

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

Vaaaamos

```
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

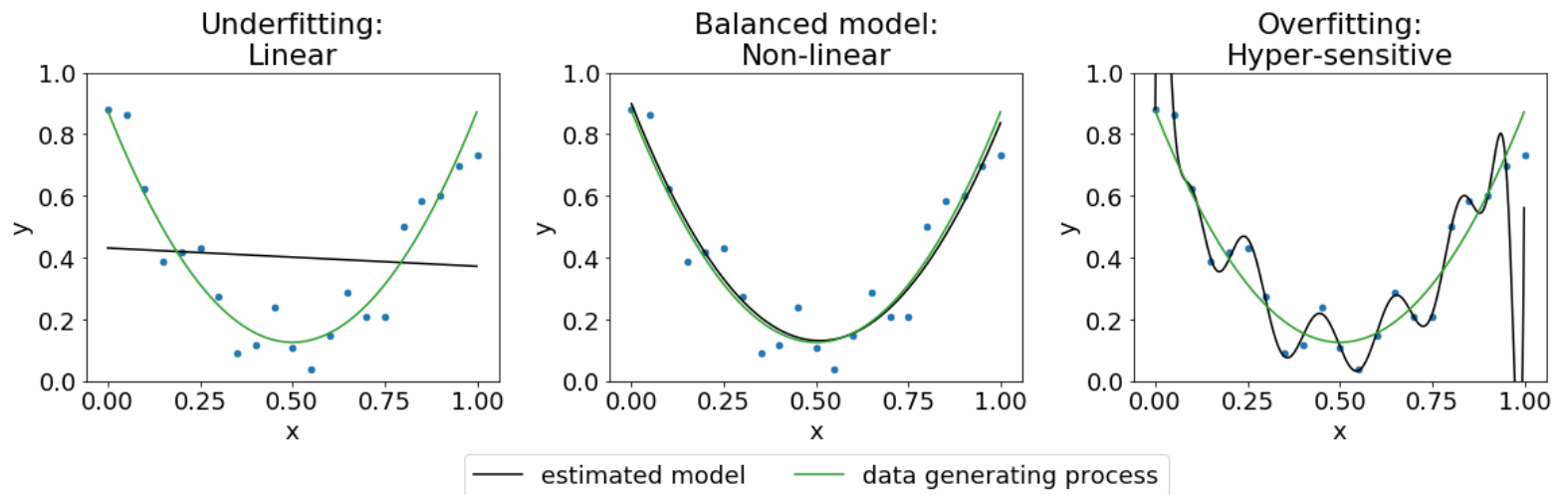
plt.style.use('default') # set style (colors, background, size, gridlines etc.)
plt.rcParams['figure.figsize'] = 10, 4 # set default size of plots
plt.rcParams.update({'font.size': 18})
```

Supervised problems (1)

What is the tradeoff for making supervised regression models?

```
In [3]: # f_bias_var['regression'] [2]
```

Out[3]:



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

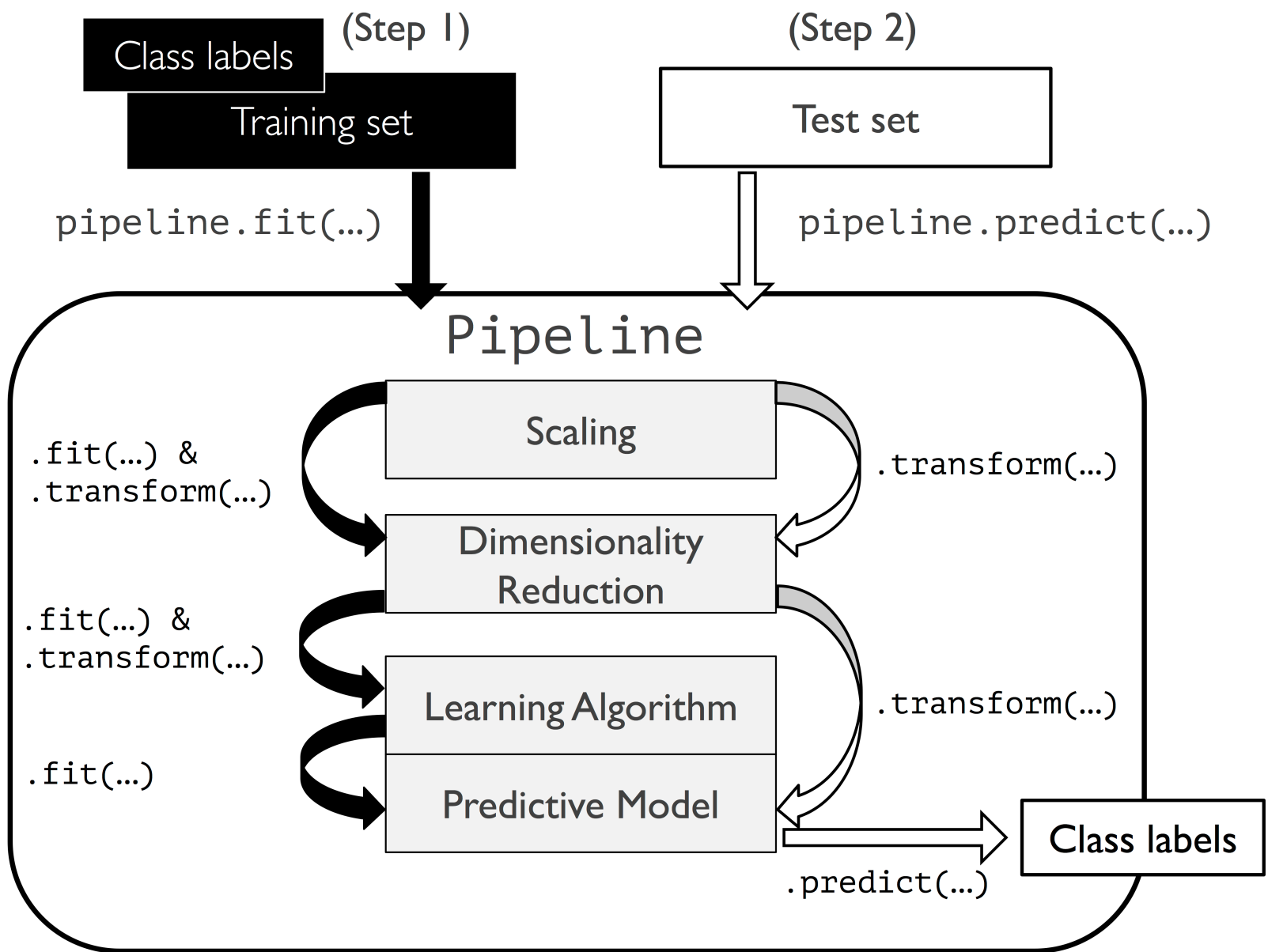
Model pipelines (1)

