### **Session 13:**

Supervised learning, part 2

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#### Agenda

- 1. model building
- 2. model selection
  - cross validation
  - tools for selection
- 3. dimensionality reduction
- 4. measures for classification

#### **V**aaaamos

```
In [51]: import warnings
    from sklearn.exceptions import ConvergenceWarning
    warnings.filterwarnings(action='ignore', category=ConvergenceWarning)

import matplotlib.pyplot as plt
    import numpy as np
    import pandas as pd
    import seaborn as sns

plt.style.use('default') # set style (colors, background, size, gridlines etc.)
    plt.rcParams['figure.figsize'] = 10, 4 # set default size of plots
    plt.rcParams.update({'font.size': 18})
```

### Supervised problems (1)

What is the tradeoff for making supervised regression models?

In [3]: # f\_bias\_var['regression'][2] Out[3]: **Underfitting:** Balanced model: Overfitting: Non-linear Hyper-sensitive Linear 1.0 1.0 1.0 0.8 0.8 0.8 0.6 0.6 0.6 > 0.4 0.4 0.4 0.2 0.2 0.2 0.0 0.00 0.0 0.0 0.00 0.25 0.50 0.75 1.00 0.25 0.50 0.75 1.00 0.25 0.50 0.75 1.00 Х Х estimated model data generating process

#### Supervised problems (2)

What was a remedy to overfitting in linear models? How do we measure overfitting?

Regularization

- Too many irrelevant features solved by L1 regularization ~ lasso
- Exploding coefficients solved by L2 regularization ~ ridge

# Model building

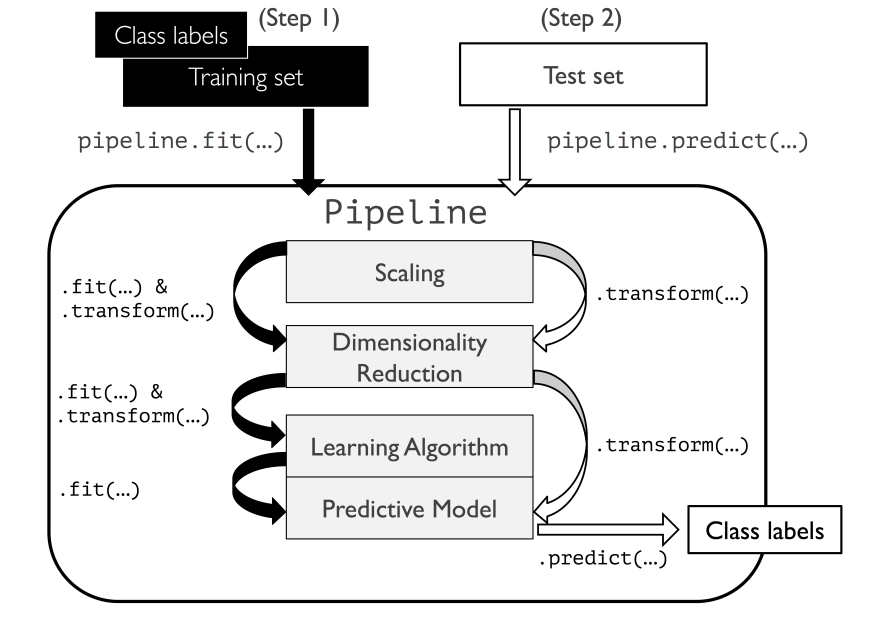
### Model pipelines (1)

Is there a smart way to build ML models?

Yes, we build a pipeline:

- Preprocessing data
  - Standard: adding polynomials, imputation, rescaling
  - Unsupervised learning (more info..)
- Supervised learning

## Model pipelines (2)



## Model pipelines (3)

What are the advantages of using a pipeline?

- Ensures good practice we only fit on training data.
- Much less code!

