

Machine Learning (CCS4340)

Lab 04: Introducing Scikit-Learn

There are several Python libraries that provide solid implementations of a range of machine learning algorithms. One of the best known is [Scikit-Learn](#), a package that provides efficient versions of a large number of common algorithms. Scikit-Learn is characterized by a clean, uniform, and streamlined API, as well as by very useful and complete online documentation. A benefit of this uniformity is that once you understand the basic use and syntax of Scikit-Learn for one type of model, switching to a new model or algorithm is straightforward.

This lab session provides an overview of the Scikit-Learn API. A solid understanding of these API elements will form the foundation for understanding the deeper practical discussion of machine learning algorithms and approaches in the following lab sessions.

We will start by covering data representation in Scikit-Learn, then delve into the Estimator API, and finally go through a more interesting example of using these tools for exploring a set of images of handwritten digits.

Data Representation in Scikit-Learn

Machine learning is about creating models from data: for that reason, we'll start by discussing how data can be represented. The best way to think about data within Scikit-Learn is in terms of *tables*.

A basic table is a two-dimensional grid of data, in which the rows represent individual elements of the dataset, and the columns represent quantities related to each of these elements. For example, consider the [Iris dataset](#), famously analyzed by Ronald Fisher in 1936. We can download this dataset in the form of a Pandas `DataFrame` using the [Seaborn](#) library, and take a look at the first few items:

```
In [1]: import seaborn as sns  
iris = sns.load_dataset('iris')  
iris.head()
```

```
Out[1]:
```

	sepal_length	sepal_width	petal_length	petal_width	species
0	5.1	3.5	1.4	0.2	setosa
1	4.9	3.0	1.4	0.2	setosa
2	4.7	3.2	1.3	0.2	setosa
3	4.6	3.1	1.5	0.2	setosa
4	5.0	3.6	1.4	0.2	setosa

Here each row of the data refers to a single observed flower, and the number of rows is the total number of flowers in the dataset. In general, we will refer to the rows of the matrix as

samples, and the number of rows as `n_samples`.

Likewise, each column of the data refers to a particular quantitative piece of information that describes each sample. In general, we will refer to the columns of the matrix as *features*, and the number of columns as `n_features`.

The Features Matrix

The table layout makes clear that the information can be thought of as a two-dimensional numerical array or matrix, which we will call the *features matrix*. By convention, this matrix is often stored in a variable named `X`. The features matrix is assumed to be two-dimensional, with shape `[n_samples, n_features]`, and is most often contained in a NumPy array or a Pandas `DataFrame`, though some Scikit-Learn models also accept SciPy sparse matrices.

The samples (i.e., rows) always refer to the individual objects described by the dataset. For example, a sample might represent a flower, a person, a document, an image, a sound file, a video, an astronomical object, or anything else you can describe with a set of quantitative measurements.

The features (i.e., columns) always refer to the distinct observations that describe each sample in a quantitative manner. Features are often real-valued, but may be Boolean or discrete-valued in some cases.