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?

```
In [55]: from sklearn.pipeline import make_pipeline
         from sklearn.preprocessing import PolynomialFeatures, StandardScaler

         pipe_preproc = make_pipeline(PolynomialFeatures(),
                                     StandardScaler())

         print(pipe_preproc.steps[0])
         print(pipe_preproc.steps[1])

('polynomialfeatures', PolynomialFeatures(degree=2, include_bias=True, interaction_on
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):

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

Applying a model pipeline (3)

And how do I apply the pipe on the data?

```
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'
          'B' 'LSTAT']
```

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. λ for Lasso, Ridge.

Model validation (1)

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

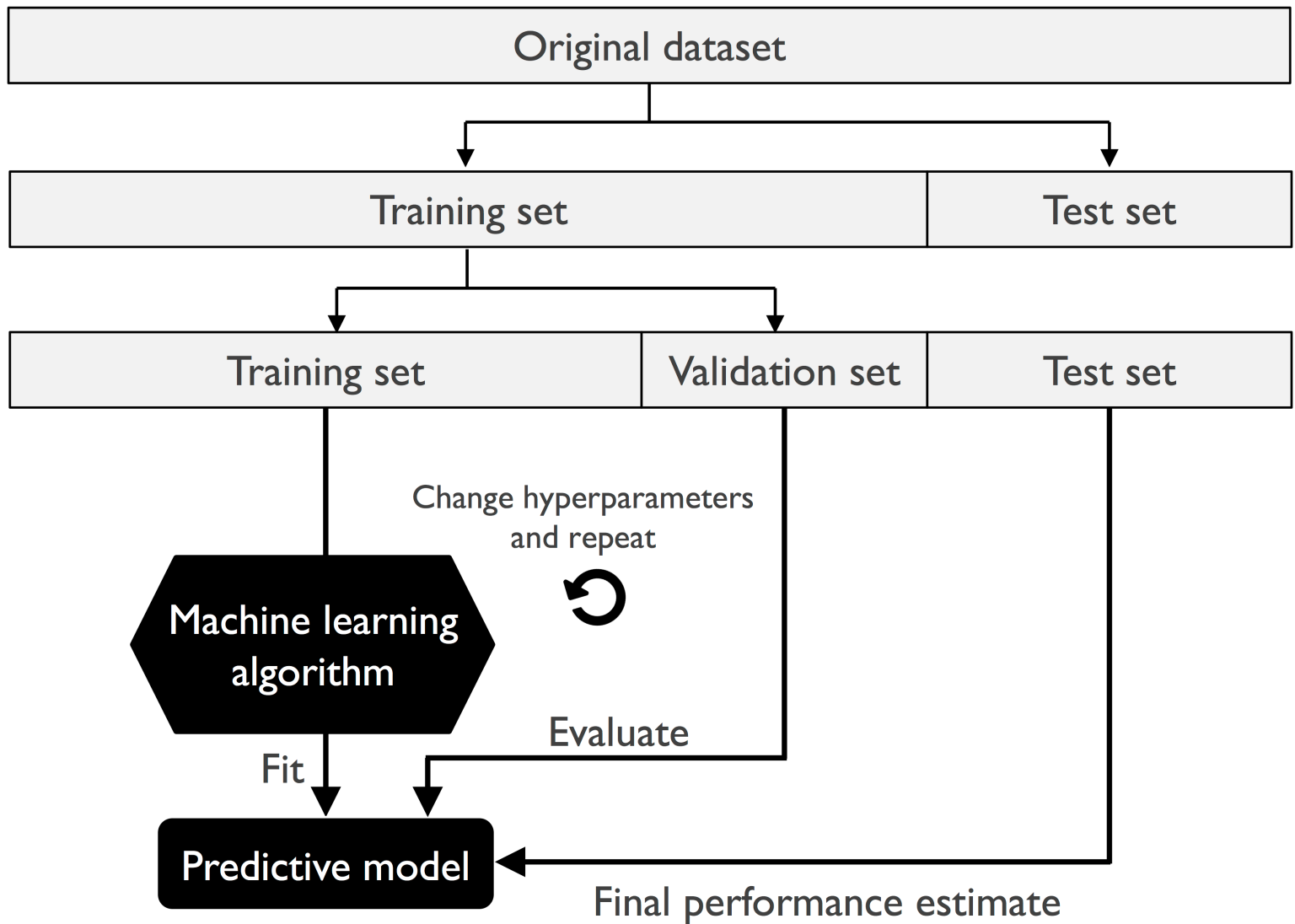
- Remember we cannot use the test set.

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