### Applying a model pipeline (1)

What would this look like in Python?

ly=False))

('standardscaler', StandardScaler(copy=True, with mean=True, with std=True))

#### Applying a model pipeline (2)

Let's some load Boston house price data

```
In [49]:
         print('\n'.join(load boston()['DESCR'].split('\n')[13:28]))
             :Attribute Information (in order):
                            per capita crime rate by town
                 - CRIM
                 - ZN
                           proportion of residential land zoned for lots over 25,000 sq.ft.
                 - INDUS
                            proportion of non-retail business acres per town
                           Charles River dummy variable (= 1 if tract bounds river; 0 otherwi
                 - CHAS
         se)
                 - NOX
                            nitric oxides concentration (parts per 10 million)
                            average number of rooms per dwelling
                 - RM
                 - AGE
                            proportion of owner-occupied units built prior to 1940
                            weighted distances to five Boston employment centres
                 - DIS
                            index of accessibility to radial highways
                 - RAD
                 - TAX
                            full-value property-tax rate per $10,000
                 - PTRATIO
                           pupil-teacher ratio by town
                            1000(Bk - 0.63)^2 where Bk is the proportion of blacks by town
                 - B
                 - LSTAT
                            % lower status of the population
                            Median value of owner-occupied homes in $1000's
                 MEDV
```

### Applying a model pipeline (3)

And how do I apply the pipe on the data?

'B' 'LSTAT']

```
In [41]: from sklearn.model_selection import train_test_split
    from sklearn.datasets import load_boston
    X = load_boston().data
    y = load_boston().target

print(load_boston().feature_names)

# splitting into train and test data
    X_train, X_test, y_train, y_test = train_test_split(X, y)

# apply preproc - fit on train
    pipe_preproc.fit(X_train) # fit to training
    X_train_prep = pipe_preproc.transform(X_train) # transform training data
    X_test_prep = pipe_preproc.transform(X_test) # transform test data

['CRIM' 'ZN' 'INDUS' 'CHAS' 'NOX' 'RM' 'AGE' 'DIS' 'RAD' 'TAX' 'PTRATIO'
```

#### Applying a model pipeline (4)

And how do I apply the pipe on the data?

```
In [56]: # THE PIPE APPLIED
         # apply preproc - fit on train
         pipe preproc = make pipeline(PolynomialFeatures(),
                                       StandardScaler())
         pipe_preproc.fit(X_train) # fit to training
         X train prep = pipe preproc.transform(X train) # transform training data
         X test prep = pipe preproc.transform(X test) # transform test data
         # WITHOUT PIPE
         poly trans = PolynomialFeatures()
         scaler = StandardScaler()
         X train poly = poly trans.fit transform(X train)
         X test poly = poly trans.fit transform(X test)
         scaler.fit(X train poly)
         X train prep alt = scaler.transform(X train poly)
         X test prep alt = scaler.transform(X test poly)
```

## **Model selection**

#### Measuring the problem

Does machine learning work out of the box?

- In some cases ML works quite well out of the box.
- Often ML requires making careful choices.
  - Note that automated machine learning packages and services exist.

Which choices are to be made?

- We need to pick model building hyperparameters.
- E.g.  $\lambda$  for Lasso, Ridge.

#### Model validation (1)

How do we measure our model's performance for different hyperparameters?

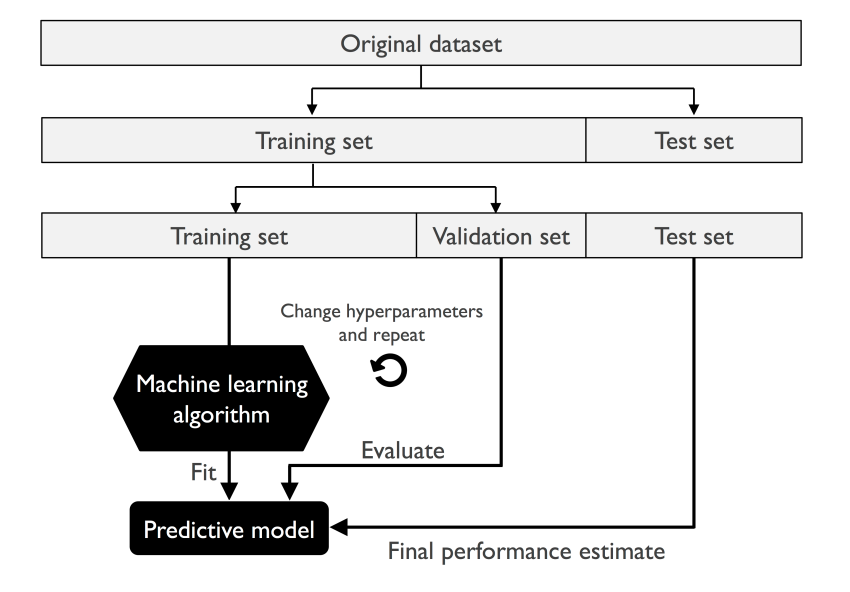
Remember we cannot use the test set.