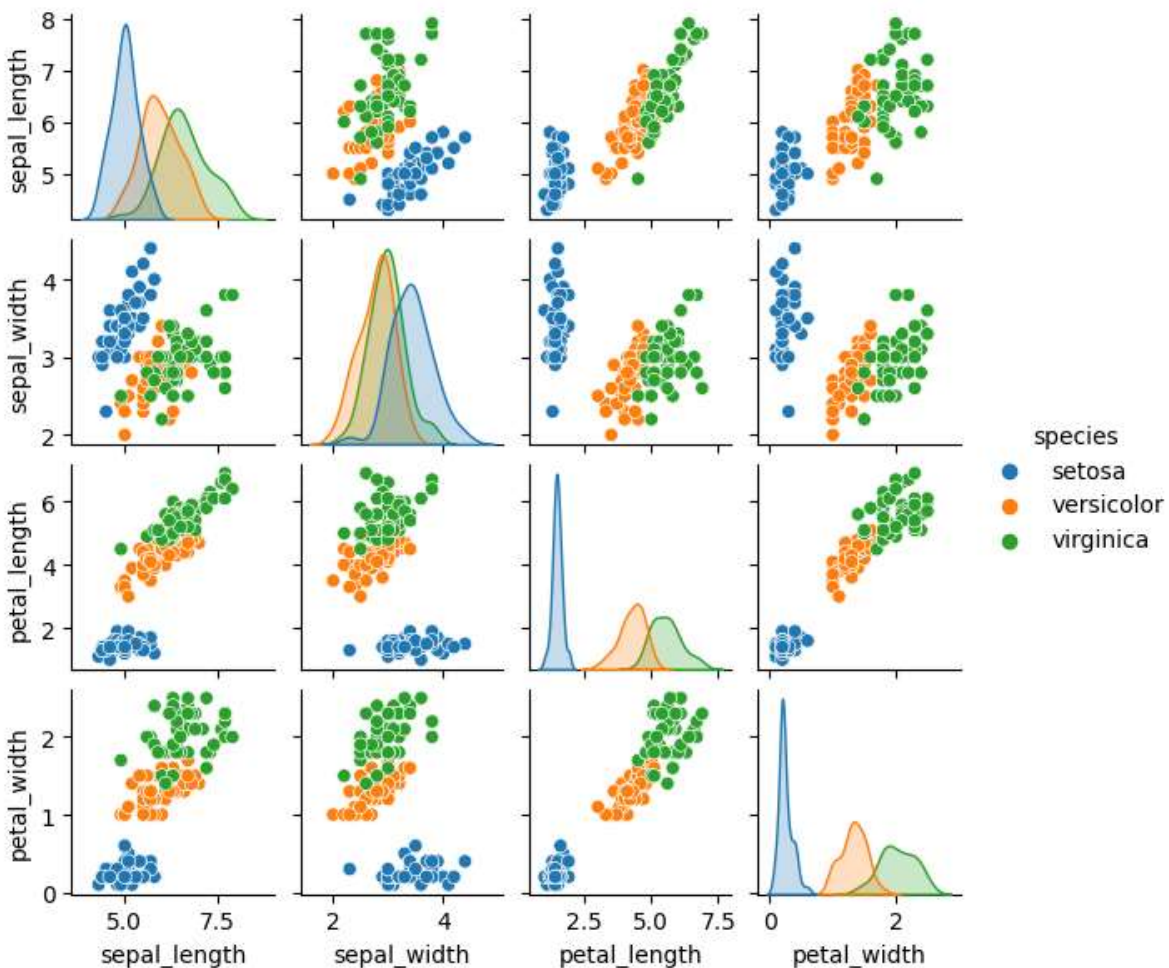
The Target Array

In addition to the feature matrix `X`, we also generally work with a *label* or *target* array, which by convention we will usually call `y`. The target array is usually one-dimensional, with length `n_samples`, and is generally contained in a NumPy array or Pandas `Series`. The target array may have continuous numerical values, or discrete classes/labels. While some Scikit-Learn estimators do handle multiple target values in the form of a two-dimensional, `[n_samples, n_targets]` target array, we will primarily be working with the common case of a one-dimensional target array.

A common point of confusion is how the target array differs from the other feature columns. The distinguishing characteristic of the target array is that it is usually the quantity we want to *predict from the features*: in statistical terms, it is the dependent variable. For example, given the preceding data we may wish to construct a model that can predict the species of flower based on the other measurements; in this case, the `species` column would be considered the target array.

With this target array in mind, we can use Seaborn to conveniently visualize the data (see the following figure):

```
In [2]: %matplotlib inline  
import seaborn as sns  
sns.pairplot(iris, hue='species', height=1.5);
```



For use in Scikit-Learn, we will extract the features matrix and target array from the `DataFrame`, which we can do using some of the Pandas `DataFrame` operations discussed.

```
In [3]: X_iris = iris.drop('species', axis=1)
X_iris.shape
```

```
Out[3]: (150, 4)
```

```
In [4]: y_iris = iris['species']
y_iris.shape
```

```
Out[4]: (150,)
```

With this data properly formatted, we can move on to consider Scikit-Learn's Estimator API.

The Estimator API

The Scikit-Learn API is designed with the following guiding principles in mind, as outlined in the [Scikit-Learn API paper](#):

- *Consistency*: All objects share a common interface drawn from a limited set of methods, with consistent documentation.
- *Inspection*: All specified parameter values are exposed as public attributes.

- *Limited object hierarchy*: Only algorithms are represented by Python classes; datasets are represented in standard formats (NumPy arrays, Pandas `DataFrame` objects, SciPy sparse matrices) and parameter names use standard Python strings.
- *Composition*: Many machine learning tasks can be expressed as sequences of more fundamental algorithms, and Scikit-Learn makes use of this wherever possible.
- *Sensible defaults*: When models require user-specified parameters, the library defines an appropriate default value.

In practice, these principles make Scikit-Learn very easy to use, once the basic principles are understood. Every machine learning algorithm in Scikit-Learn is implemented via the Estimator API, which provides a consistent interface for a wide range of machine learning applications.

Basics of the API

Most commonly, the steps in using the Scikit-Learn Estimator API are as follows:

1. Choose a class of model by importing the appropriate estimator class from Scikit-Learn.
2. Choose model hyperparameters by instantiating this class with desired values.
3. Arrange data into a features matrix and target vector, as outlined earlier in this chapter.
4. Fit the model to your data by calling the `fit` method of the model instance.
5. Apply the model to new data:
 - For supervised learning, often we predict labels for unknown data using the `predict` method.
 - For unsupervised learning, we often transform or infer properties of the data using the `transform` or `predict` method.