```
In [7]: from sklearn.linear_model import Lasso, LinearRegression
pipe_lr = make_pipeline(PolynomialFeatures(include_bias=False),
                        StandardScaler(),
                        LinearRegression())

pipe_lr.fit(X_dev, y_dev)
```

```
Out[7]: Pipeline(memory=None,
      steps=[('polynomialfeatures', PolynomialFeatures(degree=2, include_bias=False, i
nteraction_only=False)), ('standardscaler', StandardScaler(copy=True, with_mean=True,
with_std=True)), ('linearregression', LinearRegression(copy_X=True, fit_intercept=Tru
e, n_jobs=1, normalize=False))])
```

Model validation (5)

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

```
In [59]: from sklearn.metrics import mean_squared_error as mse

perform = []
lambdas = np.logspace(-4, 4, 33)
for lambda_ in lambdas:
    pipe_lasso = make_pipeline(PolynomialFeatures(include_bias=False),
                               StandardScaler(),
                               Lasso(alpha=lambda_, random_state=1))
    pipe_lasso.fit(X_train, y_train)
    y_pred = pipe_lasso.predict(X_val)
    perform.append(mse(y_pred, y_val))

hyperparam_perform = pd.Series(perform, index=lambdas)

optimal = hyperparam_perform.nsmallest(1)
print(optimal)
```

```
0.01    17.318434
dtype: float64
```

Model validation (6)

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

```
In [60]: pipe_lasso = make_pipeline(PolynomialFeatures(include_bias=False),  
                                   StandardScaler(),  
                                   Lasso(alpha=optimal.index[0]))  
  
pipe_lasso.fit(X_dev,y_dev)  
print('Lasso', round(mse(pipe_lasso.predict(X_test),y_test), 3))  
print('LinReg', round(mse(pipe_lr.predict(X_test),y_test), 3))
```

Lasso 12.382

LinReg 17.895

Bias and variance (1)

How do we describe the modelling error?

From Wikipedia Sunday, August 19, 2018

(https://en.wikipedia.org/wiki/Bias%E2%80%93variance_tradeoff):

- 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
 - training 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
 - training 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 get the most 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 get 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.

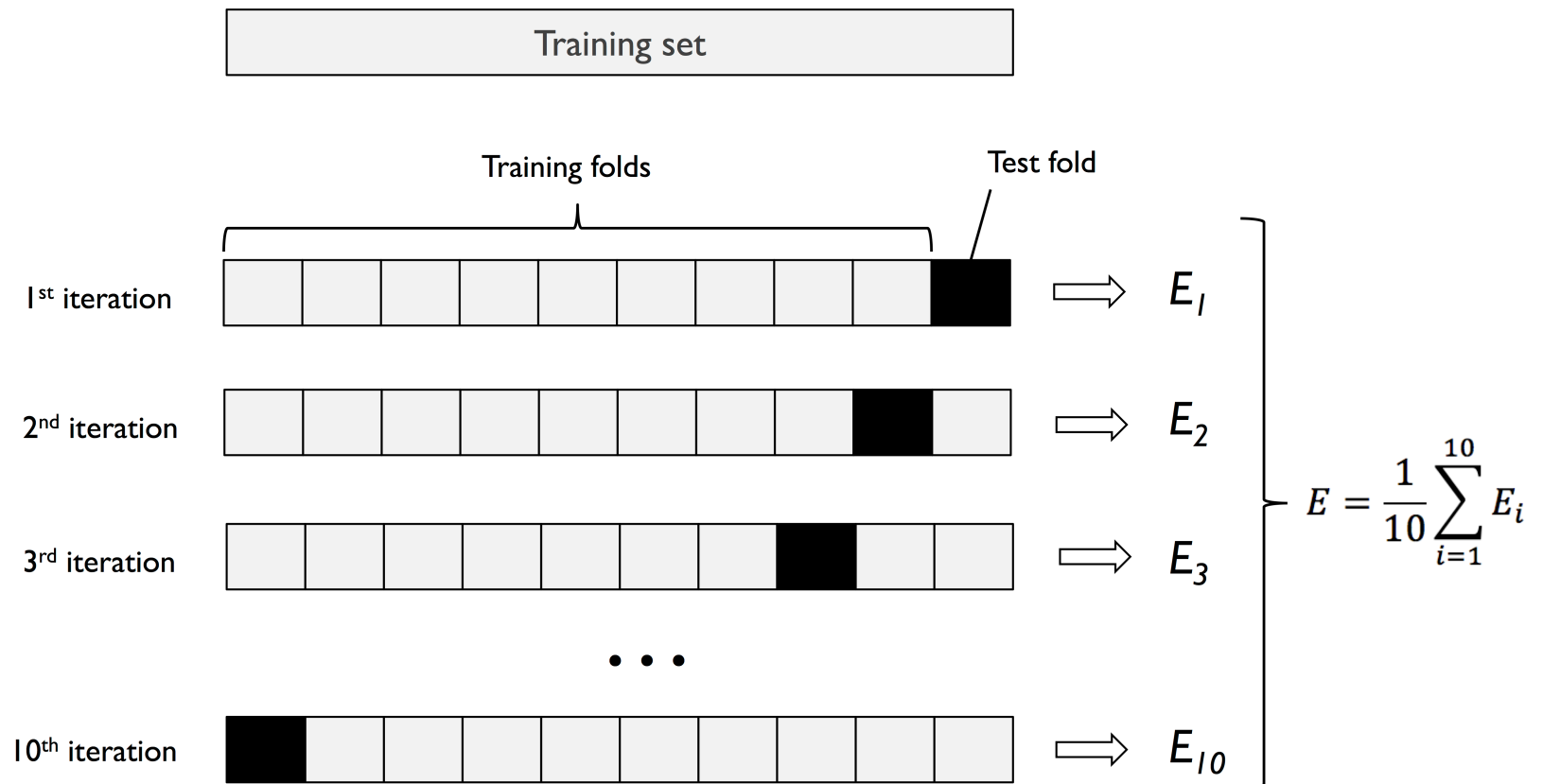
- For each test bin k we use the remaining data for training.

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)