Could we somehow mimick what we do with test data?

- Yes, we can split the remaining non-test data into training and validation data:
  - we train model for various hyperparameters on training data;
  - pick the hyperparameters which performs best on validation data.

## Model validation (2)

The non-test data is split into training and validation



#### Model validation (3)

What would this look like in Python?

```
In [57]: # splitting into development (2/3) and test data (1/3)
X_dev, X_test, y_dev, y_test = train_test_split(X, y, test_size=1/3, random_state=1)
# splitting development into train (1/3) and validation (1/3)
X_train, X_val, y_train, y_val = train_test_split(X_dev, y_dev, test_size=1/2, random_state=1)
```

#### Model validation (4)

Let's train a linear regression model

e, n jobs=1, normalize=False))])

with std=True)), ('linearregression', LinearRegression(copy X=True, fit intercept=Tru

#### Model validation (5)

Let's find the Lasso model which performs best in the validation set

0.01 17.318434 dtype: float64

#### Model validation (6)

Let's compare the performance of the Lasso vs. Linear Regression

Lasso 12.382 LinReg 17.895

#### Bias and variance (1)

How do we describe the modelling error?

From <u>Wikipedia Sunday, August 19, 2018</u> (<u>https://en.wikipedia.org/wiki/Bias%E2%80%93variance\_tradeoff</u>):

- model bias: an error from erroneous assumptions in the learning algorithm
  - oversimplification of models, cannot approximate all patterns found
- model variance: an error from sensitivity to small fluctuations in the training set
  - reacts too much to sample errors and thus finds too many spurious relations

#### Bias and variance (2)

- over fitting: low bias / high variance
  - traning our model captures all patterns but we also find some irrelevant
  - examples: Decision Trees, Support Vector Machines or Neural Networks
- under fitting: high bias / low variance
  - traning our model captures all patterns but we also find some irrelevant
  - examples: linear and logistic regression (without polynomial expansion)

### Bias and variance (3)

Not so fast.. OLS is unbiased, right?

Yes, OLS is unbiased. But ..

• Requires we know the true form of the model.

What happens if we introduce regularization? • Then model is no longer unbiased.

#### **Smarter validation**

Is this approach the smartest way for deciding on choice of hyperparameters?

#### NO

Our model choice depends a lot on which sample we pick. Could we use more of the data?

## **Cross validation**

#### The holdout method

How do we got the more out of the data?

We reuse the data in the development set repeatedly

- We test on all the data
- Rotate which parts of data is used for test and train.

#### Leave-one-out CV

How do we got the most of the data?

The most robust approach

- Each single observation in the training data we use the remaining data to train.
- Makes number of models equal to the number of observations
- Very computing intensive does not scale! LOOCV

#### K fold method (1)

How do balance computing time vs. overfitting?

We split the sample into K even sized test bins.

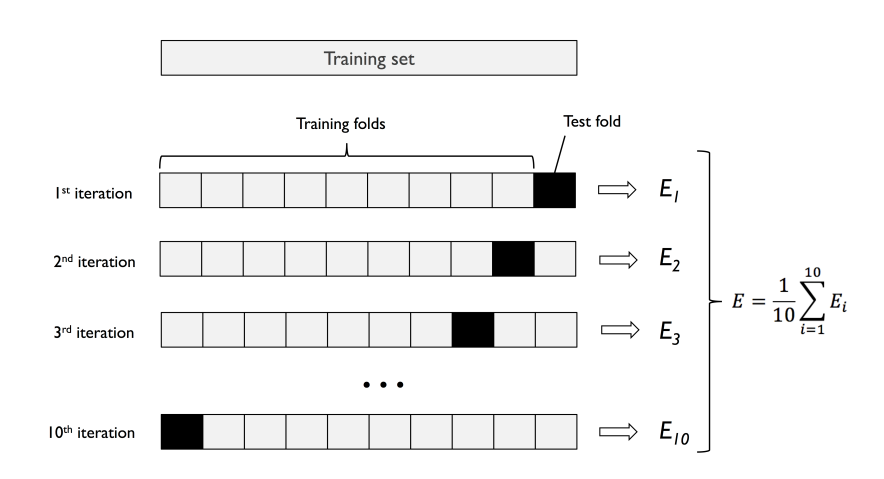
• For each test bin k we use the remaining data for training.

