Unit 10: Introduction to supervised learning

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10 Introduction to supervised learning

10.1 Simple (univariate) linear regression

Imagine the simplest linear model where the dependent variable y is assumed to be an affine function of the explanatory variable x and an error term ϵ , given by

$$y_i = \alpha + \beta x_i + \epsilon_i$$

for each observation i. In econometrics, the parameters α and β are called the intercept and slope parameters, respectively. In machine learning, the terminology often differs and you might see a simple linear model written like

$$y_i = b + wx_i + \epsilon_i$$

where *b* is called the *bias* and *w* is called a *weight*.

Our goal is to estimate the parameters α and β which is most commonly done by ordinary least squares (OLS). OLS is defined as the estimator that finds the estimates $(\widehat{\alpha}, \widehat{\beta})$ such that the sum of squared errors is minimized,

$$L(\alpha, \beta) = \frac{1}{N} \sum_{i}^{N} (y_i - \alpha - \beta x_i)^2$$

where L is the loss function that depends on the choice of parameters. Note that we use the "hat" notation $\widehat{\alpha}$ to distinguish the OLS estimate from the (usually unknown) true parameter α . The exact values of $\widehat{\alpha}$ and $\widehat{\beta}$ will vary depending the sample size and estimator used as we will see later in this unit.

For this simple model, the estimates are given by the expressions

$$\widehat{\beta} = \frac{\widehat{Cov}(y, x)}{\widehat{Var}(x)}$$

$$\widehat{\alpha} = \overline{y} - \widehat{\beta}\overline{x}$$

where $\widehat{Cov}(\bullet, \bullet)$ and $\widehat{Var}(\bullet)$ are the *sample* covariance and variance, respectively, and \overline{y} and \overline{x} are the sample means of y and x.

There is a straightforward generalization to the multivariate setting where we have a vector \mathbf{x}_i of explanatory variables (which usually include the intercept) and a parameter vector $\boldsymbol{\beta}$ so that the model is given by

$$y_i = \mathbf{x}_i' \boldsymbol{\beta} + \epsilon_i$$

If we stack all x_i in the matrix X and all y_i in the vector y, the OLS estimate of β is given by the well-known formula

$$\widehat{\boldsymbol{\beta}} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y}$$

However, we will not be estimating linear regressions based on this formula and you should never attempt this, as naively implementing such matrix operations can lead to numerical problems. Instead, always use pre-packaged least-squares solvers such as those implemented in NumPy's lstsq() or SciPy's lstsq() functions. For econometrics and machine learning, it usually makes sense to use linear regression models such as those implemented in statsmodels or scikit-learn, which is what we turn to next.

10.1.1 Univariate linear regressions with scikit-learn

We start by fitting the simple linear model

$$y_i = \alpha + \beta x_i + \epsilon_i$$

using scikit-learn. For now, we proceed using synthetically generated data where we know the true relationship, assuming that

$$y_i = 1 + \frac{1}{2}x_i + \epsilon_i$$
$$\epsilon \stackrel{\text{iid}}{\sim} N(0, 0.7^2)$$

so that the true parameters are $\alpha = 1$ and $\beta = \frac{1}{2}$. The error term ϵ is assumed to be normally distributed with mean 0 and variance $0.7^2 = 0.49$. Note that it is customary to specify the normal distribution in terms of its variance (here 0.49), but most NumPy and SciPy functions expect the standard deviation (here 0.7) to be passed as the scale parameter of a normal distribution.

We generate a sample of N = 30 observations as follows:

```
[1]: import numpy as np
from numpy.random import default_rng

rng = default_rng(123)

# Number of observations
```

```
N = 30

# True parameters
alpha = 1.0
beta = 0.5

# Use x that are uniformly spaced on [0, 10]
x = np.linspace(0, 10, N)
# Normally distributed errors
epsilon = rng.normal(scale=0.7, size=N)
# Create outcome variable
y = alpha + beta * x + epsilon
```

To fit a linear model, we use the LinearRegression class provided by scikit-learn. Before doing so, we need to import it from sklearn.linear_model. Note that most fitting routines in scikit-learn expect a *matrix* as opposed to a vector even if the model has only one explanatory variable, so we need to insert an artificial axis to create the matrix X = x[:, None].

```
from sklearn.linear_model import LinearRegression

# Create LinearRegression object
lr = LinearRegression(fit_intercept=True)

# fit() expects two-dimensional object, convert to N x 1 matrix
X = x[:, None]

# Fit model to data
lr.fit(X, y)

# Predict fitted values
yhat = lr.predict(X)
```

The following code visualises the sample as a scatter plot of (x_i, y_i) and adds the fitted line (in solid orange). Intuitively, for x = 0 the fitted line has the value $\widehat{\alpha}$ (the intercept) and its slope is equal to $\widehat{\beta}$. Note that scikit-learn models usually store the fitted model parameters in the coef_ attribute (which is an array even if there is only a single explanatory variable). If the model includes an intercept, its fitted value is stored in the attribute intercept_.

```
[3]: import matplotlib.pyplot as plt
     # Extract parameter estimates from LinearRegression object
     alpha_hat = lr.intercept_
     beta_hat = lr.coef_[o]
     fig, ax = plt.subplots(1, 1, figsize=(5, 3.5))
     # Plot true relationship
     ax.axline((0.0, alpha), slope=beta, lw=1.0, ls='--', c='black',
         label=r'$y = \alpha + \beta x
     # Plot regression line
     ax.axline((0.0, alpha_hat), slope=beta_hat, lw=1.0, c='darkorange',
         label=r'\widehat{\pha} + \widehat{\beta} x$')
     # Plot raw data
     ax.scatter(x, y, s=20, color='none', edgecolor='steelblue', alpha=1.0,
         lw=0.75, label='y')
     # Plot lines connecting true and predicted values for
     # each observation
     for i in range(len(x)):
         # Predict yhat for given x_i
```

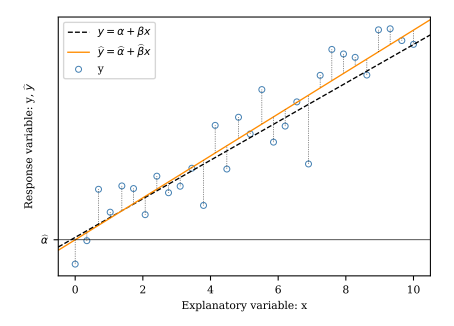
```
yhat_i = alpha_hat + beta_hat * x[i]
    ax.plot([x[i], x[i]], [y[i], yhat_i], lw=0.5, ls=':',
        c='black', alpha=0.9)

# Add annotations
ax.set_xlabel('Explanatory variable: x')
ax.set_ylabel(r'Response variable: y, $\widehat{y}$')
ax.axhline(alpha_hat, lw=0.5, c='black')
ax.legend(loc='best')
ax.set_yticks((alpha_hat, ), (r'$\widehat{\alpha}$', ))

fig.suptitle('Univariate linear regression')
```

[3]: Text(0.5, 0.98, 'Univariate linear regression')

Univariate linear regression



The dashed black line shows the true relationship which is known in this synthetic sample, but in general is unknown for real data. For small sample sizes, the true and estimated parameters need not be close as the graph illustrates. The graph also shows the prediction errors as the dotted lines between the sample points y_i and the black fitted model. OLS minimizes the sum of the squares of these distances.

10.1.2 Training and test samples

In econometrics, we usually emphasise *inference*, i.e., we are interested in testing a hypothesis about the estimated parameter, for example whether it is statistically different from zero. Conversely, in machine learning the emphasis is often on prediction, i.e., our goal is to estimate a relationship from a *training* sample and make predictions for new data. Usually, we use a *test* sample that is different from the training data to assess how well our model is able to predict outcomes for new data which has not been used for estimation.

To demonstrate the use of training and test data sets, we use as simplified variant of the Ames house price data set which can be obtained from openml.org, a repository for free-to-use data suitable for machine learning tasks.

The original data set has 80 features (explanatory variables) which are characteristics of houses in Ames, a city of about 60 thousand inhabitants in the middle of Iowa, USA. The goal is to use these features to

predict the house price (or "target" in ML terminology). The above website provides details on all 80 features, but we restrict ourselves to a small subset.

We could download the data set directly from openml.org using scikit-learn as follows:

```
from sklearn.datasets import fetch_openml
ds = fetch_openml(name='house_prices')

X = ds.data  # Get features (explanatory variables)
y = ds.target  # get dependent variable
```

which returns an object with various information about the data set. The features are stored in the data attribute, while the dependent variable is stored in the target attribute.

Instead, we will use a local copy of the simplified data set which contains only a subset of 12 features that are slightly adapted for our purposes.

```
[4]: # Uncomment this to use files in the local data/ directory
DATA_PATH = '../data'

# Load data directly from GitHub (for Google Colab)
# DATA_PATH = 'https://raw.githubusercontent.com/richardfoltyn/python-intro-PGR/main/data'
```

```
[5]: import pandas as pd

file = f'{DATA_PATH}/ames_houses.csv'
    df = pd.read_csv(file)

# List columns present in DataFrame
    df.info()
```

```
RangeIndex: 1460 entries, 0 to 1459
Data columns (total 13 columns):
# Column
                    Non-Null Count Dtype
                   1460 non-null float64
0
    SalePrice
    LotArea
                    1460 non-null float64
1
    Neighborhood 1460 non-null object
BuildingType 1386 non-null object
3
    OverallQuality 1460 non-null int64
    OverallCondition 1460 non-null int64
    YearBuilt 1460 non-null int64
6
    CentralAir
                    1460 non-null object
7
    CentralAir
LivingArea
8
                    1460 non-null
                                    float64
    Bathrooms
9
                    1460 non-null
                                     int64
                    1460 non-null
10 Bedrooms
                                     int64
11 Fireplaces
                     1460 non-null
                                     int64
12 HasGarage
                     1460 non-null
                                     int64
dtypes: float64(3), int64(7), object(3)
memory usage: 148.4+ KB
```

<class 'pandas.core.frame.DataFrame'>

In this section, we focus on the columns SalePrice which contains the house price in US dollars and LivingArea which contains the living area in m^2 for each house in the sample. To get some intuition for the data, we look at some descriptive statistics:

```
[6]: df[['SalePrice', 'LivingArea']].describe()

[6]: SalePrice LivingArea
count 1460.000000 1460.000000
mean 180921.195890 140.777444
std 79442.502883 48.813961
min 34900.000000 31.026587
```

```
25% 129975.000000 104.923743
50% 163000.000000 135.996777
75% 214000.000000 165.049367
max 755000.000000 524.107798
```

For our analysis, we convert the DataFrame columns to NumPy arrays because scikit-learn was not programmed to work with pandas DataFrames:

```
[7]: features = ['LivingArea']
  target = 'SalePrice'

# Convert DataFrame to NumPy arrays
  y = df[target].to_numpy()
  X = df[features].to_numpy()
```

Manually creating training and test samples

We next want to split the data set intro training and test sub-samples. We do this by randomly selecting a desired fraction of the data to be part of the training sample and assign the rest to the test sample. For this example, we assign 10% of observations to the test sample and the remainder to the training data set. Once we have randomly allocated 90% of observations to the training data set, we use the function <code>setdiff1()</code> to find the complementary test sample. This function returns the set difference of two one-dimensional arrays, i.e., all array elements from the first argument that are *not* present in the second array.

```
[8]: import numpy as np
      from numpy.random import default_rng
     rng = default_rng(123)
     # sample size
     N = len(y)
      # fraction of data used for test sample
     test_size = 0.1
     # training and test sample sizes
     N_{test} = int(N * test_size)
     N_{train} = N - N_{test}
     # Randomly assign observations to training samples
     itrain = rng.choice(np.arange(N), size=N_train, replace=False)
     # Test sample is complement of training sample
     itest = np.setdiff1d(np.arange(N), itrain)
     # Select training sample
     X_train = X[itrain]
     y_train = y[itrain]
     # Select test sample
     X_test = X[itest]
     y_test = y[itest]
```

Once we have split the sample, we estimate the model on the training sample.

```
[9]: from sklearn.linear_model import LinearRegression

# Fit model on training sample
lr = LinearRegression()
lr.fit(X_train, y_train)
```

```
print('Regression coefficients')
print(f' Intercept: {lr.intercept_:.1f}')
print(f' Slope: {lr.coef_[0]:.1f}')
```

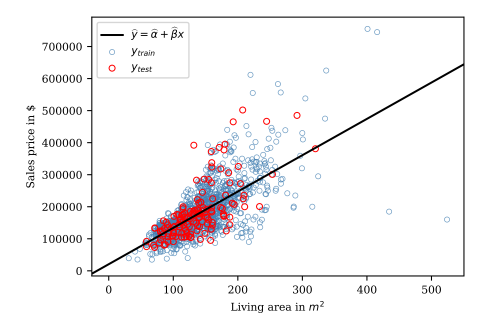
```
Regression coefficients
Intercept: 20675.0
Slope: 1133.9
```

The fitted coefficients show that for each additional square meter of living area, the sale price on average increases by \$1,134.