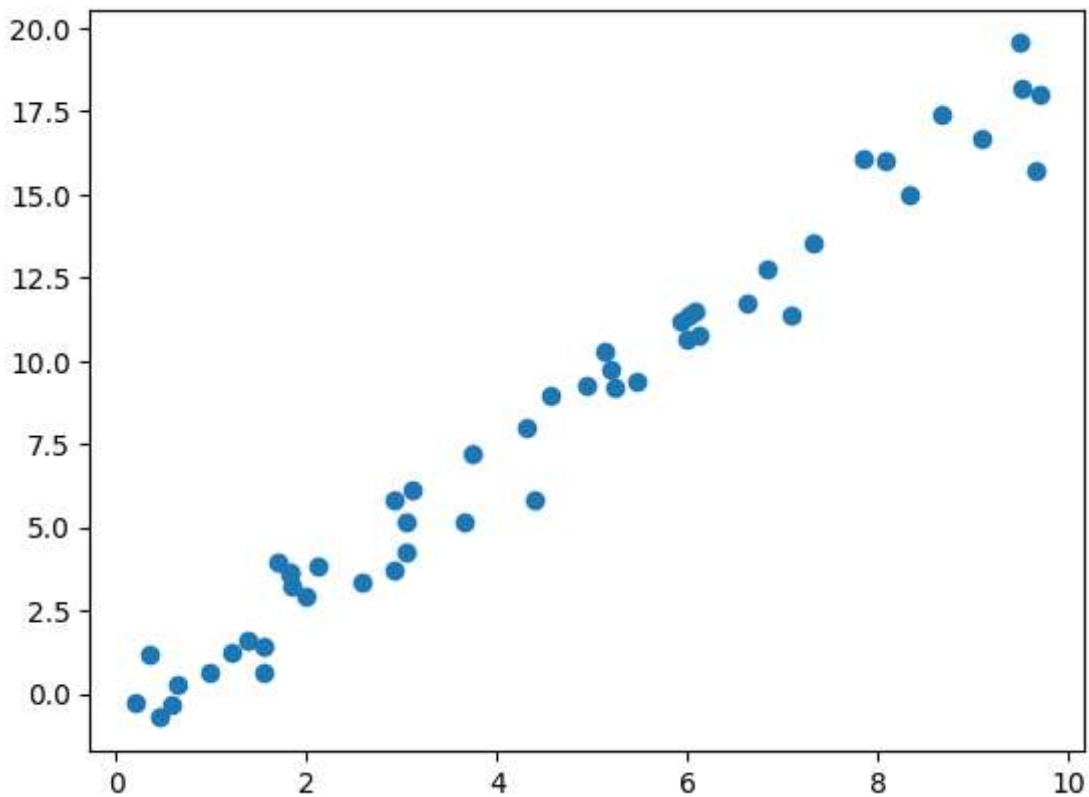
We will now step through several simple examples of applying supervised and unsupervised learning methods.

Supervised Learning Example: Simple Linear Regression

As an example of this process, let's consider a simple linear regression—that is, the common case of fitting a line to (x, y) data. We will use the following simple data for our regression example (see the following figure):

```
In [5]: import matplotlib.pyplot as plt
import numpy as np

rng = np.random.RandomState(42)
x = 10 * rng.rand(50)
y = 2 * x - 1 + rng.randn(50)
plt.scatter(x, y);
```



With this data in place, we can use the recipe outlined earlier. Let's walk through the process:

1. Choose a class of model

In Scikit-Learn, every class of model is represented by a Python class. So, for example, if we would like to compute a simple `LinearRegression` model, we can import the linear regression class:

```
In [6]: from sklearn.linear_model import LinearRegression
```

Note that other more general linear regression models exist as well; you can read more about them in the `sklearn.linear_model` [module documentation](#).

2. Choose model hyperparameters

An important point is that *a class of model is not the same as an instance of a model*.

Once we have decided on our model class, there are still some options open to us. Depending on the model class we are working with, we might need to answer one or more questions like the following:

- Would we like to fit for the offset (i.e., y -intercept)?
- Would we like the model to be normalized?
- Would we like to preprocess our features to add model flexibility?
- What degree of regularization would we like to use in our model?
- How many model components would we like to use?

These are examples of the important choices that must be made *once the model class is selected*. These choices are often represented as *hyperparameters*, or parameters that must

be set before the model is fit to data. In Scikit-Learn, hyperparameters are chosen by passing values at model instantiation. We will explore how you can quantitatively choose hyperparameters in next lab session.

For our linear regression example, we can instantiate the `LinearRegression` class and specify that we would like to fit the intercept using the `fit_intercept` hyperparameter:

```
In [7]: model = LinearRegression(fit_intercept=True)  
model
```

```
Out[7]: ▾ LinearRegression  
LinearRegression()
```

Keep in mind that when the model is instantiated, the only action is the storing of these hyperparameter values. In particular, we have not yet applied the model to any data: the Scikit-Learn API makes very clear the distinction between *choice of model* and *application of model to data*.