#### Advantages:

- We use all our data for testing.
- Training is done with 100-(100/K) pct. of the data, i.e. 90 pct. for K=10.

### K fold method (2)

In K-fold cross validation we average the errors.



#### K fold method (3)

How would we use K-fold cross validation to select our model?

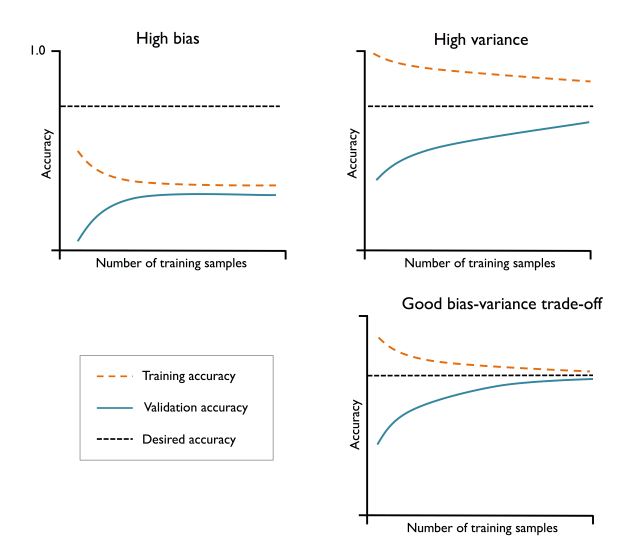
```
In [61]:
         from sklearn.model selection import KFold
         kfolds = KFold(n_splits=10)
         mseCV = []
         for lambda in lambdas:
             pipe lassoCV = make pipeline(PolynomialFeatures(degree=3, include bias=False),
                                           StandardScaler(),
                                           Lasso(alpha=lambda , random state=1))
             mseCV = []
             for train idx, val idx in kfolds.split(X dev, y dev):
                 X train, y train, = X dev[train idx], y dev[train idx]
                 X val, y val = X dev[val idx], y dev[val idx]
                  pipe lassoCV.fit(X train, y train)
                 mseCV .append(mse(pipe lassoCV.predict(X val), y val))
             mseCV.append(mseCV )
```

#### K fold method (4)

Lasso 12.4 Lasso CV 12.4 LinReg 17.9

### Learning curves (1)

What does a balanced model look like?

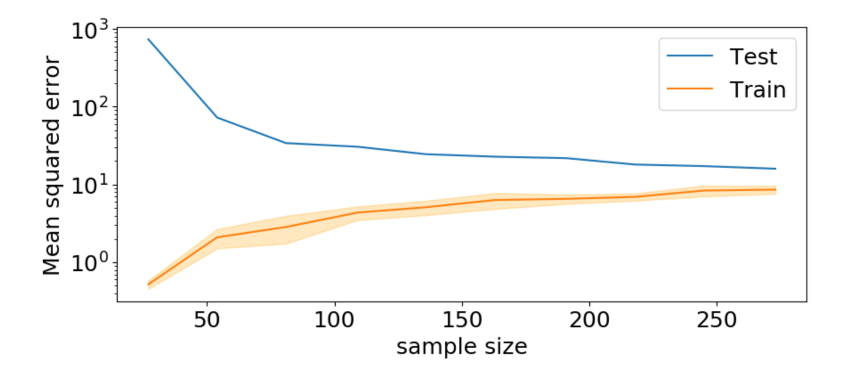


#### Learning curves (2)

	Test	Train
sample size		
20	327.856169	0.040538
40	67.607438	0.836417
60	52.208482	1.592246
80	50.194690	1.911472
101	36.249970	3.419180