The following code creates a scatter plot showing the training and test samples and adds the fitted line.

```
[10]: intercept = lr.intercept_
       slope = lr.coef_[0]
       fig, ax = plt.subplots(1, 1, figsize=(5, 3.5))
       # Plot regression line
       ax.axline((0.0, intercept), slope=slope, lw=1.5, c='black',
           label=r'$\widehat{y} = \widehat{\alpha} + \widehat{\beta} x$')
       # Plot training data
       ax.scatter(X_train, y_train, s=15, color='none', edgecolor='steelblue',
           alpha=0.8, lw=0.5, label=r'$y_{train}$')
       # Plot test data
       ax.scatter(X_test, y_test, s=20, color='none', edgecolor='red',
           alpha=1.0, lw=0.75, label=r'$y_{test}')
       # Add annotations
       ax.set_xlabel('Living area in $m^2$')
       ax.set_ylabel(r'Sales price in $')
       ax.legend(loc='best')
```

[10]: <matplotlib.legend.Legend at 0x7fca8d47ef50>



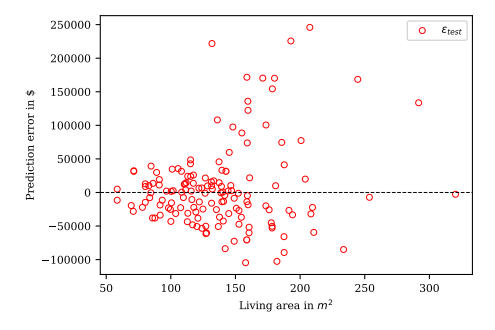
Since this is a univariate model, we can also plot the prediction error against the explanatory variable x. For this, we first need to compute the predicted values in the test sample and then the prediction error

for each observations,

$$\epsilon_i = y_i - \widehat{y}_i = y_i - \widehat{\alpha} - \widehat{\beta}x_i$$

for all *i* that are part of the test sample.

[11]: <matplotlib.lines.Line2D at 0x7fca8d227130>



As the graph shows, the errors are reasonably centred around 0 as we would expect from a model that contains an intercept. However, the error variance seems to be increasing in x which indicates that our model might be missing some explanatory variables.

Automatically creating training and test samples

A lot of code was required to create the training and test samples manually. Instead, we can use scikit-learn's train_test_split() to considerably simply this task. We need to either specify the test_size or train_size arguments to determine how the sample should be (randomly) split. The random_state argument is used to seed the RNG and get reproducible results.

The following code repeats the sample splitting we performed above using this simpler approach.

Regression coefficients Intercept: 21364.5 Slope: 1125.4

Note that the estimated coefficients need not be the same we got when splitting the sample manually. This is because scikit-learn uses a different way to randomly allocate the observations to training or test samples, and thus the composition of the training sample will differ.

10.1.3 Evaluating the model fit

One of the commonly used metrics to evaluate models is the *mean squared error* (MSE), defined as

$$MSE = \frac{1}{N} \sum_{i=1}^{N} (y_i - \widehat{y}_i)^2$$

which computes the average squared prediction error $y - \hat{y}$. The magnitude of the MSE is usually hard to interpret, so we often compute the *root mean squared error* (*RMSE*),

$$RMSE = \sqrt{MSE} = \left(\frac{1}{N} \sum_{i=1}^{N} (y_i - \hat{y}_i)^2\right)^{\frac{1}{2}}$$

which can be interpreted in units of the response variable y. Lastly, another measure of a model's fit is the *coefficients of determination* or R^2 , which is a normalized version of the MSE and usually takes on values between [0,1]. The R^2 is defined as

$$R^2 = 1 - \frac{MSE}{\widehat{Var}(y)}$$

where $\widehat{Var}(y)$ is the sample variance of the response y. Intuitively, an R^2 of 1 means that the model predicts the response for each observation perfectly (which is unlikely), whereas an R^2 of 0 implies that the model possesses no explanatory power relative to a model that includes only the sample mean. Note that in a test sample, the R^2 could even be negative.

While these measures are easy to implement ourselves, we can just as well use the functions provided in scikit-learn.metrics to do the work for us: $mean_squared_error()$ for the MSE and $r2_score()$ for the R^2 .

```
[13]: from sklearn.metrics import mean_squared_error, r2_score

# Compute predicted values for test sample
y_test_hat = lr.predict(X_test)
```

```
# Mean squared error (MSE)
mse = mean_squared_error(y_test, y_test_hat)

# Coefficient of determination (R²)
r2 = r2_score(y_test, y_test_hat)

print(f'Mean squared error: {mse:.1f}')
print(f'Root mean squared error {np.sqrt(mse):.1f}')
print(f'Coefficient of determination (R^2): {r2:.2f}')
```

```
Mean squared error: 2821058288.6
Root mean squared error 53113.6
Coefficient of determination (R^2): 0.58
```

Since we estimated a model with intercept, the $R^2 = 0.58$ implies that the model explains 58% of the variance in the test sample.

10.2 Multivariate linear regression

10.2.1 Data with several explanatory variables

Multivariate (or multiple) linear regression extends the simple model to multiple explanatory variables or regressors. To illustrate, we'll load the Ames housing data again but use several (continuous) explanatory variables.

```
[14]: # Uncomment this to use files in the local data/ directory

DATA_PATH = '../data'

# Load data directly from GitHub (for Google Colab)

# DATA_PATH = 'https://raw.githubusercontent.com/richardfoltyn/python-intro-PGR/main/data'
```

```
[15]: import pandas as pd

# Load data from CSV file
file = f'{DATA_PATH}/ames_houses.csv'
df = pd.read_csv(file)

# Print info about columns contained in DataFrame
df.info()
```

```
RangeIndex: 1460 entries, 0 to 1459
Data columns (total 13 columns):
                 Non-Null Count Dtype
# Column
                        _____
    SalePrice 1460 non-null float64
LotArea 1460 non-null float64
Neighborhood 1460 non-null object
BuildingType 1386 non-null object
OverallQuality 1460 non-null int64
0
    OverallCondition 1460 non-null int64
    YearBuilt 1460 non-null int64
                       1460 non-null object
    CentralAir
                    1460 non-null
 8 LivingArea
                                        float64
                      1460 non-null int64
 9 Bathrooms
 10 Bedrooms
                       1460 non-null int64
11 Fireplaces
                      1460 non-null
                                          int64
12 HasGarage
                       1460 non-null
                                          int64
dtypes: float64(3), int64(7), object(3)
memory usage: 148.4+ KB
```

<class 'pandas.core.frame.DataFrame'>

Compared to our earlier analysis, we'll now add the lot area (in m^2) as a feature to the model, which is thus given by

```
SalePrice_i = \alpha + \beta_0 LivingArea_i + \beta_1 LotArea_i + \epsilon_i
```

We first inspect the descriptive statistics of the newly added explanatory variable to get some idea about its distribution.

```
[16]: df[['SalePrice', 'LivingArea', 'LotArea']].describe()
                   SalePrice LivingArea
[16]:
                                                  LotArea
       count
                1460.000000 1460.000000 1460.000000
       mean 180921.195890 140.777444 976.949947
       std
             79442.502883 48.813961 927.199358

    34900.000000
    31.026587
    120.762165

    129975.000000
    104.923743
    701.674628

       min
       25%
       50%
              163000.000000 135.996777
                                              880.495526
              214000.000000
       75%
                               165.049367
                                             1077.709432
               755000.000000 524.107798 19994.963289
       max
```

As before, we convert the DataFrame columns to NumPy arrays, split the data into training and test samples and estimate a linear model.

```
[17]: features = ['LivingArea', 'LotArea']
  target = 'SalePrice'

y = df[target].to_numpy()
X = df[features].to_numpy()
```

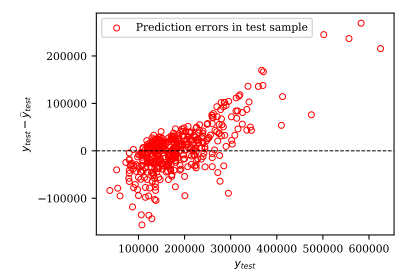
```
Intercept: 20025.027199480886
Coefficients: [1085.68252338 7.09932726]
```

The coefficient array $coef_now$ stores two values, the first one for LivingArea and the second one for LotArea. These coefficients are in the same order as the features in the feature matrix X passed when fitting the model.

Since we now have multiple explanatory variables, we can no longer easily plot prediction errors against one feature unless we fix the remaining features at some value or use 3D scatter plots. The latter of course does not help if we have more than two explanatory variables. Instead, we can plot the prediction errors against *y* which we do below.

```
plt.axhline(0.0, lw=0.75, ls='--', c='black')
plt.legend()
plt.xlabel(r'$y_{test}$')
plt.ylabel(r'$y_{test} - \widehat{y}_{test}$')
```

[19]: Text(0, 0.5, '\$y_{test} - \\widehat{y}_{test}\$')



The scatter plot indicates that the prediction errors differ systematically for different levels of y. The model on average overpredicts the sale price for low y (hence the error is negative) and underpredicts the price for large y (hence the error is positive).

10.2.2 Creating polynomial features

Instead of including additional explanatory variables, we can also create additional terms that are functions of variables. The most common way to do this is to include a polynomial in x (or polynomials in multiple explanatory variables). Thus a simple model could be turned into a model with several terms such as

$$y_i = \alpha + \beta_0 x_i + \beta_1 x_i^2 + \beta_2 x_i^3 + \epsilon_i$$

where y is modelled as a cubic polynomial in x. Note that this model is still called *linear* despite the fact that the mapping between x and y is obviously *non-linear*. However, what matters for estimation is that the model is linear in the model parameters $(\alpha, \beta_0, \beta_1, \beta_2)$. Linear models are thus quite flexible since they can include almost arbitrary non-linear transformations of explanatory and response variables.

To illustrate, we extend the previous model which included LivingArea and LotArea to now include a polynomial of degree 2 in both variables (often the terms "degree" and "order" are used interchangeably, so this might be called a 2nd-order polynomial). Specifically, if we have two variables x and z, such a polynomial would include all terms with exponents summing up to 2 or less:

$$p(x,z) = \beta_0 + \beta_1 x + \beta_2 z + \beta_3 x^2 + \beta_4 x \cdot z + \beta_5 z^2$$

We can use these six terms as explanatory variables in our linear models and estimate the parameters β_0, \dots, β_5 .

It would be quite error-prone to create such polynomials ourselves, so we are going to use scikit-learn's PolynomialFeatures class to accomplish this tasks. As the name implies, this transformation creates new features that are polynomials of a given input matrix. We can request that the resulting data should include an intercept ("bias") by specifying include_bias=True. Note that if an intercept is included in the feature matrix, we should fit the linear model without an intercept (by specifying fit_intercept=False) as otherwise the model would contain two constant terms.

```
from sklearn.preprocessing import PolynomialFeatures

# Create polynomials of degree 2 or less, including an intercept (bias)
poly = PolynomialFeatures(degree=2, include_bias=True)
Xpoly_train = poly.fit_transform(X_train)

# print polynomial exponents
poly.powers_
```

We can use the powers_ attribute to get a list of exponents for each generated feature. The above output tells us that the first feature was constructed as $\mathbf{X}_{(1)}^0 + \mathbf{X}_{(2)}^0$ since the first row of exponents is $[\mathfrak{o}, \mathfrak{o}]$ and is thus the intercept, while the second feature is given by $\mathbf{X}_{(1)}^1 + \mathbf{X}_{(2)}^0 = \mathbf{X}_{(1)}$, where the notation $\mathbf{X}_{(i)}$ refers to the *i*-th column of the input matrix \mathbf{X} .

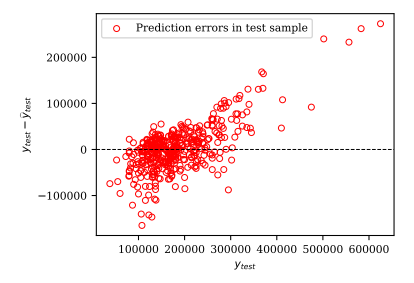
Now that we have transformed the input matrix **X**, we can estimate the linear model on the expanded feature matrix as before (adding fit_intercept=False):

```
Intercept: 0.0
Coefficients [-2.19724359e+04 1.24027918e+03 5.75108671e+01 3.95841742e-01 -2.46346339e-01 -3.77428502e-04]
```

The fitted model has 6 coefficients and the intercept is 0 as the LinearRegression model did not explicitly included one.

As earlier, we can plot the prediction errors as a function of the response variable. Before doing this, it is crucial to also transform the original explanatory variables in the test data set using the same polynomial transformation. We can achieve this by using the transform() method of the poly object we stored from earlier.

```
[22]: Text(0, 0.5, '$y_{test} - \\widehat{y}_{test}$')
```



10.2.3 Using scikit-learn pipelines

As you just saw, additional transformation steps before fitting the model can be tedious and errorprone (we might, for example, forget to transform the test data before computing predictions for the
test sample). For this reason, scikit-learn implements a feature called pipelines which allows us
to combine multiple transformations and a final estimation step. For this to work, all steps in the
pipeline except for the last must support fit() and transform() methods, and the final step in the
pipeline should be an estimator such as LinearRegression (for details, see the section on pipelines in
the scikit-learn user guide).