```
In [70]: optimalCV = pd.DataFrame(mseCV, index=lambdas).mean(axis=1).nsmallest(1)

pipe_lassoCV = make_pipeline(PolynomialFeatures(include_bias=False),
                             StandardScaler(),
                             Lasso(alpha=optimalCV.index[0], random_state=1))

pipe_lassoCV.fit(X_dev,y_dev)

model_pipes = ('Lasso', pipe_lasso), ('Lasso CV', pipe_lassoCV), ('LinReg', pipe_lr)

for model_name, model_pipe in model_pipes:
    score = mse(model_pipe.predict(X_test),y_test)
    print(model_name, round(score, 1))
```

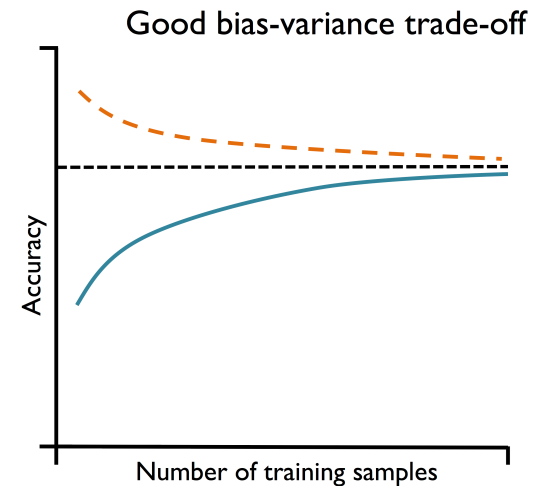
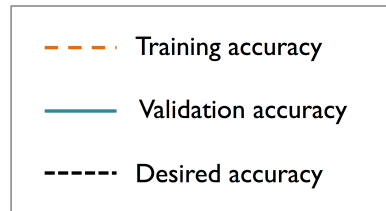
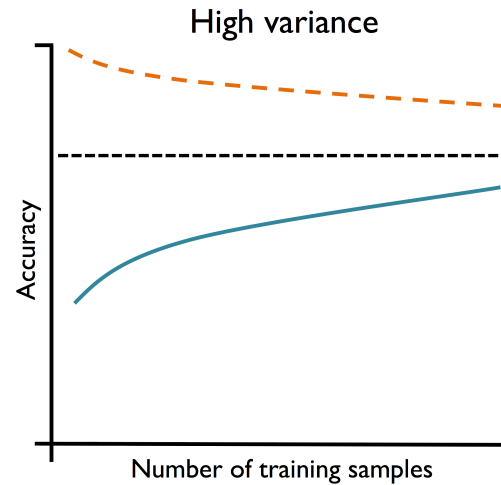
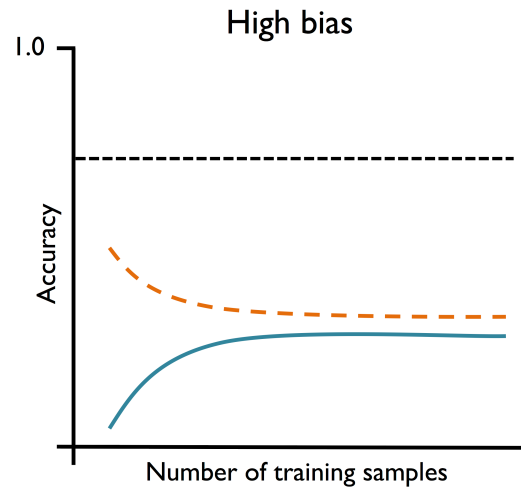
Lasso 12.4

Lasso CV 12.4

LinReg 17.9

Learning curves (1)

What does a balanced model look like?



Learning curves (2)

```
In [74]: from sklearn.model_selection import learning_curve

train_sizes, train_scores, test_scores = \
    learning_curve(estimator=pipe_lasso,
                  X=X_train,
                  y=y_train,
                  train_sizes=np.linspace(0.1, 1.0, 10),
                  scoring='neg_mean_squared_error',
                  cv=3)