Learning curves (3)

Out[38]: Text(0,0.5, 'Mean squared error')



Tools for model selection

# Validation curves (1)

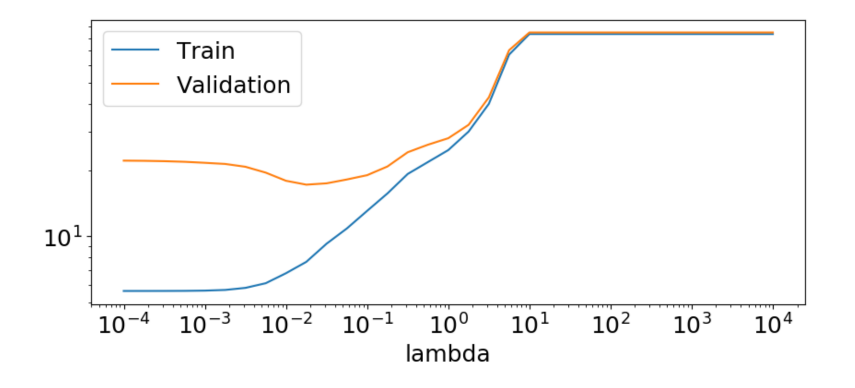
```
from sklearn.model selection import validation curve
train scores, test scores = \
    validation curve(estimator=pipe lasso,
                     X=X train,
                     y=y train,
                     param name='lasso alpha',
                     param range=lambdas,
                     scoring='neg mean squared error',
                     cv=3)
mse score = pd.DataFrame({'Train':-train scores.mean(axis=1),
                          'Validation':-test scores.mean(axis=1),
                          'lambda':lambdas})\
              .set index('lambda')
print(mse score.Test.nsmallest(1))
AttributeError
                                          Traceback (most recent call last)
<ipython-input-78-ff6698fa70dc> in <module>()
                                  'lambda':lambdas})\
     13
     14
                      .set index('lambda')
---> 15 print(mse score.Test.nsmallest(1))
~\AppData\Local\Continuum\miniconda3\envs\Python3\lib\site-packages\pandas\core\gener
ic.py in getattr (self, name)
   3612
                    if name in self. info axis:
   3613
                        return self[name]
                    return object. getattribute (self, name)
-> 3614
   3615
   3616
           def setattr (self, name, value):
AttributeError: 'DataFrame' object has no attribute 'Test'
```

In [78]:

#### Validation curves (2)

In [79]: mse\_score.plot(logx=True, logy=True)

Out[79]: <matplotlib.axes.\_subplots.AxesSubplot at 0x2910f9e4128>



#### Grid search (1)

How do we search for two or more optimal parameters?

• Goal: find the optimal parameter combination:

$$\lambda_1^*, \lambda_2^* = rg\min_{\lambda_1, \lambda_2} MSE^{CV}(X_{train}, y_{train})$$

- Option 1: We can loop over the joint grid of parameters.
  - One level for each parameter.
  - Caveats:
- Option 2: sklearn has GridSearchCV has a tool which tests all parameter combinations.

#### Grid search (2)

How does this look in Python?

- Notation: double underscore between estimator and hyperparameter, e.g. 'est\_hyperparam'
- Scoring: negative MSE as we're maximizing the score ~ minimize MSE.

#### Grid search (3)

What if we have 10,000 parameter combinations?