3. Arrange data into a features matrix and target vector

Previously we examined the Scikit-Learn data representation, which requires a two-dimensional features matrix and a one-dimensional target array. Here our target variable `y` is already in the correct form (a length-`n_samples` array), but we need to massage the data `x` to make it a matrix of size `[n_samples, n_features]`. In this case, this amounts to a simple reshaping of the one-dimensional array:

```
In [8]: X = x[:, np.newaxis]  
X.shape
```

```
Out[8]: (50, 1)
```

4. Fit the model to the data

Now it is time to apply our model to the data. This can be done with the `fit` method of the model:

```
In [9]: model.fit(X, y)
```

```
Out[9]: ▾ LinearRegression  
LinearRegression()
```

This `fit` command causes a number of model-dependent internal computations to take place, and the results of these computations are stored in model-specific attributes that the user can explore. In Scikit-Learn, by convention all model parameters that were learned during the `fit` process have trailing underscores; for example in this linear model, we have the following:

```
In [10]: model.coef_
```

```
Out[10]: array([1.9776566])
```

```
In [11]: model.intercept_
```

```
Out[11]: -0.903310725531111
```

These two parameters represent the slope and intercept of the simple linear fit to the data. Comparing the results to the data definition, we see that they are close to the values used to generate the data: a slope of 2 and intercept of -1 .

One question that frequently comes up regards the uncertainty in such internal model parameters. In general, Scikit-Learn does not provide tools to draw conclusions from internal model parameters themselves: interpreting model parameters is much more a *statistical modeling* question than a *machine learning* question. Machine learning instead focuses on what the model *predicts*. If you would like to dive into the meaning of fit parameters within the model, other tools are available, including the [statsmodels Python package](#).

5. Predict labels for unknown data

Once the model is trained, the main task of supervised machine learning is to evaluate it based on what it says about new data that was not part of the training set. In Scikit-Learn, this can be done using the `predict` method. For the sake of this example, our "new data" will be a grid of x values, and we will ask what y values the model predicts:

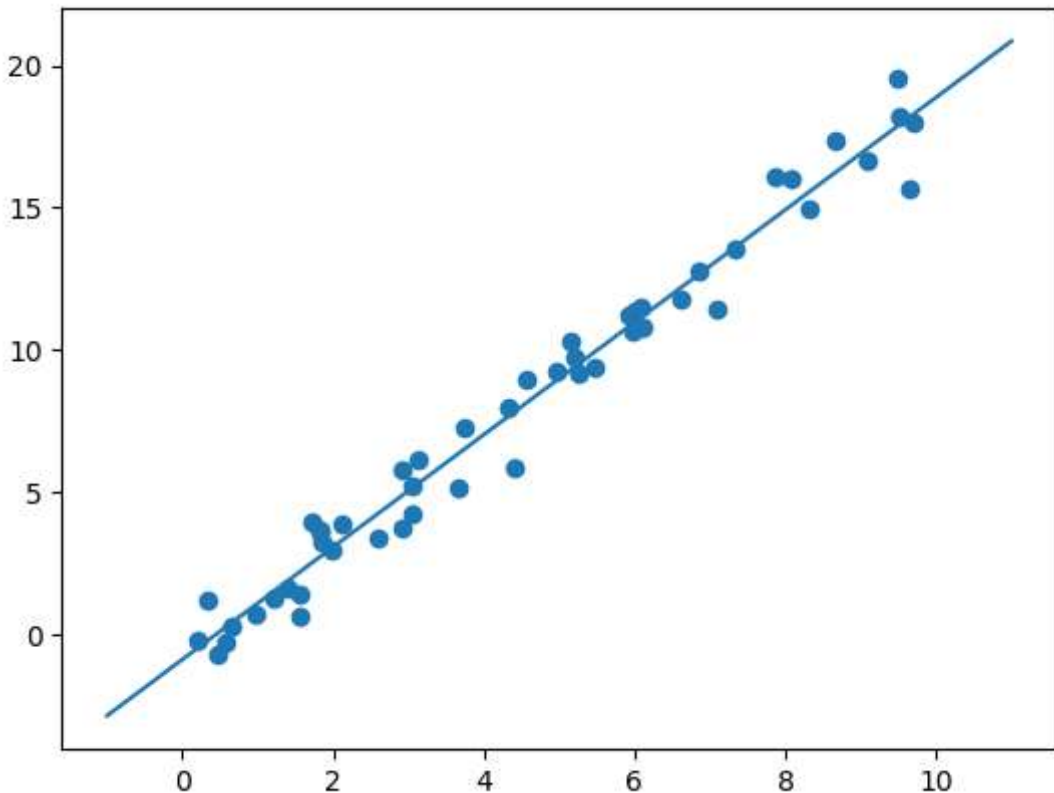
```
In [12]: xfit = np.linspace(-1, 11)
```

As before, we need to coerce these x values into a `[n_samples, n_features]` features matrix, after which we can feed it to the model:

```
In [13]: Xfit = xfit[:, np.newaxis]
yfit = model.predict(Xfit)
```

Finally, let's visualize the results by plotting first the raw data, and then this model fit (see the following figure):

```
In [14]: plt.scatter(x, y)
plt.plot(xfit, yfit);
```



Typically the efficacy of the model is evaluated by comparing its results to some known baseline, as we will see in the next example.