There are two ways to construct a pipeline:

- 1. Create an instance of the Pipeline class and specify the steps as name-value pairs.
- 2. Use the make pipeline() convenience function, which sets a default name for each step.

The first approach requires a list of tuples, where each tuple contains an (arbitrary) name and an object that implements the actions taken at this step. To compose a pipeline that creates polynomial features and fits a linear model to them, we would therefore proceed as follows:

In an interactive notebook, printing the pipe object will generate a visualisation which contains details for each step (some editors such as Visual Studio Code even make this visualisation interactive). To transform and fit the model in a single call, we simply need to invoke the fit() method. For example, using the training data from above, we run

```
[24]: # transform and fit in a single step
pipe.fit(X_train, y_train)
print(f'Coefficients: {pipe.named_steps.lr.coef_}')
```

```
Coefficients: [-2.19724359e+04 1.24027918e+03 5.75108671e+01 3.95841742e-01 -2.46346339e-01 -3.77428502e-04]
```

The second approach is to construct the pipeline using make_pipeline(). For this to work, we only need to pass the objects that constitute the individual steps of a pipeline as follows:

```
[25]: from sklearn.pipeline import make_pipeline

pipe = make_pipeline(
    PolynomialFeatures(degree=2),
    LinearRegression(fit_intercept=False)
)

# transform and fit model in single step
pipe.fit(X_train, y_train)

# print default names assigned to each step
pipe.named_steps
```

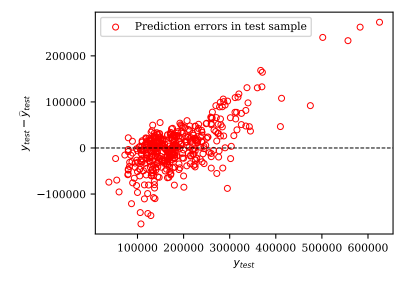
The function assigns default names to each step which are printed above (basically, these are just lowercase versions of the class defining a step). These names are occasionally required to retrieve information for a specific step. To demonstrate that the pipeline generates results that are equivalent to our manual implementation, we conclude this section by recreating the error plot from above.

```
[26]: # Predict test responses in a single step. No manual transformation required!
    y_test_hat = pipe.predict(X_test)

# Compute prediction errors
    errors = y_test - y_test_hat

plt.scatter(y_test, errors, s=20, lw=0.75,
        color='none', edgecolor='red',
        label='Prediction errors in test sample'
)
    plt.axhline(0.0, lw=0.75, ls='--', c='black')
    plt.legend()
    plt.xlabel(r'$y_{test}')
    plt.ylabel(r'$y_{test} - \widehat{y}_{test}')
```

[26]: Text(0, 0.5, '\$y_{test} - \widehat{y}_{test}\$')



10.3 Optimising hyperparameters with cross-validation

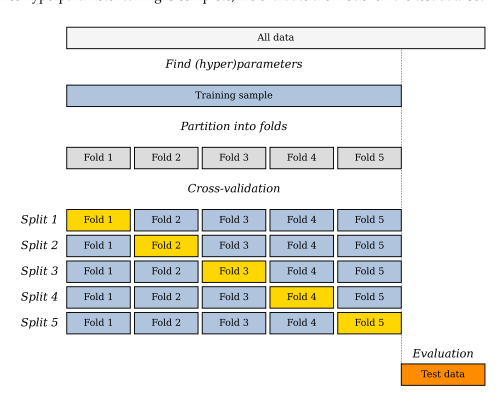
10.3.1 Outline of hyperparameter tuning

Previously, we discussed some ways to evaluate the model fit (MSE and R²) but did not specify what to to with this information. In this section, we demonstrate how we can use these measures to tune parameters which govern the estimation process, such as the polynomial degree from the previous section. Such parameters are usually called *hyperparameters* to clearly distinguish them from the parameters that are estimated by the model (e.g., the coefficients of a linear model).

Tuning hyperparameters, estimating the model and evaluating its generalisability cannot be done based on the same data set, as these steps then become interdependent (for the same reason we don't want to evaluate the model fit on the training sample as this would lead to overfitting). To this end, in machine learning we often split the data intro *three* parts: the training, validation and test data sets. Models are estimated on the training sample, the choice of hyperparameters is determined based on the validation sample and model generalisability is determined based on the test sample which was not used in estimation or tuning at all.

However, because we might not have enough data to split the sample this way, we usually perform so-called *cross-validation* which is illustrated in the figure below:

- 1. We split the overall data set into training and test sub-samples.
- 2. The training data set is further split into *K* so-called *folds*.
 - 1. For each k = 1, ..., K, a smaller training sample is formed by excluding the k-th fold and estimating the model on the remaining K 1 folds. We then compute the chosen metric of model fit on the k-th fold and store the result.
 - 2. After cycling through all *K* folds, we have *K* values of our desired metric, which we then average to get our final measure. Hyperparameters are tuned by minimising (or maximising) this averaged metric.
- 3. Once hyperparameter tuning is complete, we evaluate the model on the test data set.



In this unit we will skip the final step of assessing generalisability on the test data set. Consequently, when running cross-validation, we will only use training/validation data sets and we will use the terms "validation" and "test" interchangeably.

The scikit-learn documentation contains a wealth of additional information on cross-validation. Another method to perform hyperparameter tuning is grid search which we won't cover in this unit.

10.3.2 Example: Tuning of the polynomial degree

To illustrate the concept, we demonstrate the procedure outlined above by tuning the polynomial degree for the example covered in the previous section by minimising the root mean squared error (RMSE).

We recreate the data set in the same way as before, using LivingArea and LotArea as explanatory variables.

```
[27]: import pandas as pd

df = pd.read_csv(f'{DATA_PATH}/ames_houses.csv')

features = ['LivingArea', 'LotArea']
  target = 'SalePrice'

y = df[target].to_numpy()
X = df[features].to_numpy()
```

We now iterate over the candidate polynomial degrees $d=0,\ldots,4$ and apply k-fold cross-validation with 10 folds. There is no need to manually split the sample into training and validation/test sub-sets. Instead, we use the KFold class to automatically create the splits for us. Once we have created an instance with the desired number of folds, e.g., KFold(n_splits=10), we can call the split() method which iterates trough all possible combinations of training and test data sets and returns the array indices for each.

```
[28]: from sklearn.pipeline import Pipeline
       from sklearn.preprocessing import PolynomialFeatures
       from sklearn.linear_model import LinearRegression
       from sklearn.model_selection import KFold
       from sklearn.metrics import mean_squared_error
       degrees = np.arange(5)
       rmse_mean = []
       for d in degrees:
           # Create pipeline to transform and fit the model. Pipeline depends
           # on polynomial degree!
           pipe = Pipeline(steps=[
               ('poly', PolynomialFeatures(degree=d, include_bias=True)),
               ('lr', LinearRegression(fit_intercept=False))
           1)
           rmse_fold = []
           # Create 10 folds
           folds = KFold(n_splits=10)
           # Iterate through all combinations of training/test data
           for itrain, itest in folds.split(X, y):
               # Extract training data for this split
               X train = X[itrain]
               y_train = y[itrain]
               # Extract test (or validation) data for this split
               X_test = X[itest]
```

```
y_test = y[itest]
        # Fit model on training data for given degree
        pipe.fit(X_train, y_train)
        # Predict response on test data
        y_test_hat = pipe.predict(X_test)
        # Compute RMSE as model fit measure: function returns RMSE
        # if squared=False is passed!
        rmse = mean_squared_error(y_test, y_test_hat, squared=False)
        # Store RMSE for current split
        rmse_fold.append(rmse)
    # Store average MSE for current polynomial degree
    rmse_mean.append(np.mean(rmse_fold))
# Convert to NumPy array
rmse_mean = np.array(rmse_mean)
# Print average RMSE for all polynomial degrees
rmse_mean
```

```
[28]: array([ 79020.25214052, 55268.49840787, 55228.58262438, 55885.05422781, 437077.00283378])
```

This code returns an array of 5 averaged RMSEs, one for each d = 0, ..., 4. We can now find the optimal d by picking the one which has the lowest mean squared error in the test samples using the $\underset{\text{argmin}()}{\operatorname{argmin}()}$ function which returns the $\underset{\text{index}}{\operatorname{index}}$ of the smallest array element.

```
[29]: # Find index of polynomial degree with smallest RMSE
imin = np.argmin(rmse_mean)

# RMSE-minimising degree
dmin = degrees[imin]

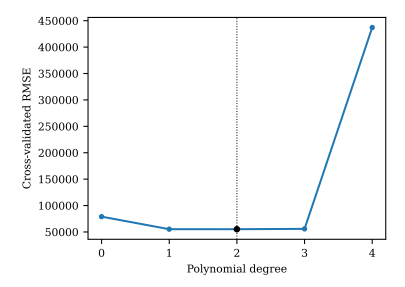
print(f'Polynomial degree with minimum RMSE: {dmin}')
```

Polynomial degree with minimum RMSE: 2

Finally, it is often instructive to visualise how the RMSE evolves as a function of the hyperparameter we want to tune.

```
plt.plot(degrees, rmse_mean, marker='o', ms=3)
plt.xlabel('Polynomial degree')
plt.ylabel('Cross-validated RMSE')
plt.scatter(degrees[imin], rmse_mean[imin], s=15, c='black', zorder=100)
plt.xticks(degrees)
plt.axvline(imin, ls=':', lw=0.75, c='black')
```

[30]: <matplotlib.lines.Line2D at 0x7fca84770070>



Here we see that for d = 0 (the intercept-only model), the model underfits the data leading to a high prediction error. However, higher d's do not always translate into a better fit. For d = 4, the model vastly overfits the data, resulting in poor performance in the test sample and a very large RMSE.

Automating cross-validation

The code implemented above to run cross-validation was needlessly complex even though we leveraged the KFold class to do the sample splitting for us. Fortunately, scikit-learn provides us with even more convenience functions that further simplify this process. For example, if we want to perform tuning based on a single score (such as the root mean squared error), we can instead use cross_val_score(). This function requires us to specify an estimator (or a pipeline), the number of folds to use for cross-validation (cv=10) and the metric to evaluate. For example, if we want to compute the RMSE, we would pass the argument scoring='neg_root_mean_squared_error'. Note that the function returns the negative RMSE which we need to correct manually.

See the documentation for a complete list of valid metrics that can be passed as scoring arguments. Alternatively, scikit-learn lists available metrics by running

```
import sklearn.metrics
sklearn.metrics.get_scorer_names()
```

The following code re-implements the k-fold cross-validation from earlier using cross val score():

```
# Function returns NEGATIVE RMSE, correct this here!
rmse_mean.append(np.mean(-score))

# Convert to NumPy array
rmse_mean = np.array(rmse_mean)

# Print average RMSE for all polynomial degrees
rmse_mean
```

```
[31]: array([ 79020.25214052, 55268.49840787, 55228.58262438, 55885.05422781, 437077.00283378])
```

The RMSE-minimising degree is of course the same as before:

```
[32]: imin = np.argmin(rmse_mean)
dmin = degrees[imin]

print(f'Polynomial degree with min. RMSE: {dmin}')
```

Polynomial degree with min. RMSE: 2

10.4 Linear models with regularisation: Ridge regression

The linear models we encountered so far are part of the standard econometrics toolbox. In this section, we look at extensions that make these models more useful for machine learning applications.

The first model we study is Ridge regression, which estimates a linear model but adds a penalty term to the loss function which is now given by

$$L(\mu, \boldsymbol{\beta}) = \underbrace{\sum_{i=1}^{N} (y_i - \mu - \mathbf{x}_i' \boldsymbol{\beta})^2}_{\text{Sum of squared errors}} + \underbrace{\alpha \sum_{k=1}^{K} \beta_k^2}_{\text{L2 penalty}}$$

Compared to ordinary least squares (OLS) we discussed initially, the penalty term increases the loss function if the estimated coefficients β are large. This term is called an L2 penalty because it corresponds to the (squared) L2 vector norm. In many textbooks, you will see the penalty term written as $\alpha \|\beta\|_2^2$ which is equivalent to the formula used above.

For any $\alpha > 0$, the loss function is increasing in the (absolute) coefficient values, thus the minimum L might be one where the elements of β are smaller than what they would have been with OLS. We therefore say that that Ridge regression applies *regularisation* or *shrinkage*. Note, however, that the intercept which we now denote by μ is not included in the penalty term, and thus no regularisation is applied to it.

Clearly, the regularisation strength depends on the value of α . For tiny (or zero) α , the estimated $\widehat{\beta}$ will be close (or identical) to OLS, while for large α the estimated coefficients will be compressed towards zero. In this setting, α is a hyperparameter and we can accordingly use cross-validation to find an "optimal" value.

Why would we ever want to use Ridge regression given that OLS is the best linear unbiased estimator? It turns out that regularisation can help in scenarios where we have a large number of (potentially multicollinear) regressors in which case OLS is prone to overfitting.