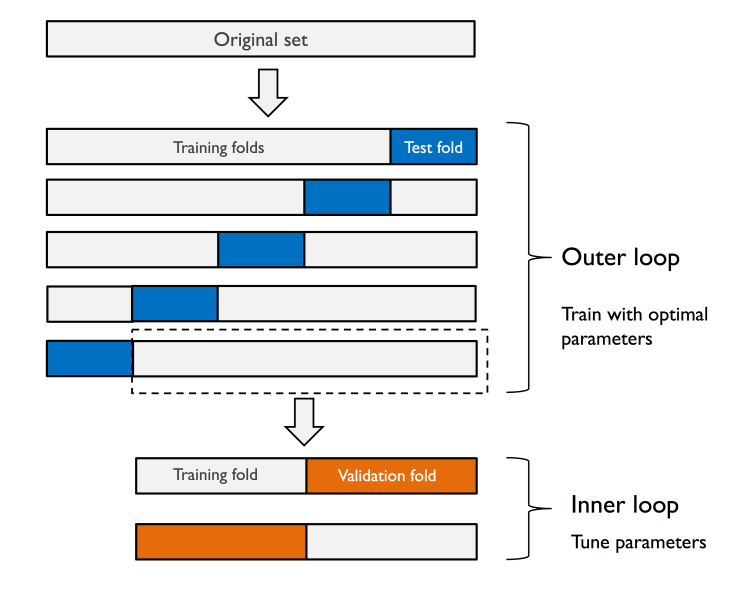
- Option 1: you buy a cluster on Amazon, learn how to parallelize across computers. Intro in last lecture.
- Option 2: you drop some of the parameter values
- Option 3: RandomizedSearchCV searches a subset of the combinations.

#### **Enhancing cross validation (1)**

- Model validation does not consider that we are also tuning hyperparameters:
  - Leads too overfitting (Varma & Simon 2006; Cawley, Talbot 2010).
- Solution is **nested cross validation**.
  - Validation step should not be modelled as 1) train; 2) test.
  - Better way is 1) model selection: train, validate; 2) test.
  - Implement as pp 204-205 in Python for Machine Learning:
    - first inner loop: GridSearchCV
    - second outer loop: cross\_val\_score

# **Enhancing cross validation (1)**

Cross-val. suffers from the fact that it models test-train



**Dimensionality reduction** 

#### Principal components analysis (1)

How can we reducing the number of features?

One solution is finding the **principal components**.

- essence: we get **fewer features** of **greater importance**.
- the new features are:
  - uncorrelated (i.e. linearly independent, orthogonal)
  - ordered so decreasing in how much variation of the feature data they explain

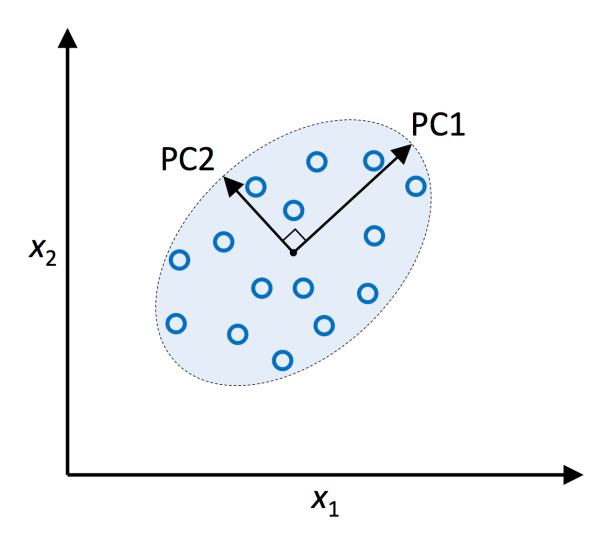
The method is called **principal components analysis** 

- corresponds to eigen decomposition of matrix into
  - principal eigenvectors (factors)
  - principal eigenvalues (factor importance)

## Principal components analysis (2)

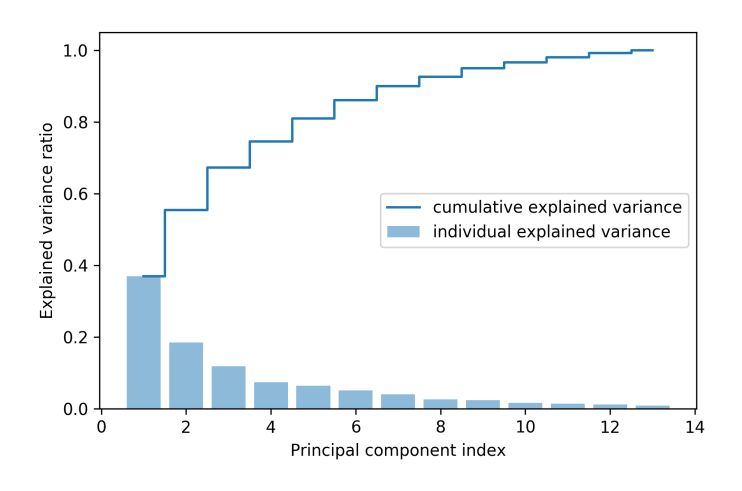
Finding principal components for two features. Notice:

- The factors are orthogonal
- The first factor explains more variation, |PC1| > |PC2|



## Principal components analysis (3)

We can plot the explained variation against the component indices, often called scree plot.