Supervised Learning Example: Iris Classification

Let's take a look at another example of this process, using the Iris dataset we discussed earlier. Our question will be this: given a model trained on a portion of the Iris data, how well can we predict the remaining labels?

For this task, we will use a simple generative model known as *Gaussian naive Bayes*, which proceeds by assuming each class is drawn from an axis-aligned Gaussian distribution.

Because it is so fast and has no hyperparameters to choose, Gaussian naive Bayes is often a good model to use as a baseline classification, before exploring whether improvements can be found through more sophisticated models.

We would like to evaluate the model on data it has not seen before, so we will split the data into a *training set* and a *testing set*. This could be done by hand, but it is more convenient to use the `train_test_split` utility function:

```
In [15]: from sklearn.model_selection import train_test_split
Xtrain, Xtest, ytrain, ytest = train_test_split(X_iris, y_iris,
                                              random_state=1)
```

With the data arranged, we can follow our recipe to predict the labels:

```
In [16]: from sklearn.naive_bayes import GaussianNB # 1. choose model class
model = GaussianNB() # 2. instantiate model
model.fit(Xtrain, ytrain) # 3. fit model to data
y_model = model.predict(Xtest) # 4. predict on new data
```

Finally, we can use the `accuracy_score` utility to see the fraction of predicted labels that match their true values:

```
In [17]: from sklearn.metrics import accuracy_score  
accuracy_score(ytest, y_model)
```

```
Out[17]: 0.9736842105263158
```

With an accuracy topping 97%, we see that even this very naive classification algorithm is effective for this particular dataset!

Unsupervised Learning Example: Iris Dimensionality

As an example of an unsupervised learning problem, let's take a look at reducing the dimensionality of the Iris data so as to more easily visualize it. Recall that the Iris data is four-dimensional: there are four features recorded for each sample.

The task of dimensionality reduction centers around determining whether there is a suitable lower-dimensional representation that retains the essential features of the data. Often dimensionality reduction is used as an aid to visualizing data: after all, it is much easier to plot data in two dimensions than in four dimensions or more!

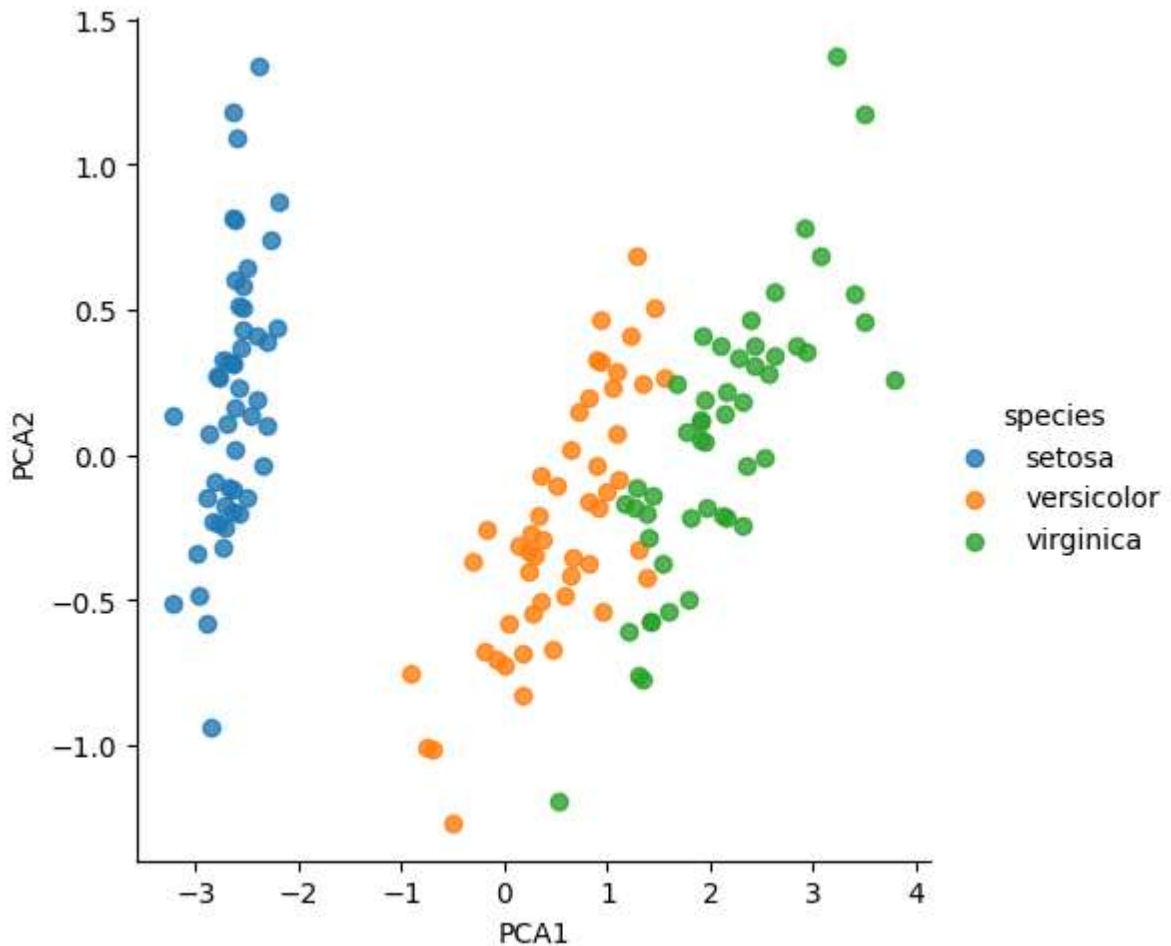
Here we will use *principal component analysis*, which is a fast linear dimensionality reduction technique. We will ask the model to return two components—that is, a two-dimensional representation of the data.