10.4.1 Example: Polynomial approximation

We illustrate such problems and the benefits of Ridge regression using a highly stylised example. Imagine we want to approximate a non-linear function using a high-order polynomial, a setting which is notoriously susceptible to overfitting (also see the optional exercises for more illustrations). Assume that our model is given by

$$y_i = \cos\left(\frac{3}{2}\pi x_i\right) + \epsilon_i$$
$$\epsilon \stackrel{\text{iid}}{\sim} N(0, 0.25)$$

where ϵ_i as an additive, normally-distributed measurement error term with mean 0 and variance $\frac{1}{4}$.

The true values of y (without measurement error) are computed using the function compute_true_y() which returns y for a given x.

```
[33]: import numpy as np

# True function (w/o errors)
def compute_true_y(x):
    return np.cos(1.5 * np.pi * x)
```

The following code creates a demo sample with N=100 observations.

```
[34]: from numpy.random import default_rng

# Initialise random number generator
rng = default_rng(1234)

# Sample size
N = 100

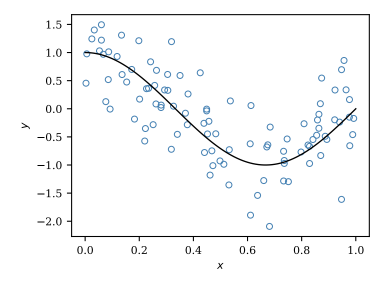
# Randomly draw explanatory variable x uniformly distributed on [0, 1]
x = rng.random(size=N)

# Draw errors from normal distribution
epsilon = rng.normal(scale=0.5, size=N)

# Compute y, add measurement error
y = compute_true_y(x) + epsilon
```

The next graph visualises the true relationship and the sample points (x_i, y_i) .

```
[35]: Text(o, o.5, '$y$')
```



Estimating the Ridge and linear regression models

We now estimate a polynomial approximation where we assume that y is a degree-K polynomial in x, i.e.,

$$y_i \approx \mu + \beta_1 x_i + \beta_2 x_i^2 + \dots + \beta_K x_i^K$$

For this example, we choose an unconventionally high K = 15 since we anticipate that this leads to problems with OLS.

A few more steps are required before we can run the Ridge regression:

- We need to create the polynomial in *x* which we do with the PolynomialFeatures transformation we already encountered.
- Moreover, regularisation methods can be susceptible to scaling issues, so we need to demean and normalise all input variables, i.e., we make sure that each feature has mean 0 and a variance of 1. We can automate this step using the StandardScaler transformation.
- Finally, the estimation step is performed by the Ridge class.
- We build a Pipeline that combines these transformation together with the estimation step. We use the function make_pipeline() to simplify this step.

To run the Ridge regression, we moreover need to specify the regularisation strength α which we set to $\alpha = 3$ for illustration.

```
[36]: from sklearn.linear_model import Ridge
       from sklearn.preprocessing import PolynomialFeatures, StandardScaler
       from sklearn.pipeline import make_pipeline
       # Polynomial degree
       degree = 15
       # Build pipeline of transformations and Ridge estimation.
       # We create the polynomial transformation w/o the intercept so we
       # need to include an intercept in the fitting step.
       pipe_ridge = make_pipeline(
           PolynomialFeatures(
               degree=degree,
               include_bias=False
           StandardScaler(),
                                                        # standardise features
           Ridge(alpha=3.0, fit_intercept=True)
                                                        # Fit Ridge regression
       )
```

```
# Make sure X is a matrix
X = x[:, None]

# Fit model
pipe_ridge.fit(X, y)
```

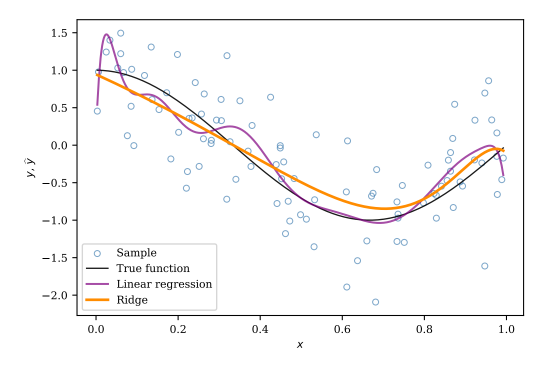
It is instructive to estimate the same model using linear regression and compare the results:

To illustrate the difference for this artificial example, we add the model predictions from the Ridge and linear regressions to the sample scatter plot we created earlier.

```
[38]: import matplotlib.pyplot as plt
       fig, ax = plt.subplots(1, 1, figsize=(6, 4))
       # Grid on which to evaluate true model and predictions
       xvalues = np.linspace(np.amin(x), np.amax(x), 500)
       # Sample points scatter plot
       ax.scatter(x, y, s=20, color='none', edgecolor='steelblue',
           lw=0.75, alpha=0.7, label='Sample')
       ax.plot(xvalues, compute_true_y(xvalues), color='black', lw=1.0, alpha=0.9,
           label='True function')
       # LR prediction
       ax.plot(xvalues, pipe_lr.predict(xvalues[:, None]), color='purple',
           alpha=0.7, label='Linear regression')
       # Ridge prediction
       ax.plot(xvalues, pipe_ridge.predict(xvalues[:, None]),
           color='darkorange', lw=2.0,
           label='Ridge'
       )
```

```
ax.legend()
ax.set_xlabel('$x$')
ax.set_ylabel('$y$, $\widehat{y}$')
```

[38]: $Text(0, 0.5, '$y$, $\\\widehat{y}$')$



As the graph shows, the OLS model vastly overfits the data which is what we would expect in this settings. Conversely, the prediction of the Ridge regression is reasonably close to the true function and much better behaved despite the high polynomial degree used here.

To gain some intuition for what is going on, it is instructive to plot the estimated Ridge coefficients for a whole range of α values which we do in the code below. Note that we choose the grid of α to be uniformly spaced in logs since we want to zoom in on what happens when α is small. This is accomplished by using np.logspace() instead of np.linspace().

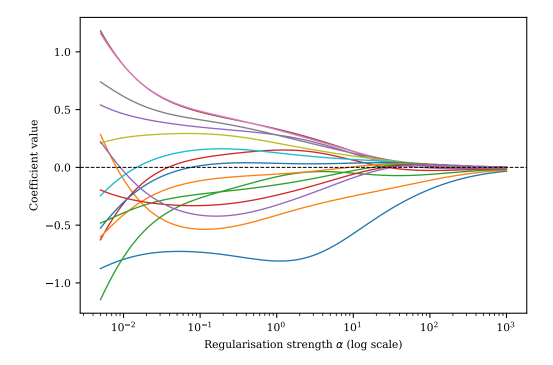
```
[39]: # Create grid of alphas spaced uniformly in logs
       alphas = np.logspace(start=np.log10(5.0e-3), stop=np.log10(1000.0), num=100)
       # Re-create pipeline w/o Ridge estimator, estimation step differs for each alpha
       transform = make_pipeline(
           PolynomialFeatures(
               degree=degree,
               include_bias=False
           ),
           StandardScaler()
       )
       # Create polynomial features
       Xtrans = transform.fit_transform(x[:, None])
       # Array to store coefficients for all alphas
       coefs = np.empty((len(alphas), Xtrans.shape[1]))
       # Estimate Ridge for each alpha, store fitted coefficients
       for i, alpha in enumerate(alphas):
           ridge = Ridge(alpha=alpha, fit_intercept=True)
           # Fit model for given alpha
```

```
ridge.fit(Xtrans, y)
coefs[i] = ridge.coef_
```

The following plot shows each coefficient (one color corresponds to one coefficients) for different values of α on the *x*-axis. Note that the *x*-axis is plotted on a \log_{10} scale which allows us to zoom in on smaller values of α .

```
plt.figure(figsize=(6,4))
  plt.plot(alphas, coefs, lw=1.0)
  plt.xscale('log', base=10)
  plt.axhline(0.0, ls='--', lw=0.75, c='black')
  plt.xlabel(r'Regularisation strength $\alpha$ (log scale)')
  plt.ylabel('Coefficient value')
```

[40]: Text(0, 0.5, 'Coefficient value')



For small α (on the left) the estimated coefficients are close to the (standardised) OLS coefficients, but their values shrink towards zero as α increases. For a very large $\alpha = 10^3$ the estimated coefficients are basically zero since the penalty dominates the sum of squared errors in the loss function.

10.4.2 Tuning the regularisation parameter via cross-validation

In the previous example, we picked an arbitrary regularisation strength α when fitting the Ridge regression. In applied work, we would want to tune α (which is a hyperparameter) using cross-validation instead. To this end, we could use the generic cross-validation functionality we studied earlier in this unit since that one works for all types of estimators. However, scikit-learn implements a cross-validation class specifically for Ridge regression called RidgeCV which we use in the code below to find an optimal α .

```
[41]: from sklearn.linear_model import RidgeCV

# RidgeCV does not support pipelines, so we need to transform x before
# cross-validation.

transform = make_pipeline(
    PolynomialFeatures(
```

```
degree=degree,
    include_bias=False
),
StandardScaler()
)

# Create standardised polynomial features
Xtrans = transform.fit_transform(x[:, None])

# Set of candidate alphas on [1.0e-5, 5] to cross-validate.
# Spaced uniformly in logs to get denser grid for small alphas.
alphas = np.logspace(start=np.log1o(1.0e-5), stop=np.log1o(5), num=100)

# Create and run Ridge cross-validation
rcv = RidgeCV(alphas=alphas, store_cv_values=True).fit(Xtrans, y)
```

By default, RidgeCV uses the (negative) mean squared error (MSE) to find the best α which we can then recover from the alpha_attribute.

```
[42]: # Recover best alpha that minimizes MSE
alpha_best = rcv.alpha_

# Best MSE is stored as negative score!
MSE_best = - rcv.best_score_

print(f'Best alpha: {alpha_best:.3g} (MSE: {MSE_best:.3g})')
```

Best alpha: 0.6 (MSE: 0.267)

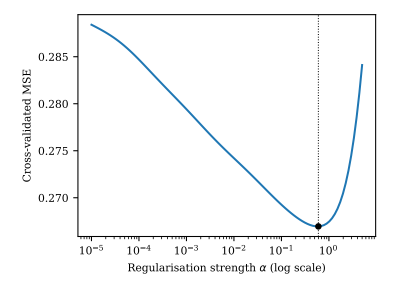
Because we fitted RidgeCV with the argument store_cv_values=True, the fitted object stores the MSE for each CV split in the attribute cv_values_ which we can use to plot the MSE as a function of α .

```
[43]: import matplotlib.pyplot as plt

# Compute average MSE for each alpha value across all folds
mse_mean = np.mean(rcv.cv_values_, axis=0)

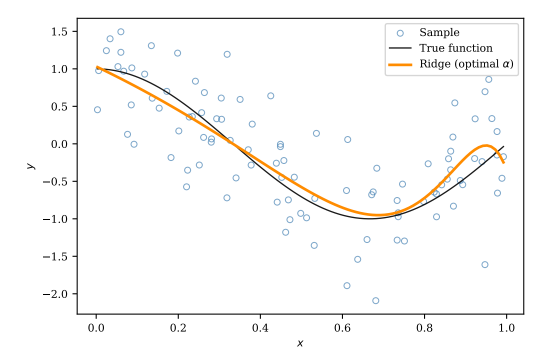
# Index of MSE-minimising alpha
imin = np.argmin(mse_mean)

# Plot MSE against alphas, highlight minimum MSE
plt.plot(alphas, mse_mean)
plt.xlabel(r'Regularisation strength $\alpha$ (log scale)')
plt.ylabel('Cross-validated MSE')
plt.scatter(alphas[imin], mse_mean[imin], s=15, c='black', zorder=100)
plt.axvline(alphas[imin], ls=':', lw=0.75, c='black')
plt.xscale('log')
```



Now that we have identified the optimal α , we can re-fit the Ridge regression and plot the prediction from this model. Note that this is not strictly necessary as the return value of RidgeCV can be used to do prediction based on coefficients estimated for the best α , but because RidgeCV does not support pipelines, we'd have to apply any transformations manually before doing so.

[45]: Text(0, 0.5, '\$y\$')



10.5 Linear models with regularisation: Lasso

Another widely used estimator with shrinkage is LASSO (least absolute shrinkage and selection operator) which adds an L1 penalty term to the objective function:

$$L(\mu, \boldsymbol{\beta}) = \underbrace{\sum_{i=1}^{N} (y_i - \mu - \mathbf{x}_i' \boldsymbol{\beta})^2}_{\text{Sum of squared errors}} + \underbrace{\alpha \sum_{k=1}^{K} |\beta_k|}_{\text{L1 penalty}}$$

As with the Ridge regression, this additional term penalises large coefficient values. This term is called an L1 penalty because it corresponds to the L1 vector norm which can equivalently be written as $\alpha \|\boldsymbol{\beta}\|_1$.

While the objective looks very similar to Ridge regression, using the L1 instead of the L2 norm can produce much more parsimonious models because many coefficients end up being exactly zero and the corresponding features are thus eliminated from the model. We will see this in the example below.