## Principal components analysis (4)

How do we choose the number of components?

• Standard is to look for an *elbow* in the previous scree plot.

What might go wrong about this approach??

• The number of feautures should be a hyperparameter in the model building!!!

## Principal components analysis (5)

How does this look in Python?

Out[22]: {'lasso\_alpha': 0.1778279410038923, 'pca\_n\_components': 13}

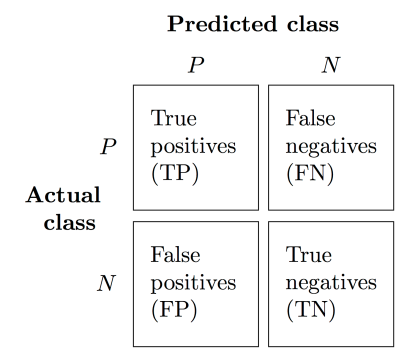
**Measures for classification** 

#### Breakdown by error type (1)

We measure the accaracy as the rate of true predictions, i.e.

$$ACC = rac{TP + TN}{TP + TN + FP + FN} = rac{True}{True + False}$$

where our measures are



# Breakdown by error type (2)

Some powerful measures:

• Precision: share of predicted positive that are true

■ PRE = 
$$\frac{TP}{TP+FP}$$

• Recall: share of actual positive that are true

• REC = 
$$\frac{TP}{TP+FN} = \frac{TP}{AP}$$

- Also known as True Positive Rate, TPR
- F1: mix recall and precision:  $\frac{2 \cdot PRE \cdot REC}{PRE \cdot REC}$
- False Positive Rate: share of actual negatives that are true

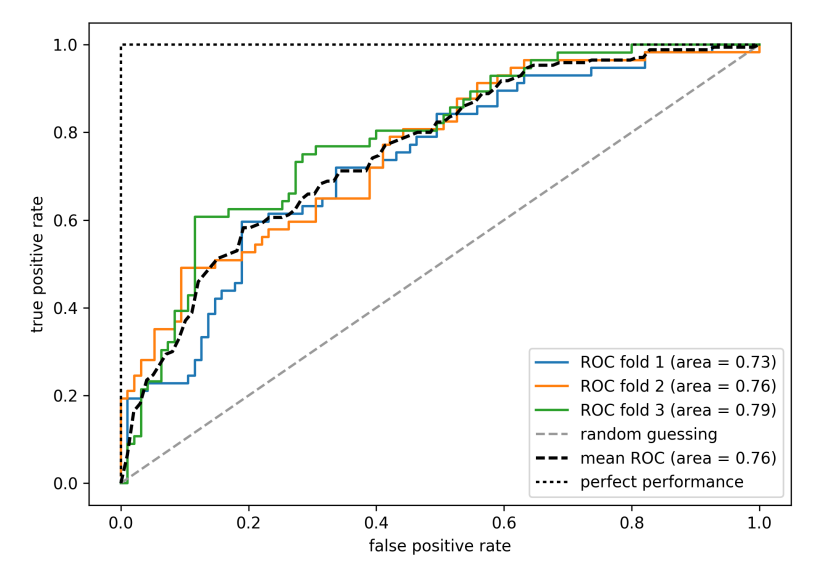
■ 
$$FPR = \frac{FP}{FP + TN} = \frac{FP}{AN}$$

In [ ]: from sklearn.metrics import precision\_score, recall\_score, f1\_score

## Breakdown by error type (3)

Classification models provide a predicted likelihood of being in the class or not:

- Receiver Operating Characteristic (ROC) curve by varying thresholds for predicted true.
  - ROC is a *theoretical* measure of model performance based on probabilities.
  - AUC: Area Under the (ROC) Curve.



# The end

Return to agenda