Following the sequence of steps outlined earlier, we have:

```
In [18]: from sklearn.decomposition import PCA # 1. Choose the model class  
model = PCA(n_components=2) # 2. Instantiate the model  
model.fit(X_iris) # 3. Fit to data  
X_2D = model.transform(X_iris) # 4. Transform the data
```

Now let's plot the results. A quick way to do this is to insert the results into the original Iris `DataFrame`, and use Seaborn's `lmplot` to show the results (see the following figure):

```
In [19]: iris['PCA1'] = X_2D[:, 0]  
iris['PCA2'] = X_2D[:, 1]  
sns.lmplot(x="PCA1", y="PCA2", hue='species', data=iris, fit_reg=False);
```



We see that in the two-dimensional representation, the species are fairly well separated, even though the PCA algorithm had no knowledge of the species labels! This suggests to us that a relatively straightforward classification will probably be effective on the dataset, as we saw before.

Unsupervised Learning Example: Iris Clustering

Let's next look at applying clustering to the Iris data. A clustering algorithm attempts to find distinct groups of data without reference to any labels. Here we will use a powerful clustering method called a *Gaussian mixture model* (GMM), discussed in more detail in future lab sessions.

A GMM attempts to model the data as a collection of Gaussian blobs.

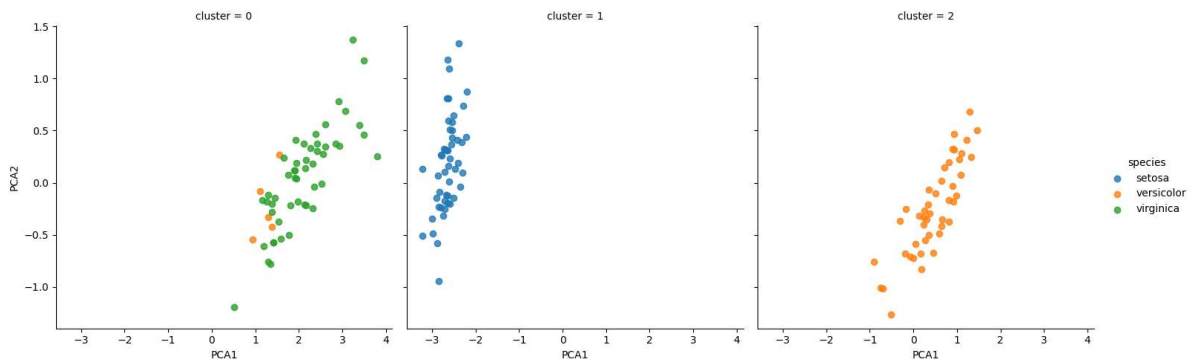
We can fit the Gaussian mixture model as follows:

```
In [20]: from sklearn.mixture import GaussianMixture      # 1. Choose the model class
model = GaussianMixture(n_components=3,                 # 2. Instantiate the model
                       covariance_type='full')    # 3. Fit to data
model.fit(X_iris)                                     # 4. Determine labels
y_gmm = model.predict(X_iris)
```

```
C:\ProgramData\anaconda3\lib\site-packages\sklearn\cluster\_kmeans.py:1382: UserWarning: KMeans is known to have a memory leak on Windows with MKL, when there are 1 less chunks than available threads. You can avoid it by setting the environment variable OMP_NUM_THREADS=1.
warnings.warn(
```

As before, we will add the cluster label to the Iris `DataFrame` and use Seaborn to plot the results (see the following figure):

```
In [21]: iris['cluster'] = y_gmm
sns.lmplot(x="PCA1", y="PCA2", data=iris, hue='species',
            col='cluster', fit_reg=False);
```



By splitting the data by cluster number, we see exactly how well the GMM algorithm has recovered the underlying labels: the *setosa* species is separated perfectly within cluster 0, while there remains a small amount of mixing between *versicolor* and *virginica*. This means that even without an expert to tell us the species labels of the individual flowers, the measurements of these flowers are distinct enough that we could *automatically* identify the presence of these different groups of species with a simple clustering algorithm! This sort of algorithm might further give experts in the field clues as to the relationships between the samples they are observing.