10.5.1 Example: Polynomial approximation

We apply Lasso to the same random sample as in the section on Ridge which allows us to compare the two methods. The following code recreates that data, making the same functional form and distributional assumptions as in the previous section.

```
[46]: import numpy as np

# True function (w/o errors)
def compute_true_y(x):
    return np.cos(1.5 * np.pi * x)
[47]: from numpy random import default rng
```

```
[47]: from numpy.random import default_rng

# Initialise random number generator
```

```
rng = default_rng(1234)

# Sample size
N = 100

# Randomly draw explanatory variable x uniformly distributed on [0, 1]
x = rng.random(size=N)

# Draw errors from normal distribution
epsilon = rng.normal(scale=0.5, size=N)

# Compute y, add measurement error
y = compute_true_y(x) + epsilon
```

Estimating the Lasso and linear regression models

As with Ridge, we need to standardise the explanatory variables before fitting Lasso. The code below repeats these steps, but we now use Lasso to perform the model estimation. For now, we set the regularisation strength to $\alpha=0.015$ and will use cross-validation to find the optimal value later.

Note that for Lasso it might be necessary to increase the number of iterations by increasing the max_iter parameter (from the default of 1,000).

```
[48]: from sklearn.linear_model import Lasso
      from sklearn.preprocessing import PolynomialFeatures, StandardScaler
      from sklearn.pipeline import make_pipeline
       # Polynomial degree
      degree = 15
       # Build pipeline of transformations and Lasso estimation.
       # We create the polynomial transformation w/o the intercept so we
       # need to include an intercept in the fitting step.
      pipe_lasso = make_pipeline(
           PolynomialFeatures(
              degree=degree,
               include_bias=False
           ),
           StandardScaler(),
           Lasso(alpha=0.015, fit_intercept=True, max_iter=10000)
      # Make sure X is a matrix
      X = x[:, None]
      pipe_lasso.fit(X, y)
```

For completeness, let's also recreate the linear regression estimation and plot the model predictions alongside the Lasso.

```
[49]: from sklearn.linear_model import LinearRegression

# Create pipeline for linear model (linear regression does not require
    # standardisation!)
pipe_lr = make_pipeline(
    PolynomialFeatures(
```

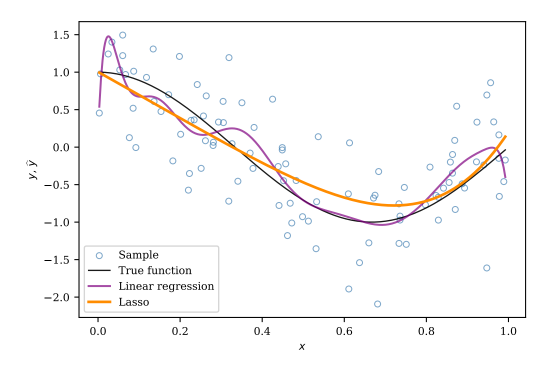
```
degree=degree,
    include_bias=False
),
LinearRegression(fit_intercept=True)
)

# Make sure X is a matrix
X = x[:, None]
pipe_lr.fit(X, y)
```

The following code plots the sample data, the true functional relationship, and the predictions from the linear regression and Lasso models.

```
[50]: import matplotlib.pyplot as plt
       fig, ax = plt.subplots(1, 1, figsize=(6, 4))
       # Sample points scatter plot
       ax.scatter(x, y, s=20, color='none', edgecolor='steelblue',
          lw=0.75, alpha=0.7, label='Sample')
       # Grid on which to evaluate true model and predictions
       xvalues = np.linspace(np.amin(x), np.amax(x), 500)
       # True model
       ax.plot(xvalues, compute_true_y(xvalues),
           color='black', lw=1.0, alpha=0.9,
           label='True function'
       # Linear regression prediction
       ax.plot(xvalues, pipe_lr.predict(xvalues[:, None]),
           color='purple', alpha=0.7,
           label='Linear regression'
       # Lasso prediction
       ax.plot(xvalues, pipe_lasso.predict(xvalues[:, None]),
           color='darkorange', lw=2.0,
           label='Lasso'
       )
       ax.legend()
       ax.set_xlabel('$x$')
       ax.set_ylabel('$y$, $\widehat{y}$')
```

[50]: Text(0, 0.5, '\$y\$, \$\\widehat{y}\$')



You might be wondering why we chose $\alpha = 0.015$ whereas we initially used $\alpha = 3$ for the Ridge regression. The reason is that the scikit-learn implementation of Lasso uses a slightly different loss function than the one given above, namely

$$L(\mu, \boldsymbol{\beta}) = \frac{1}{2N} \sum_{i=1}^{N} (y_i - \mu - \mathbf{x}_i' \boldsymbol{\beta})^2 + \alpha \|\beta_k\|_1$$

which scales the sum of squared errors term by a factor of of $(2N)^{-1}$. This makes no difference for the optimisation, but changes the interpretation of the regularisation strength α compared to Ridge regression. For our sample size of N=100, an α_{Ridge} used for the Ridge estimator approximately corresponds to $\alpha_{Lasso}=\frac{\alpha}{200}$ when plugged into Lasso.

We next fit the model for different values of α on the interval $[5 \times 10^{-3}, 1]$ where we again space the α uniformly in logs.

```
[51]: # Create grid of alphas spaced uniformly in logs
       alphas = np.logspace(start=np.log10(5.0e-3), stop=np.log10(1.0), num=100)
       # Re-create pipeline w/o Lasso estimator, estimation step differs for each alpha
       transform = make_pipeline(
           PolynomialFeatures(
               degree=degree,
               include_bias=False
           StandardScaler()
       )
       # Create polynomial features
       Xtrans = transform.fit_transform(x[:, None])
       # Array to store coefficients for all alphas
       coefs = np.empty((len(alphas), Xtrans.shape[1]))
       # Estimate Lasso for each alpha, store fitted coefficients
       for i, alpha in enumerate(alphas):
           lasso = Lasso(alpha=alpha,
               fit_intercept=True,
               max iter=10000
```

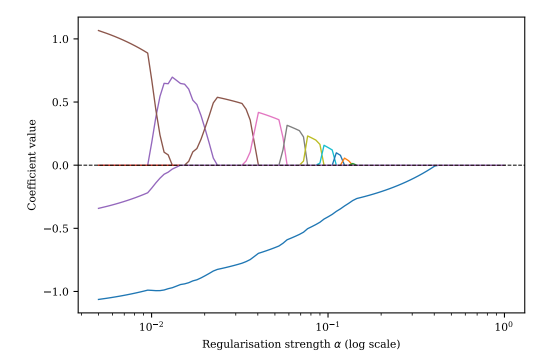
```
# Fit model for given alpha
lasso.fit(Xtrans, y)

coefs[i] = lasso.coef_
```

Plotting the fitted coefficients against α on a log scale looks as follows:

```
[52]: plt.figure(figsize=(6, 4))
   plt.plot(alphas, coefs, lw=1.0)
   plt.xscale('log', base=10)
   plt.axhline(0.0, ls='--', lw=0.75, c='black')
   plt.xlabel(r'Regularisation strength $\alpha$ (log scale)')
   plt.ylabel('Coefficient value')
```

[52]: Text(0, 0.5, 'Coefficient value')

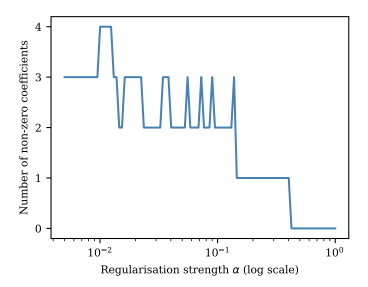


The graph shows that the coefficient estimates are quite different than what we obtained from the Ridge estimator. In fact, most of them are exactly zero for most values of α . We highlight this result in the graph below which plots the number of non-zero coefficients against α .

```
[53]: # Number of non-zero coefficients for each alpha.
nonzero = np.sum(np.abs(coefs) > 1.0e-6, axis=1).astype(int)

plt.plot(alphas, nonzero, lw=1.5, c='steelblue')
plt.xscale('log', base=10)
plt.yticks(np.arange(0, np.amax(nonzero) + 1))
plt.xlabel(r'Regularisation strength $\alpha$ (log scale)')
plt.ylabel('Number of non-zero coefficients')
```

[53]: Text(0, 0.5, 'Number of non-zero coefficients')



Clearly, in this case the model estimated by Lasso is substantially less complex than the linear regression or even Ridge regression. For most values of α , only 2-3 features out of the original 15 are retained in the model!

10.5.2 Tuning the regularisation parameter via cross-validation

In the previous example, we picked an arbitrary regularisation strength α when fitting the Lasso. In this section, we again find an optimal α using cross-validation. Just like in the case of Ridge regression, scikit-learn implements a cross-validation class specifically for Lasso called LassoCV which we use in the code below to find an optimal α .

LassoCV optionally accepts a grid of candidate α just like RidgeCV, but the default way to run cross-validation is to specify the fraction $\epsilon = \frac{\alpha_{min}}{\alpha_{max}}$ (default: 10^{-3}) and the grid size (default: 100). LassoCV then automatically determines an appropriate grid which is stored in the alphas_ attribute once fitting is complete. Moreover, we use the cv=... argument to set the desired number of CV folds (default: 5).

After fitting, we can recover the best alpha from the alpha_attribute and the MSE for each α on the grid and each CV fold from the mse_path_ attribute.

```
[55]: # Recover best alpha that minimizes MSE
alpha_best = lcv.alpha_

# MSE for each alpha, averaged over folds
```

```
mse_mean = np.mean(lcv.mse_path_, axis=1)

# Index of min. MSE
imin = np.argmin(mse_mean)

mse_best = mse_mean[imin]

print(f'Best alpha: {alpha_best:.4g} (MSE: {mse_best:.3g})')
```

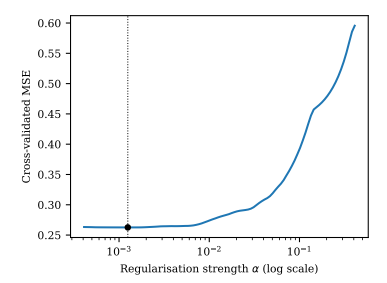
Best alpha: 0.001251 (MSE: 0.263)

The next plot visualises the average MSE over the entire range of candidate α values.

```
[56]: import matplotlib.pyplot as plt

# Recover grid of alphas used for CV
alphas = lcv.alphas_

# Plot MSE against alphas, highlight minimum MSE
plt.plot(alphas, mse_mean)
plt.xlabel(r'Regularisation strength $\alpha$ (log scale)')
plt.ylabel('Cross-validated MSE')
plt.scatter(alphas[imin], mse_mean[imin], s=15, c='black', zorder=100)
plt.axvline(alphas[imin], ls=':', lw=0.75, c='black')
plt.xscale('log')
```

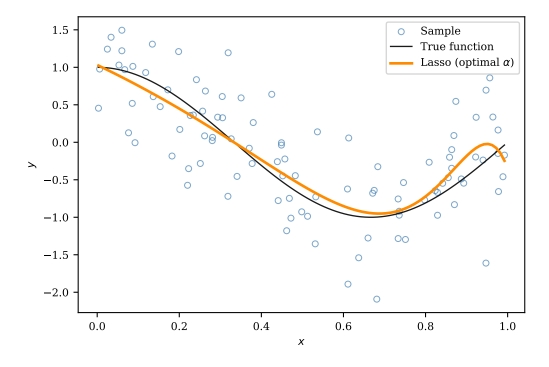


Now that we have identified the optimal α , we can re-fit the Lasso and plot the prediction from this model.

```
[57]: # Create pipeline with Lasso using optimal alpha
pipe_lasso = make_pipeline(
    PolynomialFeatures(
         degree=degree,
         include_bias=False
    ),
    StandardScaler(),
    Lasso(alpha=alpha_best, fit_intercept=True, max_iter=100000)
)
pipe_lasso.fit(x[:, None], y)
```

Finally, we visually compare the true model to the Lasso prediction using the optimal value of α .

[58]: Text(0, 0.5, '\$y\$')



10.6 Dealing with categorical data (optional)

So far in this unit, we only dealt with continuous data, i.e., data that can take on (almost any) real value. However, many data sets contain categorical variables which take on a finite number of admissible values or even binary variables (often called dummy or indicator variables) which can be either 0 or 1.

Dealing with such data in scikit-learn is less straightforward than in most other software packages, so in this section we illustrate how to work with categorical variables.