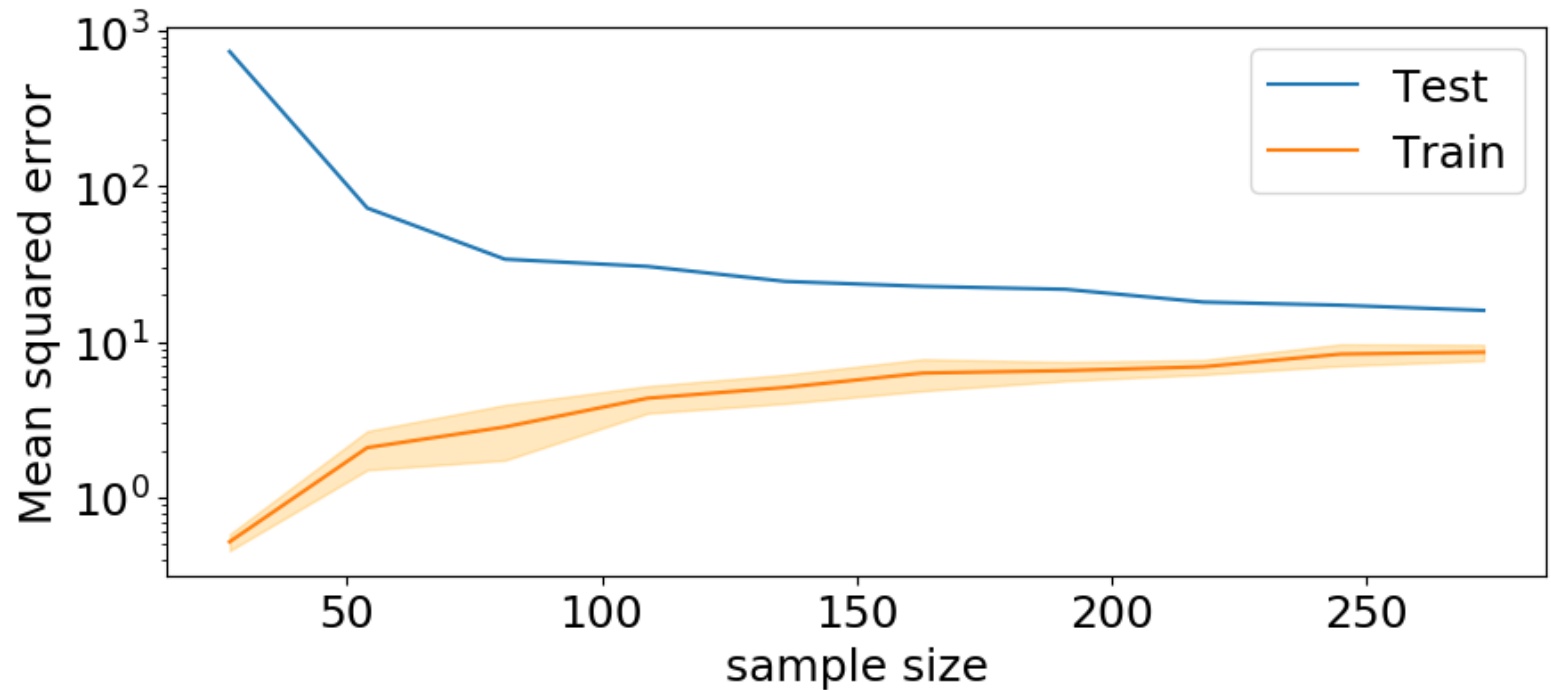
mse_ = pd.DataFrame({'Train':-train_scores.mean(axis=1),
                    'Test':-test_scores.mean(axis=1)})\
    .set_index(pd.Index(train_sizes,name='sample size'))
print(mse_.head())
```

	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)

```
In [38]: f_learn, ax = plt.subplots(figsize=(10,4))
mse_.plot(ax=ax, logy=True)
ax.fill_between(train_sizes,
               -train_scores.mean(1) + train_scores.std(1)*1.96,
               -train_scores.mean(1) - train_scores.std(1)*1.96,
               alpha=0.25,
               color='orange')
ax.set_ylabel('Mean squared error')
```

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



Tools for model selection

Validation curves (1)

```
In [78]: 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=lamdbas,
                    scoring='neg_mean_squared_error',
                    cv=3)

mse_score = pd.DataFrame({'Train':-train_scores.mean(axis=1),
                        'Validation':-test_scores.mean(axis=1),
                        'lambda':lamdbas})\
    .set_index('lambda')
print(mse_score.Test.nsmallest(1))
```

```
-----
AttributeError                                Traceback (most recent call last)
```

