.5$$

$$\text{Recall}_{\text{White}} = \frac{50}{50 + 100} = 0.33$$

$$f_1 \text{ White} = \frac{2 * .5 * .33}{.5 + .33} = 0.39$$



# Performance metrics

In this example you are going to have a single Accuracy value=0.85

Then you could take the **average of the f1 scores** for the classes as another (and more reliable) measure of the performance of the algorithm

In our case:  $(.91+.39)/2=.65$

You see the difference here between Accuracy and the averaged f1 score. This difference is due that we are doing well with the Black class, but relative poorly with the White class



# Performance metrics

If you have a very imbalanced datasets (that is, when the numbers of observations in different classes vary greatly - a point on which we will return later on), besides the average value of f1, you could also decide to focus on **Balanced Accuracy**

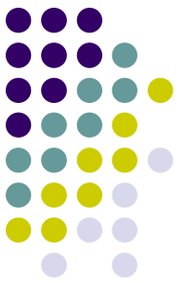
It is defined as the macro-average of the recall obtained on each class

In our previous example of cats & dogs:

Accuracy: 0.85

Balanced Accuracy:  $(0.94+0.33)/2=0.635$

# Another example with 3 categories



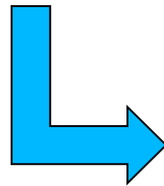
		Actual class		
		Cat	Dog	Rabbit
Predicted class	Cat	5	2	0
	Dog	3	3	2
	Rabbit	0	1	11

Here **Accuracy** is equal to the ratio between the sum of the diagonal (i.e., the sum of «True Positive») and the total number of observations, i.e.,  $(5+3+11)/(5+2+3+3+2+1+11)=0.704$

# Another example with 3 categories

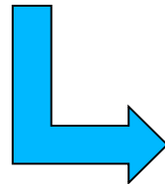


For each category  $k$  we can move from here



		Actual class		
		Cat	Dog	Rabbit
Predicted class	Cat	5	2	0
	Dog	3	3	2
	Rabbit	0	1	11

to here (example for the “cat” category)



		Actual class	
		Cat	Non-cat
Predicted class	Cat	5 True Positives	2 False Positives
	Non-cat	3 False Negatives	17 True Negatives

# Another example with 3 categories



Then:

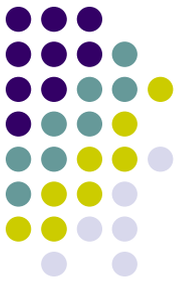
In the case, Precision for the cat class is:  $5/(5+2)=0.71$

Recall for the cat class is:  $5/(5+3)=0.625$

f1 for the cat class is:  $2*(0.625*0.71)/(0.625+0.71)=0.66$

You can do the same thing for the dog and the rabbit cases, and then averaging across values to have a sense of the overall performance of your model

# Validation



Depending on the application, scholars may conclude that the supervised method is able to sufficiently replicate human coders. A largely employed rule-of-thumb is getting accuracy  $>.85$  for example (or average f1  $>.75$ )

Or, additional steps can be taken to improve accuracy, including trying to apply other ML algorithms



# Validation

Most algorithms also have a range of “**hyper-parameters**” (or “tuning parameters”)– assumptions and modifiers which are used to fine-tune the model and which can be set to different values prior to training – that can significantly impact performance (remember about  $C$  in SVM or the number of trees in RF)

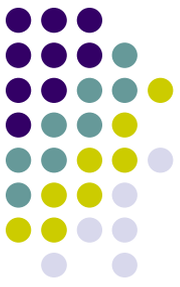
Finding the right set of hyper-parameters for a certain task and a specific data set is also largely a case of trial and error, and it can only be done via cross-validation!



# Validation

Some packages in R (such as `Caret` or `h2o` or the same `Quanteda` with the library `quanteda.classifiers`) provide ways to automate this task; this is known as a “grid search”, allowing researchers to exhaustively search through every combination of a set of hyper-parameters to find the best performing model

This process can take a lot of time – often in the order of several hours for algorithms with complex sets of parameters – but often yields better performance than the default parameter set



# Validation

Moreover remember: the purpose of cross-validation is **model checking!**

Accordingly, cross-validation allows also **to select among different machine-learning algorithms!**

The **No Free Lunch Theorem** states that no machine learning algorithm is always better at predicting new, unobserved, data points universally. So...

Which is the machine-learning algorithms to prefer given your specific training-test?!? The one that fares better in cross-validation!!!

So trust this latter one, when you want to classify the unlabeled test-set!



# Validation: a summary

Two possible routes in this regard according to how you want to deal with the **hyper-parameters**:

First route (to success...)

- a) You keep the default hyper-parameters of your ML algorithms;
- b) you run a CV on each of such ML algorithms
- c) you select the one (or two) with the best performance on CV
- d) you fine-tune the hyper-parameters on such model(s)
- e) you re-run CV just on them
- f) you keep the ML algorithm that performs better in the CV



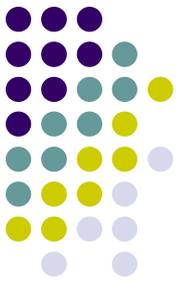
# Validation: a summary

Two possible routes in this regard according to how you want to deal with the **hyper-parameters**:

Second route (to success...)

- a) You fine-tune the hyper-parameters on each of the your ML algorithms you want to test
- b) you run CV on each of them
- c) you keep the ML algorithm that performs better in the CV

# R packages to install



```
install.packages("cvTools", repos='http://cran.us.r-project.org')
```