To do this, we again load the Ames house data which contains several categorical and binary variables.

```
[59]: import pandas as pd

df = pd.read_csv(f'{DATA_PATH}/ames_houses.csv')
```

For example, consider the BuildingType variable, which (in this simplified version of the data) takes on three values, 'Single-family', 'Townhouse' and 'Two-family' and has on top a few missing observations. We use the value_counts() method to tabulate the number of observations falling into each category. The argument dropna=False also includes the number of missing values in the tabulation.

```
[60]: df['BuildingType'].value_counts(dropna=False).sort_index()

[60]: Single-family 1220
    Townhouse 114
    Two-family 52
    NaN 74
    Name: BuildingType, dtype: int64
```

There are several ways to deal with such data (see the official documentation for details). First, since the data are stored as strings, we could use OrdinalEncoder to map these strings to integers.

```
from sklearn.preprocessing import OrdinalEncoder

# Drop rows with NA
df = df.dropna(subset='BuildingType')
enc = OrdinalEncoder()

# Transform string variable into integers
bldg_int = enc.fit_transform(df[['BuildingType']].to_numpy())

# Print unique values and histogram
print(f'Unique values: {np.unique(bldg_int)}')
print(f'Histogram: {np.bincount(bldg_int.astype(int).flatten())}')
```

```
Unique values: [0. 1. 2.]
Histogram: [1220 114 52]
```

This, however, is still not particularly useful if we want to use these categories as explanatory variables in a LinearRegression because scikit-learn will simply treat them as a continuous variable that happens to take on the values 0, 1 or 2. There is nothing to enforce the categorical nature of this data.

An alternative encoding strategy is to create a binary dummy variable for each possible value of a categorical variable. This is achieved using the OneHotEncoder transformation as illustrated by the following code:

```
[62]: from sklearn.preprocessing import OneHotEncoder

# Drop rows with NA
df = df.dropna(subset='BuildingType')

# List of unique categories
bldg_uniq = list(np.sort(df['BuildingType'].unique()))

# Create dummy variable encoder
enc = OneHotEncoder(categories=[bldg_uniq], sparse_output=False)

# Convert string variable into binary indicator variables
```

```
bldg_dummies = enc.fit_transform(df[['BuildingType']].to_numpy())
```

When creating a OneHotEncoder, we can optionally pass the list of possible values using the categories=... argument. By default, this transformation returns a sparse matrix since each row will have exactly one element that is 1 while the remaining elements are 0. Using a sparse matrix saves memory but makes interacting with the resulting array more cumbersome. For small data sets, there is no need to use sparse matrices.

The transformed data now looks as follows:

In this particular example, the first five observations fall into the first category, hence the first column contains ones while the remaining elements are zero.

We can sum the number of ones in each column to verify that the frequency of each category remains the same:

```
[64]: bldg_dummies.sum(axis=0)

[64]: array([1220., 114., 52.])
```

The category labels taken from the original data are stored in the categories_ attribute:

```
[65]: enc.categories_[0]

[65]: array(['Single-family', 'Townhouse', 'Two-family'], dtype=object)
```

Now that we have converted the categories into a dummy matrix, we can append it to the continuous explanatory variables and fit the linear model as usual:

```
[66]: from sklearn.linear_model import LinearRegression
       import pandas as pd
       # Name of dependent variable
       target = 'SalePrice'
       # Continuous explanatory variables to include
       continuous = ['LivingArea', 'LotArea']
       y = df[target].to_numpy()
       # Feature matrix with dummies appended
       X = np.hstack((df[continuous].to_numpy(), bldg_dummies))
       # Create and fit linear model
       lr = LinearRegression(fit intercept=False)
       lr.fit(X, y)
       # Create DataFrame containing estimated coefficients
       labels = continuous + [f'BuildingType: {s}' for s in enc.categories_[0]]
       coefs = pd.DataFrame(lr.coef_, index=labels, columns=['Coef'])
       coefs
```

```
[66]: Coef
LivingArea 1152.605777
```

```
LotArea 8.385689
BuildingType: Single-family 12173.902329
BuildingType: Townhouse 37735.490189
BuildingType: Two-family -41842.492374
```

Finally, you should be aware that pandas provides the function <code>get_dummies()</code> which accomplishes something very similar to <code>OneHotEnconder</code> but may not work as well with the <code>scikit-learn</code> API.

```
[67]: pd.get_dummies(df['BuildingType']).head(5)
       Single-family Townhouse Two-family
[67]:
              1 0
     1
               1
                       0
                                  0
                       Θ
     2
               1
                                  0
                        Θ
                                  0
     3
                1
```

10.7 Optional exercises

Exercise 1: Interactions of explanatory variables

In applied work, we frequently encounter interactions of variables, i.e., an explanatory variable is formed as the product of two other variables. For example, if our data contains the variables x and z, we might add $x \cdot z$ as an independent variable to the model.

To illustrate, consider the following linear model which contains the interaction terms $d \cdot x$, $d \cdot z$ and $x \cdot z$:

$$y_i = \alpha + \beta_1 d_i + \beta_2 x_i + \beta_3 z_i + \beta_4 d_i x_i + \beta_5 d_i z_i + \beta_6 x_i z_i + \epsilon_i$$

where y is a function of (d, x, z) but is only observed with an additive error term ϵ_i . Assume that these variables are mutually independent and distributed as follows:

- d_i is a dummy variable that is 1 with a probability of 60% and zero otherwise;
- x_i is uniformly distributed on the interval [0,1];
- z_i is normally distributed with mean 1 and variance 4; and
- The error term ϵ_i is normally distributed with mean 0 and variance 0.3².

Create an artificial data set of size N=200 and compute y assuming for the following model parameters:

```
[68]: # Intercept
alpha = 0.1

# array containing beta_1, beta_2, ..., beta_6
betas = np.array([0.5, 1.5, 2.0, -0.5, -0.75, -0.1])
```

Perform the following tasks:

- 1. Split the sample into a training data set that comprises the first 150 observations and use the remainder as the test sample.
- 2. Create the matrix of features $\mathbf{X} = [d, x, z, d \cdot x, d \cdot z, x \cdot z]$. Use the PolynomialFeatures preprocessor to create the last three interaction terms.
 - *Hint:* You can use PolynomialFeatures(degree=(2, 2), interaction_only=True) which creates interaction terms where the sum of exponents sums to 2.
- 3. Fit the model as a LinearRegression. Compute the predicted values on the test sample and plot the prediction error against the test sample values of *y*.

4. Create a figure with two panels and plot the model predictions for d = 0 and the left and d = 1 on the right. Each panel should display the predictions for a grid of 101 values for x uniformly spaced on [0,1] on the x-axis. Plot three different lines, one for z evaluated at the 25th, 50th and 75th percentile of the sample distribution. Contrast the predictions with the true values of y.

Exercise 2: Polynomial under- and overfitting

Consider the following non-linear model,

$$y_i = \cos\left(\frac{3}{2}\pi x_i\right) + \epsilon_i$$

where y_i is a trigonometric function of x_i but is measured with an additive error ϵ_i . In this exercise, we are going to approximate y_i using polynomials in x_i of varying degrees:

- 1. Create a sample of size N=50 where the x_i are randomly drawn from a uniform distribution on the interval [0,1] and $\epsilon_i \stackrel{\text{iid}}{\sim} N(0,0.2^2)$. Then generate y_i according to the equation given above.
- 2. Create a scatter plot of the sample (x_i, y_i) and add a line depicting the true non-linear relationship (without measurement error).
- 3. Use the PolynomialFeatures transformation and LinearRegression to approximate y as a polynomial in x. Fit this model using the polynomial degrees $d \in \{0, 1, 2, 3, 6, 10\}$.
- 4. Create a figure with 6 panels, one for each polynomial degree. Each panel should show the sample scatter plot, the true function y = f(x) and the polynomial approximation of a given degree.
- 5. How does the quality of the approximation change as you increase *d*? Do higher-order polynomials always perform better?

Hint: When creating polynomials with PolynomialFeatures(..., include_bias=True), you need to fit the model without an additional intercept as the intercept is already included in the polynomial.

Hint: The cosine function and the constant π are implemented as np.cos() and np.pi in NumPy.

Exercise 3: Polynomial fitting with scikit-learn pipelines

In this exercise, we explore how to simplify fitting models to polynomials using a feature of scikit-learn called pipelines. These allow us to fuse several transformations and an estimation step into a single model.

Continuing with the setup from the previous exercise, we now focus on the model with polynomial degree d = 5. Use either make_pipeline() or the Pipeline class to create a model that combines the PolynomialFeatures transformation and the fitting of a linear model in a single step.

Fit the model using this pipeline. Plot the sample, the true relationship and the predicted value for d = 5 on a grid of 101 x-values that are uniformly spaced on [0,1].

Exercise 4: Optimal polynomial degree with cross-validation

In the previous exercises, we fitted polynomials of varying degrees but did not make any systematic attempt to assess the model fit and choose the preferred polynomial degree (which in this setting is a so-called *hyperparameter*). In this exercise, we explore how cross-validation can be used to tune a hyperparameter in a systematic way.

Consider the following model where y is a non-linear trigonometric function of x and includes an additive error term ϵ ,

$$y_i = \sin(2\pi(x_i + 0.1)) + \epsilon_i$$

and x and ϵ are assumed to be independent.

Proceed as follows:

- 1. Create a sample of N=100 observations. Draw the x_i from the standard uniform distribution on [0,1] and let $\epsilon \stackrel{\text{iid}}{\sim} N(0,0.25)$. Compute y_i according to the equation above.
 - *Hint:* The sine function and the constant π are implemented as np.sin() and np.pi in NumPy.
- 2. Create a sample scatter plot of (x_i, y_i) and add a line showing the true relationship between x and y (without measurement error).
- 3. Program a function compute_splits_mse(d, x, y, n_splits) which takes as arguments the polynomial degree d, the sample observations (x,y) and the number of splits n_splits and returns the mean squared error (MSE) for the test sample for each of the splits. Thus the function should return an array of n_split MSEs.
 - For this exercise, ignore the three-way split into training/validation/test samples illustrated in the lecture and only focus on the training/test samples within each split.
 - *Hint:* You do not need to assign training/test samples manually. Use the KFold class and call its split() method to do the work for you!
 - *Hint:* To compute the MSE for each test sample, you can use mean_squared_error().
- 4. Using the function you just wrote, compute the MSEs for polynomial degrees d = 0, ..., 15 using n_split=10 splits. For each d, compute the average MSE (averaging over 10 splits) and its standard deviation, and use these to create a plot of average MSE on the y-axis against d on the x-axis.
- 5. Which degree *d* results in the lowest average MSE? Explain the intuition behind your findings.
- 6. Add the fitted line using the optimal *d* into the scatter plot you created earlier to visualise your results.

Exercise 5: Automating cross-validation with scikit-learn

In the previous exercise, we performed fitting and cross-validation mostly manually. This exercise explores how most of these steps can be automated using the scikit-learn API.

Using the same setup (functional form for y, sample size and distributional assumptions for x and ϵ) as in the previous exercise, perform the following tasks:

- 1. For each polynomial degree d = 0, ..., 15, define a Pipeline that transforms x into a polynomial and fits a linear model.
 - *Hint:* To do this, you can either create an instance of Pipeline or use the convenience function make_pipeline().
- 2. For each *d*, perform the sample split and computation of MSEs using the function cross_val_score(), which takes as arguments the pipeline you just created, the sample, the metric used to compute the store (scoring=...) and the number of folds (cv=10).
 - *Hint:* To compute the MSE, you need to specify scoring='neg_mean_squared_error'. Note that this returns the *negative* MSE, so you need to correct for this manually.
- 3. Perform the remaining tasks as in the previous exercise, i.e., compute the mean MSE for each *d*, find the *d* that gives the lowest average MSE and plot the average MSE against *d*.

Exercise 6: Optimising hyperparameters with validation curves

This exercise asks you implemented an even faster way to find the optimal polynomial degree d discussed in the previous exercises.

Using the same setup (functional form for y, sample size and distributional assumptions for x and ϵ) as in the previous exercise, perform the following tasks:

- 1. As before, create a Pipeline that performs the polynomial transformation and the fitting of the linear model. You can use any value for PolynomialFeatures(degree=...).
- 2. Use the convenience function validation_curve() to compute the MSE for each d = 0, ..., 15 and all cross-validation splits in a single call.

To do this, the argument param_name specifies which parameter needs to be varied, and param_range determines the set of parameter values that are evaluated.

Since our estimator is a pipeline, the parameter name needs to be specified as 'STEP__ARGUMENT' where STEP is the name of the step in the pipeline we defined (see Pipeline for details), and ARGUMENT determines the argument name to be varied when re-creating the pipeline, which in our case is degree.

Hint: For pipelines created using make_pipeline(), the name of each step in the pipeline is the class name converted to lower case, i.e., for PolynomialFeatures the step name is polynomialfeatures.

3. Perform the remaining tasks as in the previous exercise, i.e., compute the mean MSE for each *d*, find the *d* that gives the lowest average MSE and plot the average MSE against *d*.

The scikit-learn documentation provides additional information on validation curves which you might want to consult.