Application: Exploring Handwritten Digits

To demonstrate these principles on a more interesting problem, let's consider one piece of the optical character recognition problem: the identification of handwritten digits. In the wild, this problem involves both locating and identifying characters in an image. Here we'll take a shortcut and use Scikit-Learn's set of preformatted digits, which is built into the library.

Loading and Visualizing the Digits Data

We can use Scikit-Learn's data access interface to take a look at this data:

```
In [22]: from sklearn.datasets import load_digits  
        digits = load_digits()  
        digits.images.shape
```

Out[22]: (1797, 8, 8)

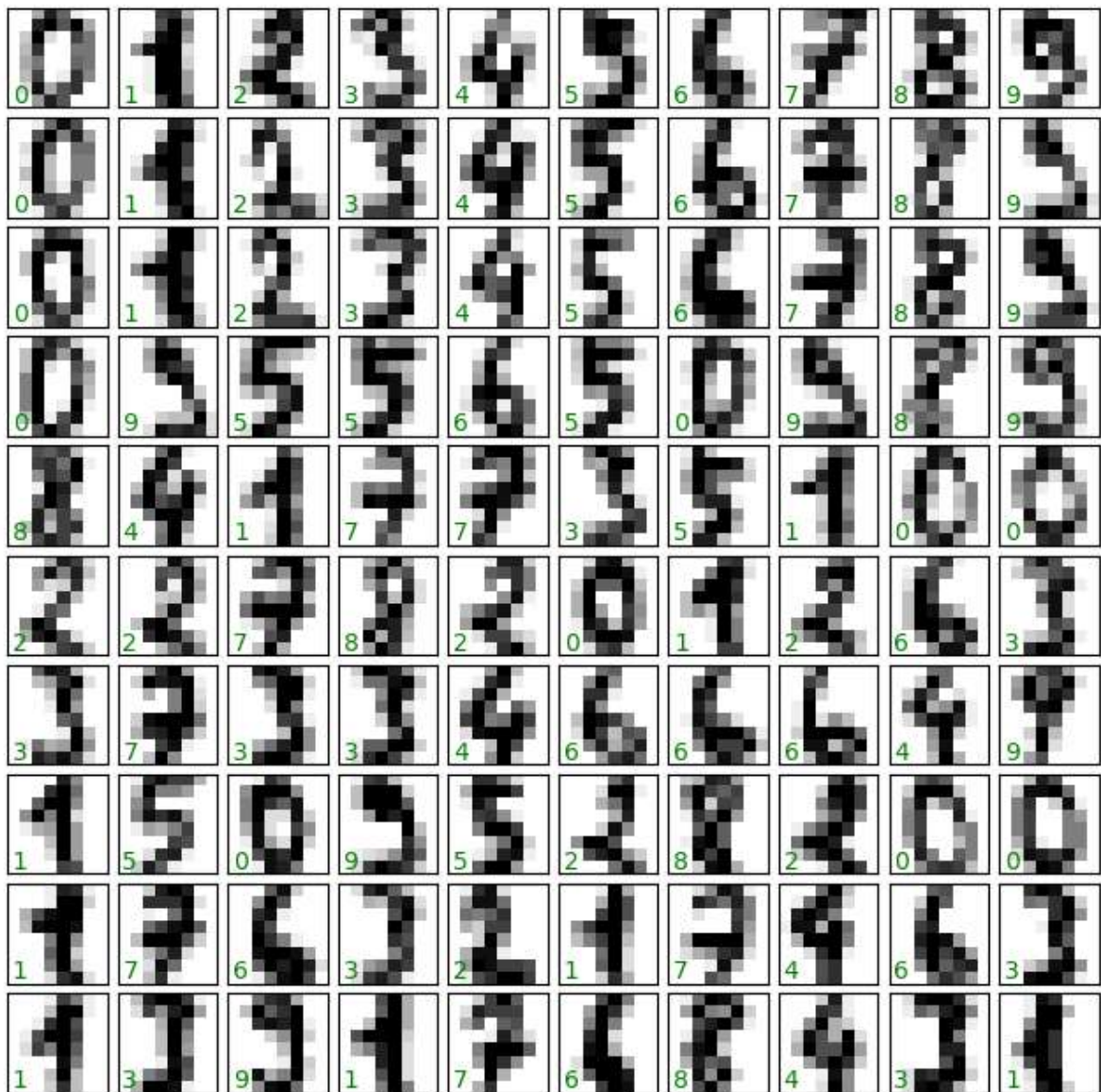
The images data is a three-dimensional array: 1,797 samples each consisting of an 8×8 grid of pixels. Let's visualize the first hundred of these (see the following figure):

```

gridspec_kw=dict(hspace=0.1, wspace=0.1))

for i, ax in enumerate(axes.flat):
    ax.imshow(digits.images[i], cmap='binary', interpolation='nearest')
    ax.text(0.05, 0.05, str(digits.target[i]),
            transform=ax.transAxes, color='green')