```
<ipython-input-78-ff6698fa70dc> in <module>()
    13         'lambda':lamdbas})\
    14         .set_index('lambda')
--> 15 print(mse_score.Test.nsmallest(1))
```

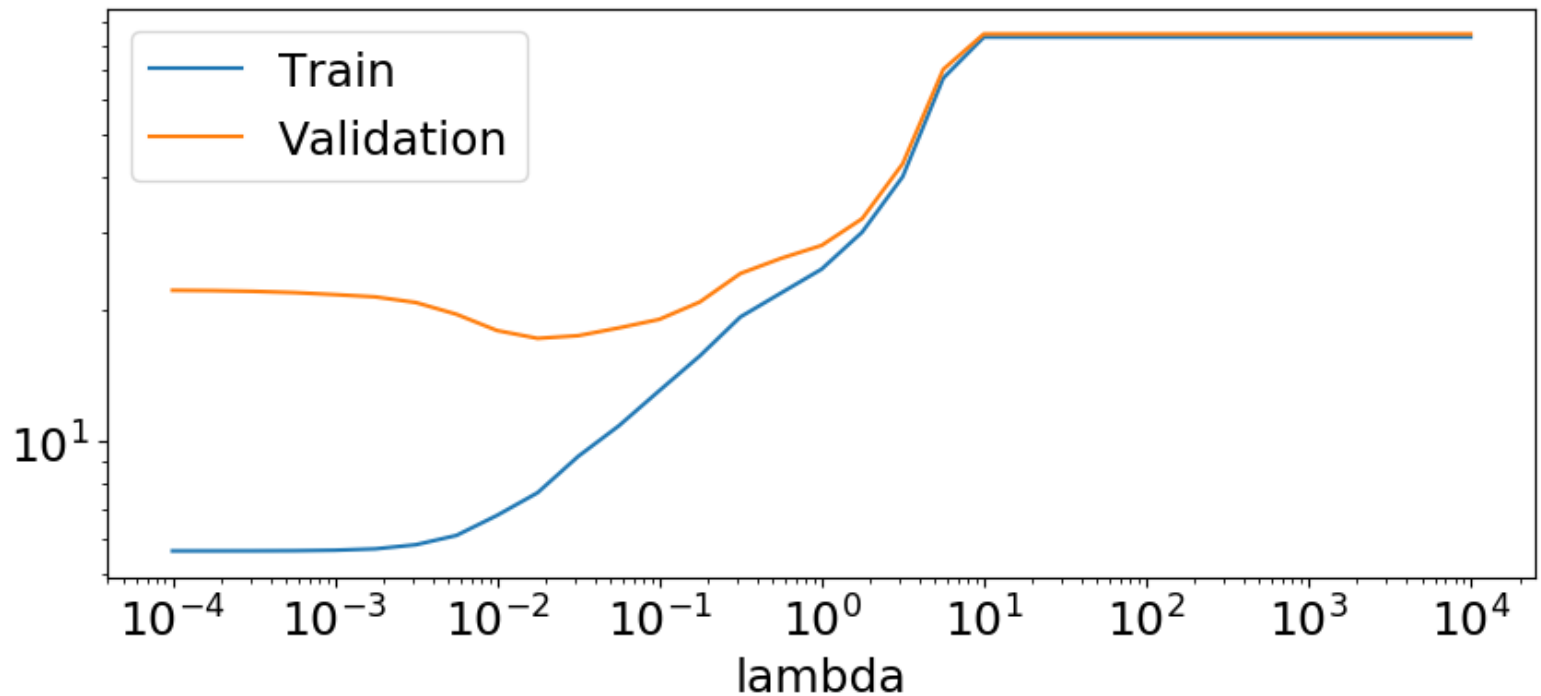
```
~\AppData\Local\Continuum\miniconda3\envs\Python3\lib\site-packages\pandas\core\gener
ic.py in __getattr__(self, name)
    3612         if name in self._info_axis:
    3613             return self[name]
-> 3614         return object.__getattribute__(self, name)
    3615
    3616     def __setattr__(self, name, value):
```

```
AttributeError: 'DataFrame' object has no attribute 'Test'
```

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:

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

- 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?

```
In [77]: from sklearn.model_selection import GridSearchCV

gs = GridSearchCV(estimator=pipe_lasso,
                  param_grid={'lasso__alpha': lambdas},
                  scoring='neg_mean_squared_error',
                  cv=10)