10.8 Solutions

Exercise 1: Interactions of explanatory variables

Creating the data set

```
[69]: import numpy as np
      import pandas as pd
      from numpy.random import default_rng
      rng = default_rng(1234)
      N = 200
       # Intercept
      alpha = 0.1
       # array containing beta_1, beta_2, ..., beta_6
      betas = np.array([0.5, 1.5, 2.0, -0.5, -0.75, -0.1])
      d = rng.binomial(n=1, p=0.6, size=N)
      x = rng.uniform(0.0, 1.0, size=N)
      z = rng.normal(loc=1.0, scale=2.0, size=N)
      error = rng.normal(loc=0.0, scale=0.3, size=N)
      def compute_y(d, x, z):
           return alpha + betas[0]*d + betas[1]*x + betas[2]*z + betas[3]*d*x \
               + betas[4]*d*z + betas[5]*x*z
      y = compute_y(d, x, z) + error
```

The easiest way to visualise the data set is to convert it to a DataFrame and print the first few rows.

```
[70]: df = pd.DataFrame({'y': y, 'd': d, 'x': x, 'z': z})
df.head(5)
```

```
[70]: y d x z
0 2.611021 0 0.747273 1.030816
1 2.820446 1 0.002071 1.695387
2 5.234120 0 0.809356 2.074067
3 2.525688 1 0.806809 0.901620
4 -0.065962 1 0.054533 -0.594632
```

Creating interactions

The PolynomialFeatures transformer creates multivariate polynomials from a feature matrix \mathbf{X} of a given degree. For example, if we have two variables x and z and set degree=2, it would return a matrix with the columns $[x, x^2, z, z^2, x \cdot z]$. The quadratic terms are not part of our model, so we pass interactions_only=True to keep only the interaction terms.

```
[71]: from sklearn.preprocessing import PolynomialFeatures
from sklearn.model_selection import train_test_split

X = df[['d', 'x', 'z']].to_numpy()

# Number of obs. in training dataset
N_train = 150

# Split into training / test data
X_train, X_test = X[:N_train], X[N_train:]
y_train, y_test = y[:N_train], y[N_train:]

poly = PolynomialFeatures(degree=(2, 2), include_bias=False, interaction_only=True)
X_train_interactions = poly.fit_transform(X_train)
```

It is convenient to use a DataFrame to print a few observations of the interaction terms that were created.

```
[72]: pd.DataFrame(X_train_interactions, columns=['d*x', 'd*z', 'x*z']).head(5)

[72]: d*x d*z x*z
0 0.0000000 0.0000000 0.770301
1 0.002071 1.695387 0.003511
2 0.0000000 0.0000000 1.678660
3 0.806809 0.901620 0.727435
4 0.054533 -0.594632 -0.032427
```

The complete feature matrix **X** of course contains both the original non-interacted variables as well as the interactions, so we stack the two matrices horizontally.

```
[73]: X_train_all = np.hstack((X_train, X_train_interactions))
X_test_all = np.hstack((X_test, poly.transform(X_test)))
```

Fitting a linear regression model

We fit the model in the usual way and use predict() to obtain the predicted values \hat{y}_i for the test data set.

```
[74]: from sklearn.linear_model import LinearRegression

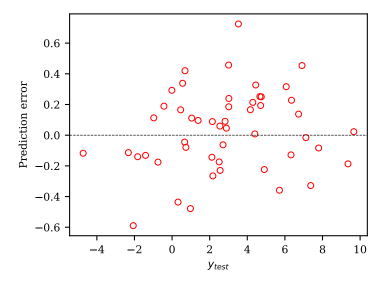
lr = LinearRegression(fit_intercept=False)
lr.fit(X_train_all, y_train)

# Predict response variable on test sample
y_test_hat = lr.predict(X_test_all)
error = y_test - y_test_hat
```

```
[75]: import matplotlib.pyplot as plt

# Plot prediction errors against response variable on test sample
plt.scatter(y_test, error, lw=0.75, color='none', edgecolor='red', s=20)
plt.axhline(0.0, lw=0.5, ls='--', c='black')
plt.xlabel('$y_{test}$')
plt.ylabel('Prediction error')
```

[75]: Text(0, 0.5, 'Prediction error')



Alternative way to construct interactions

Using the preprocessing and pipeline features of scikit-learn, we can also construct the feature matrix in a more elegant way using a FeatureUnion together with a FunctionTransformer.

The FunctionTransformer is initialized using the identity function lambda x: x and hence just returns the argument. The second component of the FeatureUnion is the PolynomialFeatures transformer which creates the interaction term in the same way as before.

```
// Independent of the manually constructed feature matrix

// Verify that these are identical to the manually constructed feature matrix

// x2 = transform_x.fit_transform(X_train)

// np.all(x2 == X_train_all)

// Provided the manually constructed feature matrix

// x2 = transform_x.fit_transform(X_train)

// np.all(x2 == X_train_all)

// Provided the manually constructed feature matrix

// x2 = transform_x.fit_transform(X_train)

// np.all(x2 == X_train_all)

// results for the manually constructed feature matrix

// x2 = transform_x.fit_transform(X_train)

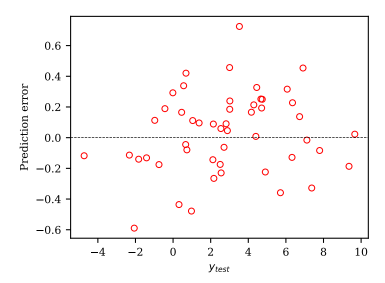
// np.all(x2 == X_train_all)

// x2 = transform_x.fit_train_all(x2 == X_train_all(x2 ==
```

[76]: True

We can now combine this transformer with the LinearRegression into a pipeline to get an object which directly fits the linear model for *y*.

[77]: Text(0, 0.5, 'Prediction error')



Plotting predicted vs. true values

To compute the predicted values on a grid, we first need to create the 3-dimensional feature space. One easy way to do this is to create the 1-dimensional grids for each dimension and then form the Cartesian product using pandas.merge(..., how='cross') which creates the cross-product of two DataFrame objects.

```
[78]: dgrid = np.array([0, 1])
    xgrid = np.linspace(0.0, 1.0, 101)
# Obtain grid for z as 25, 50, 75 percentiles of data
    zgrid = np.percentile(z, q=[25.0, 50.0, 75.0])

df_pred = pd.DataFrame({'d': dgrid})
    df_pred = df_pred.merge(pd.DataFrame({'x': xgrid}), how='cross')
    df_pred = df_pred.merge(pd.DataFrame({'z': zgrid}), how='cross')

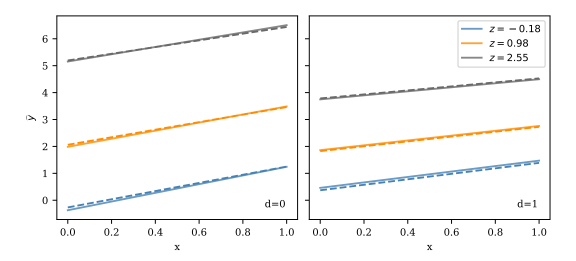
df_pred.head(6)
```

```
[78]: d x z
0 0 0.00 -0.183693
1 0 0.00 0.980220
2 0 0.00 2.546668
3 0 0.01 -0.183693
4 0 0.01 0.980220
5 0 0.01 2.546668
```

We plot the predictions for d = 0 on the left and the ones for d = 1 on the right. Each plot contains three lines, one for each value of z. The model predictions are plotted using solid lines while the true values are dashed.

```
[79]: fig, axes = plt.subplots(1, 2, figsize=(6, 2.75), sharex=True, sharey=True,
           constrained_layout=True)
       colors = ['steelblue', 'darkorange', 'dimgrey']
       for i, d in enumerate(dgrid):
           ax = axes[i]
           df_pred_d = df_pred[df_pred['d'] == d]
           X_pred = df_pred_d.to_numpy()
           y_pred = model.predict(X_pred)
           y_true = compute_y(d, df_pred_d['x'].to_numpy(), df_pred_d['z'].to_numpy())
           y_pred = y_pred.reshape((len(xgrid), len(zgrid)))
           y_true = y_true.reshape(y_pred.shape)
           for j, zj in enumerate(zgrid):
               ax.plot(xgrid, y_pred[:, j], c=colors[j], label=f'$z = {zj:.2f}$', alpha=0.8)
               ax.plot(xgrid, y_true[:, j], c=colors[j], ls='--')
           ax.set_xlabel('x')
           ax.text(0.95, 0.05, f'd={d}', transform=ax.transAxes, va='bottom', ha='right')
       # Add legend to right panel
       axes[1].legend()
       # Add y-labels only to left panel
       axes[0].set_ylabel(r'$\widehat{y}$')
```

[79]: Text(0, 0.5, '\$\\widehat{y}\$')



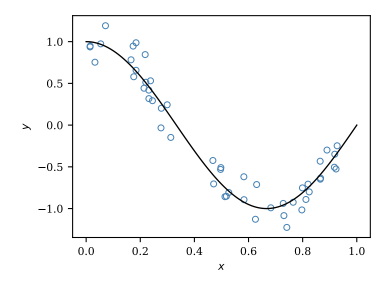
Exercise 2: Polynomial under- and overfitting

Creating and plotting the sample

[82]: Text(0, 0.5, '\$y\$')

We randomly draw values for x_i and ε_i for each observation. We define a function fcn(x) which returns the true value for y without measurement error.

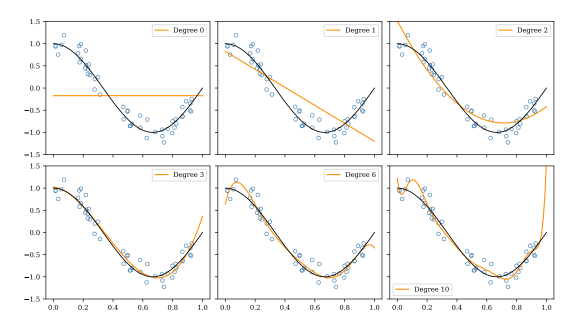
```
[80]: import numpy as np
       # True function (w/o errors)
       def fcn(x):
           return np.cos(1.5 * np.pi * x)
[81]: from numpy.random import default_rng
       # Initialise random number generator
       rng = default_rng(123)
       # Sample size
       N = 50
       # Randomly draw explanatory variable x
       x = rng.random(size=N)
       epsilon = rng.normal(scale=0.2, size=N)
       y = fcn(x) + epsilon
[82]: import matplotlib.pyplot as plt
       xvalues = np.linspace(0.0, 1.0, 101)
       plt.plot(xvalues, fcn(xvalues), color='black', lw=1.0, label='True function')
       plt.scatter(x, y, s=20, color='none', edgecolor='steelblue',
           lw=0.75, label='Sample')
       plt.xlabel('$x$')
       plt.ylabel('$y$')
```



Polynomial approximations

To plot the predicted values for each value of the polynomial degree d, we create a 2-by-3 figure and iterate over the axes objects. For each panel and corresponding d, we create a transformation as PolynomialFeatures(degree=d, include_bias=True) and use it to create a polynomial in x.

```
[83]: from sklearn.preprocessing import PolynomialFeatures
       from sklearn.linear_model import LinearRegression
       degrees = np.array([0, 1, 2, 3, 6, 10])
       ncol = 3
       nrow = int(np.ceil(len(degrees) / ncol))
       fig, axes = plt.subplots(nrow, ncol, figsize=(9, 5),
           sharex=True, sharey=True,
           constrained_layout=True
       axes = axes.flatten()
       for i, ax in enumerate(axes):
           d = degrees[i]
           poly = PolynomialFeatures(degree=d, include_bias=True)
           poly.fit(x[:, None])
           Xpoly = poly.transform(x[:, None])
           lr = LinearRegression(fit_intercept=False)
           lr.fit(Xpoly, y)
           y_hat = lr.predict(poly.transform(xvalues[:, None]))
           ax.plot(xvalues, y_hat, lw=1.25, c='darkorange', label=f'Degree {d}')
           ax.plot(xvalues, fcn(xvalues), color='black', lw=1.0)
           ax.scatter(x, y, s=20, color='none', edgecolor='steelblue', lw=0.75)
           ax.set_ylim((-1.5, 1.5))
           ax.legend()
```



As the figure shows, a degree-0 polynomial is just a constant, while a polynomial of degree 1 is linear in x. Neither fits the true relationship very well ("underfitting"), but the fit initially improves as we increase d. For high d, on the other hand, the polynomial becomes too flexible and responds strongly to local "noise" introduced by measurement error ("overfitting").

Exercise 3: Polynomial fitting with scikit-learn pipelines

We create the sample in the same way as in exercise 2.

```
[84]: import numpy as np

# True function (w/o errors)
def fcn(x):
    return np.cos(1.5 * np.pi * x)

[85]: from numpy.random import default_rng
```

```
from numpy.random import default_rng

# Initialise random number generator
rng = default_rng(123)

# Sample size
N = 100

# Randomly draw explanatory variable x
x = np.sort(rng.random(size=N))
epsilon = rng.normal(scale=0.2, size=N)
y = fcn(x) + epsilon
```

To transform x and fit the linear model in a single step, we create a pipepline using make_pipeline() as follows:

```
[86]: from sklearn.preprocessing import PolynomialFeatures
    from sklearn.pipeline import make_pipeline
    from sklearn.linear_model import LinearRegression

degree = 5

# Create PipeLine object that consist of a polynomial transformation and
# a linear regression model
pipe_lr = make_pipeline(
```

```
PolynomialFeatures(
    degree=degree,
    include_bias=False
),
LinearRegression(fit_intercept=True)
)
```

In Jupyter notebooks, printing the pipeline object produces a nice graphical representation that can be unfolded to see any parameters for each step of the pipeline (this will not be visible in the PDF version of this notebook).

```
[87]: # Show graphical representation of pipeline (only in Jupyter notebook)
pipe_lr
```

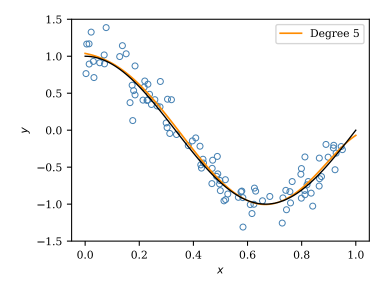
The input data can be transformed and the model fit in a single step by calling the fit() method of the pipeline object.

```
[88]: # Use pipeline to fit model
pipe_lr.fit(x[:, None], y)

# Compute predicted values on uniform grid
xvalues = np.linspace(0.0, 1.0, 101)
y_hat = pipe_lr.predict(xvalues[:, None])

# Plot predicted values
plt.plot(xvalues, y_hat, lw=1.25, c='darkorange', label=f'Degree {degree}')
# Plot true function
plt.plot(xvalues, fcn(xvalues), color='black', lw=1.0)
# Plot sample data
plt.scatter(x, y, s=20, color='none', edgecolor='steelblue', lw=0.75)
plt.ylim((-1.5, 1.5))
plt.ylabel('$y$')
plt.xlabel('$x$')
plt.legend()
```

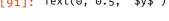
[88]: <matplotlib.legend.Legend at 0x7fca6641b3d0>

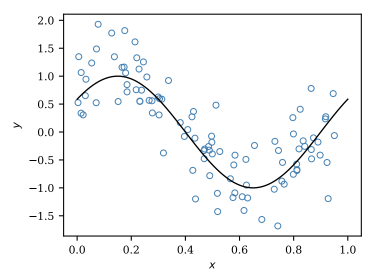


Exercise 4: Optimal polynomial degree with cross-validation

Creating and plotting the sample

```
[89]: import numpy as np
       # True function (w/o errors)
       def fcn(x):
           return np.sin(2.0 * np.pi * (x + 0.1))
[90]: from numpy.random import default_rng
       # Initialise random number generator
       rng = default_rng(123)
       # Sample size
       N = 100
       # Randomly draw explanatory variable x
       x = rng.random(size=N)
       epsilon = rng.normal(scale=0.5, size=N)
       y = fcn(x) + epsilon
[91]: import matplotlib.pyplot as plt
       xvalues = np.linspace(0.0, 1.0, 101)
       plt.plot(xvalues, fcn(xvalues), color='black', lw=1.0, label='True function')
       plt.scatter(x, y, s=20, color='none', edgecolor='steelblue',
           lw=0.75, label='Sample'
       plt.xlabel('$x$')
       plt.ylabel('$y$')
[91]: Text(0, 0.5, '$y$')
```





Function to perform fitting and MSE computation

The following implementation uses two distinct functions:

- 1. fit_poly() takes a polynomial degree d and the sample data, creates the polynomial and fits a linear model.
- 2. compute_splits_mse() splits the sample into n_splits folds, fits the model on the training subset of each split and computes the MSE on the test (or validation) sample.

```
[92]: from sklearn.preprocessing import PolynomialFeatures
from sklearn.linear_model import LinearRegression

# Function to perform polynomial transformation and model fitting.
# Alternatively, we could use a Pipeline (see next exercise)
def fit_poly(d, X, y):
    poly = PolynomialFeatures(degree=d, include_bias=True)
    poly.fit(X)
    Xpoly = poly.transform(X)
    lr = LinearRegression(fit_intercept=False)
    lr.fit(Xpoly, y)

# Return poly transformer (required for prediction) and fitted model
    return poly, lr
```

```
[93]: from sklearn.model_selection import KFold
       from sklearn.metrics import mean_squared_error
       # Function to compute MSE for given number of splits
      def compute_splits_mse(d, x, y, n_splits=10):
           # Split sample into train/test blocks for k-fold validation
           kf = KFold(n_splits=n_splits)
           # list to store MSE for each CV split
           mse_splits = []
           # Manually iterate over folds (train/test combinations)
           for itrain, itest in kf.split(x):
              x_train = x[itrain]
              x_{test} = x[itest]
              y_train = y[itrain]
              y_test = y[itest]
               poly, lr = fit_poly(d, x_train[:, None], y_train)
              Xpoly_test = poly.transform(x_test[:, None])
              y_test_hat = lr.predict(Xpoly_test)
              mse = mean_squared_error(y_test, y_test_hat)
              mse splits.append(mse)
           # Convert to array
           mse_splits = np.array(mse_splits)
           return mse_splits
```

We can test this function for a degree-0 polynomial (i.e., a constant function) on our sample data. The functions returns 10 different MSE values, one for the test sample of each split.

```
[94]: compute_splits_mse(0, x, y, n_splits=10)

[94]: array([1.05594738, 1.19357336, 0.68425812, 0.21465842, 0.64290297, 0.89604156, 0.44140114, 0.51936866, 0.83432109, 0.59097661])
```

Computing the MSE for each hyperparameter

We now use the functions we defined to evaluate the MSEs for each polynomial degree d. For each d, compute_splits_mse() returns 10 values, so we compute the average MSE and its standard deviation.

```
[95]: degrees = np.arange(16)

mse_mean = []
mse_std = []

for d in degrees:

    mse_splits = compute_splits_mse(d, x, y, n_splits=10)

    mse_mean.append(np.mean(mse_splits))
    mse_std.append(np.std(mse_splits))

# Convert to NumPy arrays
mse_mean = np.array(mse_mean)
mse_std = np.array(mse_std)

# Print MSEs for each degree
mse_mean
```

```
[95]: array([0.70734493, 0.42671672, 0.33501395, 0.23900256, 0.21559544, 0.22029082, 0.22568436, 0.22577954, 0.22433218, 0.23387504, 0.23689982, 0.24623706, 0.25145758, 0.26281712, 0.27462137, 0.29084427])
```

Finding the optimal parameter

We use np.argmin() to find the *index* of the smallest average MSE.

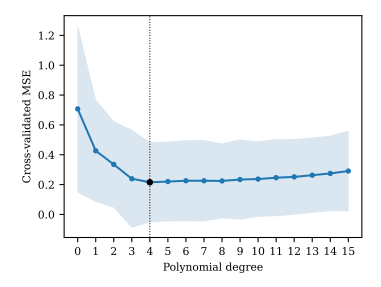
```
[96]: # Polynomial degree that minimises MSE
imin = np.argmin(mse_mean)
dmin = degrees[imin]

print(f'Polynomial degree with min. MSE: {dmin}')
```

Polynomial degree with min. MSE: 4

The results can be visualised by plotting the average MSE by polynomial degree. As the graph shows, the minimum is obtained at d = 4. Intuitively, for low d the model underfits the data, whereas overfitting occurs for high values of d.

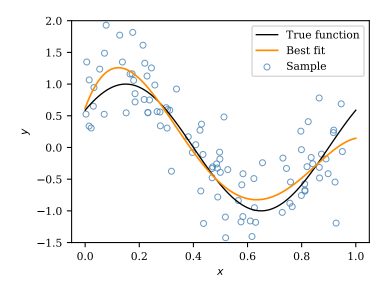
[97]: <matplotlib.lines.Line2D at 0x7fca664b8670>



Plotting the fitted model

We use the function $fit_poly()$ to re-fit the model at the optimal d and we plot the predicted values on a uniformly spaced grid of x-values.

[98]: <matplotlib.legend.Legend at 0x7fca663c6440>



Exercise 5: Automating cross-validation with scikit-learn

Creating the sample

[99]: import numpy as np

We create the sample exactly as in the previous exercise.

```
# True function (w/o errors)
def fcn(x):
    return np.sin(2.0 * np.pi * (x + 0.1))

[100]: from numpy.random import default_rng

# Initialise random number generator
rng = default_rng(123)

# Sample size
N = 100

# Randomly draw explanatory variable x
x = rng.random(size=N)
epsilon = rng.normal(scale=0.5, size=N)
y = fcn(x) + epsilon
```

Computing the MSE for each hyperparameter

For each d, we create a pipeline. Note that step one of the pipeline, polynomial transformation, depends on d so we need to recreate the pipeline in each iteration of the loop!

We then use <code>cross_val_score()</code> to compute the MSEs for each split. This replaces the function we implemented in the previous exercise with much shorter code.

```
[101]: from sklearn.model_selection import cross_val_score
    from sklearn.pipeline import make_pipeline
    from sklearn.preprocessing import PolynomialFeatures
    from sklearn.linear_model import LinearRegression

degrees = np.arange(16)
```

```
mse_mean = []
        mse\_std = []
        for d in degrees:
            pipe = make_pipeline(
                PolynomialFeatures(
                    degree=d,
                    include_bias=True
                ),
                LinearRegression(fit_intercept=False)
            scores = cross_val_score(
                pipe,
                x[:, None], y,
                scoring='neg_mean_squared_error',
            # Stores contain NEGATIVE MSE
            mse_mean.append(np.mean(-scores))
            mse_std.append(np.std(-scores))
        # Convert to NumPy arrays
        mse_mean = np.array(mse_mean)
        mse_std = np.array(mse_std)
        # Print avg MSEs for each degree
        mse_mean
[101]: array([0.70734493, 0.42671672, 0.33501395, 0.23900256, 0.21559544,
                \hbox{\tt 0.22029082, 0.22568436, 0.22577954, 0.22433218, 0.23387504, } 
               0.23689982, 0.24623706, 0.25145758, 0.26281712, 0.27462137,
               0.29084427])
```

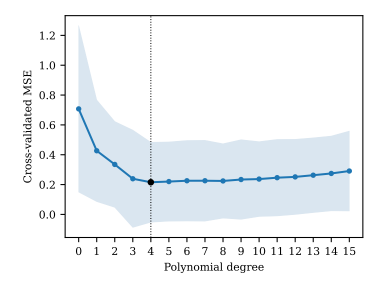
The remained of the exercise proceeds in the same way as before. Assuringly, we also find the same results.

```
[102]: imin = np.argmin(mse_mean)
       dmin = degrees[imin]
       print(f'Polynomial degree with min. MSE: {dmin}')
```

Polynomial degree with min. MSE: 4

```
[103]: plt.plot(degrees, mse_mean, marker='o', ms=3)
       plt.fill_between(degrees, mse_mean-2.0*mse_std, mse_mean+2.0*mse_std,
            lw=0.25, color='steelblue', alpha=0.2, zorder=-1)
       plt.xlabel('Polynomial degree')
       plt.ylabel('Cross-validated MSE')
       plt.scatter(degrees[imin], mse_mean[imin], s=15, c='black', zorder=100)
       plt.xticks(degrees)
       plt.axvline(imin, ls=':', lw=0.75, c='black')
```

[103]: <matplotlib.lines.Line2D at 0x7fca662af700>



Exercise 6: Optimising hyperparameters with validation curves

Creating the sample

We create the sample in the same way we did in exercises 4 and 5.

```
[104]: import numpy as np

# True function (w/o errors)
def fcn(x):
    return np.sin(2.0 * np.pi * (x + 0.1))

[105]: from numpy.random import default_rng

# Initialise random number generator
rng = default_rng(123)

# Sample size
N = 100

# Randomly draw explanatory variable x
x = rng.random(size=N)
epsilon = rng.normal(scale=0.5, size=N)
y = fcn(x) + epsilon
```

Computing the MSE for each hyperparameter

To use validation_curve(), we need to correctly specify the param_name argument. In our case this is given by 'polynomialfeatures__degree' since we are asking the function to vary the argument degree of the pipeline step called polynomialfeatures. Note that the __ serves as a delimiter between the pipeline step name and the argument name.

```
[106]: from sklearn.model_selection import validation_curve

# Set of polynomial degrees to cross-validate
degrees = np.arange(16)

# Create pipeline. The degree value does not matter at this point, will be
# automatically updated.
```

```
pipe = make_pipeline(
    PolynomialFeatures(
        degree=⊙,
        include_bias=True
    LinearRegression(fit_intercept=False)
\# Compute the MSEs for each degree and each split, returning arrays of
# size 15 x 10.
train_scores, test_scores = validation_curve(
    estimator=pipe,
   X=x[:, None], y=y,
    param_name='polynomialfeatures__degree',
    param_range=degrees,
    scoring='neg_mean_squared_error',
)
# Compute mean and std for each degree (scored returned by function are
# NEGATIVE MSEs)
mse_mean = np.mean(-test_scores, axis=1)
mse_std = np.std(-test_scores, axis=1)
```

The remainder of this exercise proceeds as before.

```
[107]: imin = np.argmin(mse_mean)
    dmin = degrees[imin]

print(f'Polynomial degree with min. MSE: {dmin}')
```

Polynomial degree with min. MSE: 4

[108]: <matplotlib.lines.Line2D at 0x7fca66113d30>