gs = gs.fit(X_train, y_train)
gs.best_params_
```

```
Out[77]: {'lasso__alpha': 0.01}
```

- 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?

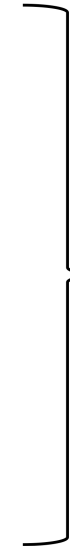
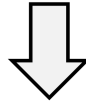
- Option 1: you buy a cluster on Amazon, learn how to parallelize across computers. Intro in last lecture.
- Option 2: you drop some of the parameter values
- Option 3: RandomizedSearchCV searches a subset of the combinations.

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

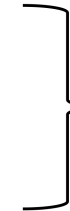
Enhancing cross validation (1)

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



Outer loop

Train with optimal
parameters



Inner loop

Tune parameters

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

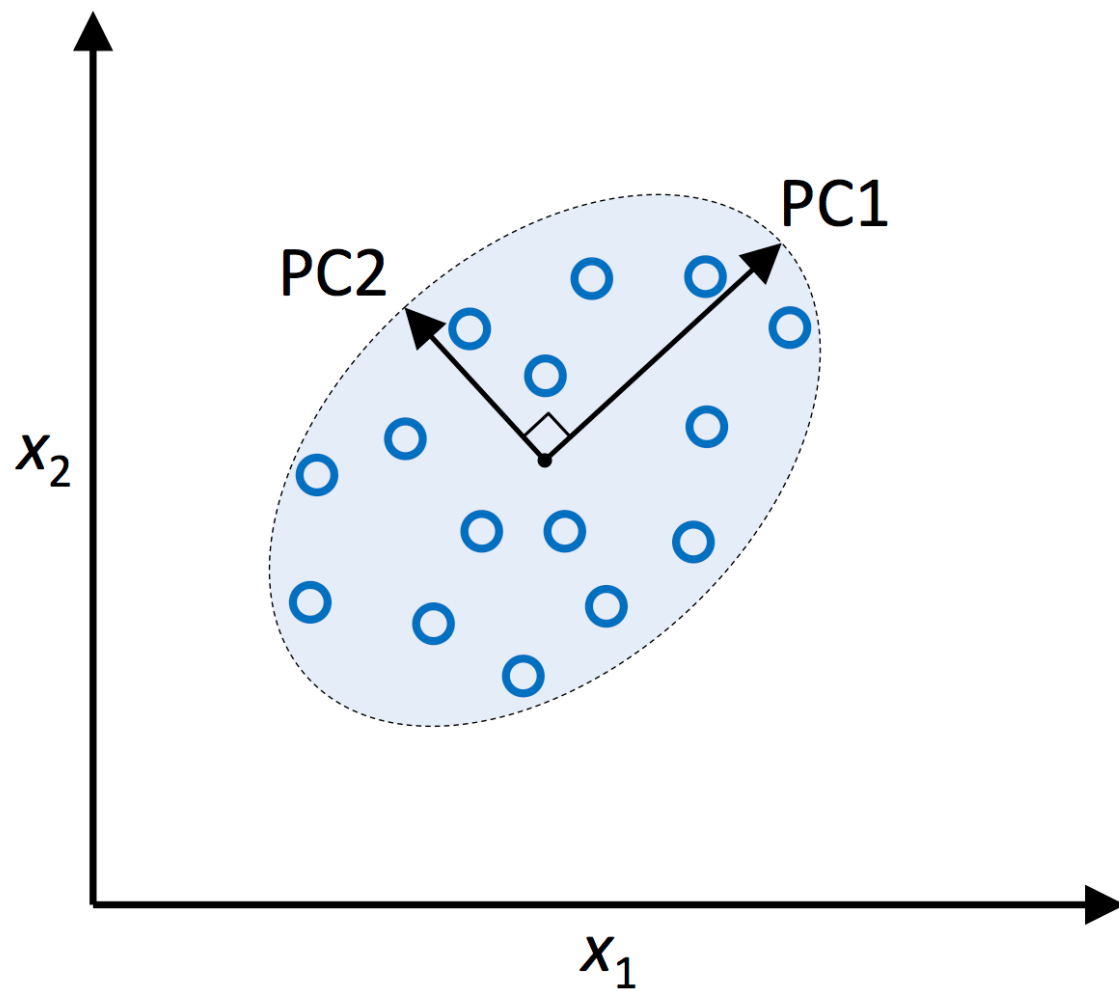
The method is called **principal components analysis**

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

Principal components analysis (2)

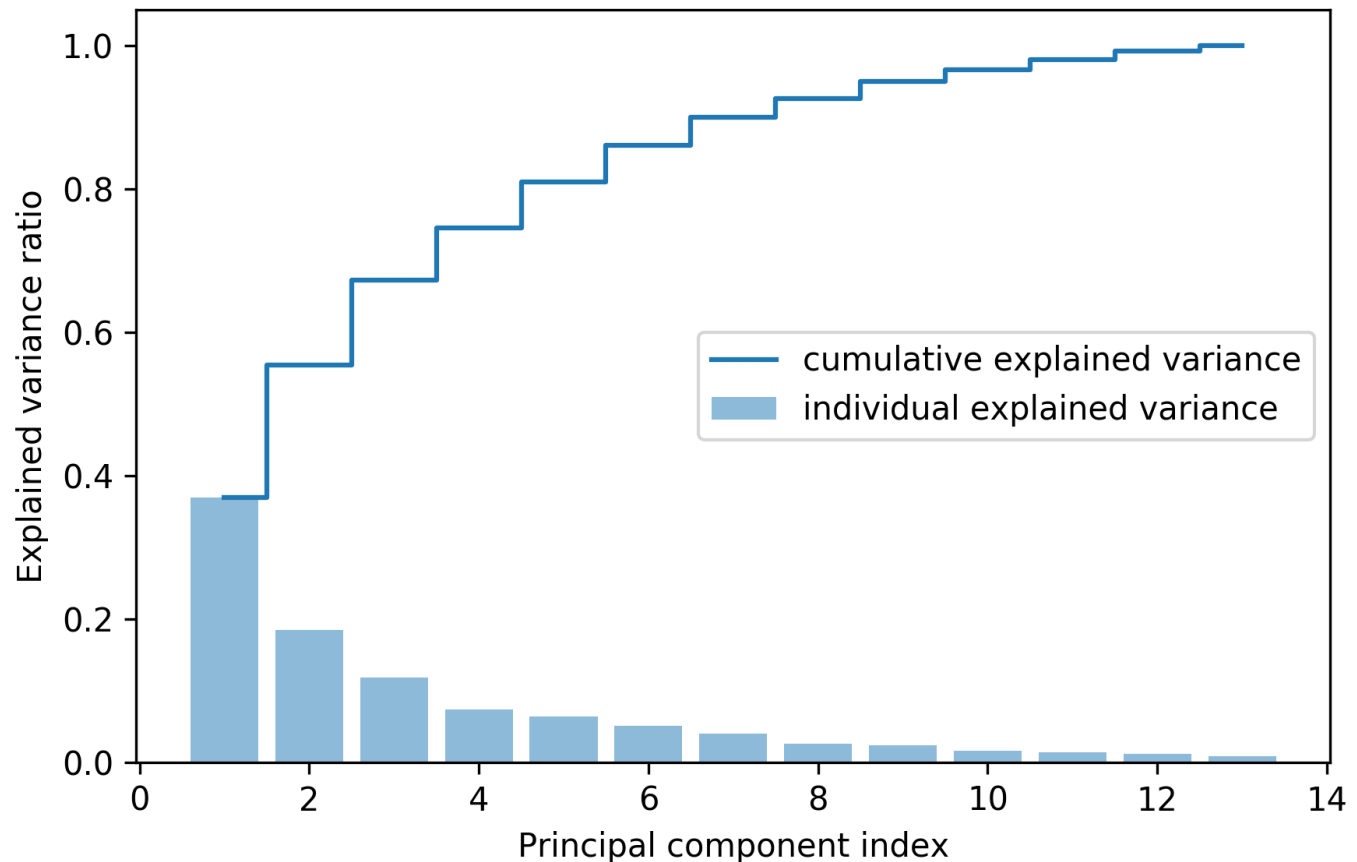
Finding principal components for two features. Notice:

- 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 features should be a hyperparameter in the model building!!!

Principal components analysis (5)

How does this look in Python?

```
In [22]: from sklearn.decomposition import PCA

pipe_pca_lasso = make_pipeline(PolynomialFeatures(include_bias=False),
                               StandardScaler(),
                               PCA(),
                               Lasso())

gs = GridSearchCV(estimator=pipe_pca_lasso,
                  param_grid={'lasso__alpha': lambdas,
                              'pca__n_components': range(1, X_train.shape[1]+1)},
                  scoring='neg_mean_squared_error',
                  cv=10)

gs = gs.fit(X_train, y_train)
gs.best_params_
```

```
Out[22]: {'lasso__alpha': 0.1778279410038923, 'pca__n_components': 13}
```

Measures for classification

Breakdown by error type (1)

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

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

where our measures are

		Predicted class	
		<i>P</i>	<i>N</i>
Actual class	<i>P</i>	True positives (TP)	False negatives (FN)
	<i>N</i>	False positives (FP)	True negatives (TN)

Breakdown by error type (2)

Some powerful measures:

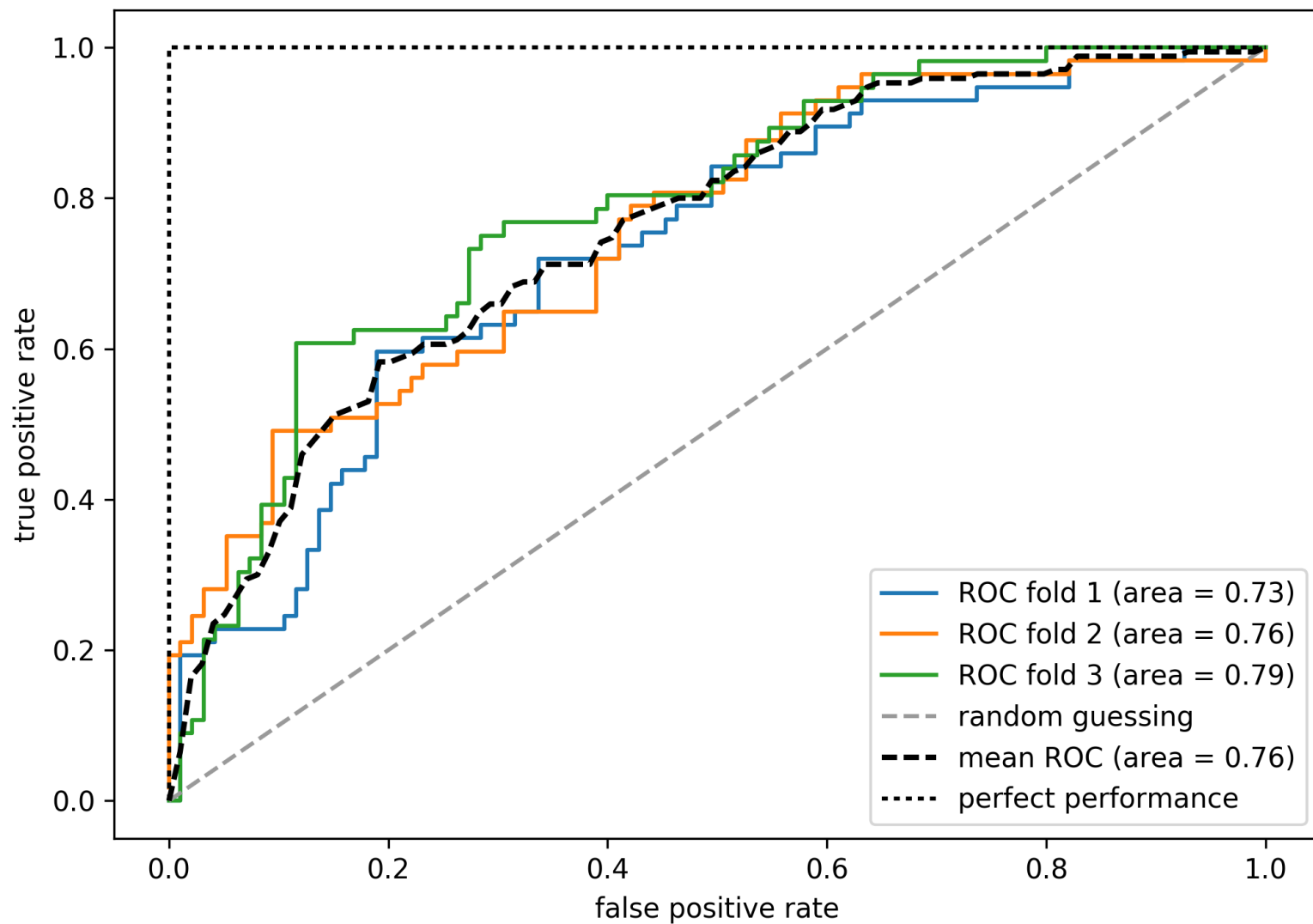
- Precision: share of predicted positive that are true
 - $PRE = \frac{TP}{TP+FP}$
- Recall: share of actual positive that are true
 - $REC = \frac{TP}{TP+FN} = \frac{TP}{AP}$
 - Also known as True Positive Rate, TPR
- F1: mix recall and precision: $\frac{2 \cdot PRE \cdot REC}{PRE + REC}$
- False Positive 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](#)