```



In order to work with this data within Scikit-Learn, we need a two-dimensional, [n_samples, n_features] representation. We can accomplish this by treating each pixel in the image as a feature: that is, by flattening out the pixel arrays so that we have a length-64 array of pixel values representing each digit. Additionally, we need the target array, which gives the previously determined label for each digit. These two quantities are built into the digits dataset under the `data` and `target` attributes, respectively:

In [24]:

```
X = digits.data
X.shape
```

Out[24]:

```
(1797, 64)
```

In [25]:

```
y = digits.target
y.shape
```

Out[25]:

```
(1797,)
```

We see here that there are 1,797 samples and 64 features.

Unsupervised Learning Example: Dimensionality Reduction

We'd like to visualize our points within the 64-dimensional parameter space, but it's difficult to effectively visualize points in such a high-dimensional space. Instead, we'll reduce the number of dimensions, using an unsupervised method. Here, we'll make use of a manifold learning algorithm called Isomap and transform the data to two dimensions:

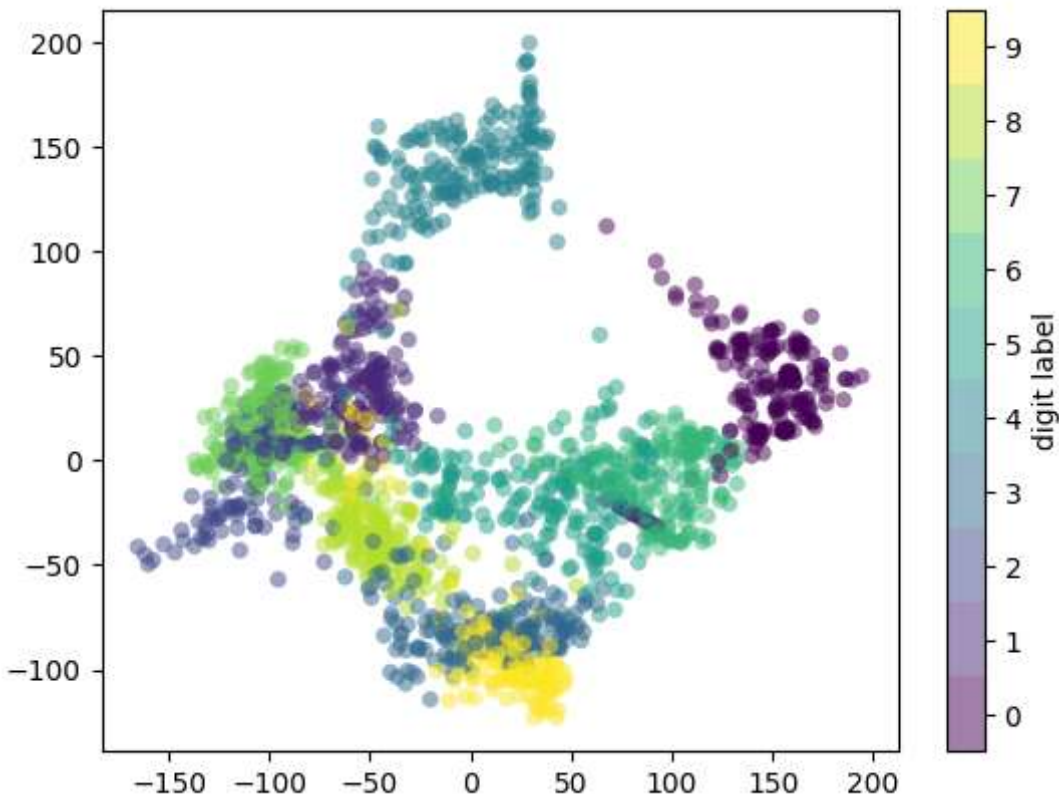
```
In [26]: from sklearn.manifold import Isomap
iso = Isomap(n_components=2)
iso.fit(digits.data)
data_projected = iso.transform(digits.data)
print(data_projected.shape)
```

```
C:\ProgramData\anaconda3\lib\site-packages\sklearn\manifold\_isomap.py:352: UserWarning: The number of connected components of the neighbors graph is 2 > 1. Completing the graph to fit Isomap might be slow. Increase the number of neighbors to avoid this issue.
    self._fit_transform(X)
C:\Users\NUWAN\AppData\Roaming\Python\Python310\site-packages\scipy\sparse\_index.py:103: SparseEfficiencyWarning: Changing the sparsity structure of a csr_matrix is expensive. lil_matrix is more efficient.
    self._set_intXint(row, col, x.flat[0])
(1797, 2)
```

We see that the projected data is now two-dimensional. Let's plot this data to see if we can learn anything from its structure (see the following figure):

```
In [27]: plt.scatter(data_projected[:, 0], data_projected[:, 1], c=digits.target,
                  edgecolor='none', alpha=0.5,
                  cmap=plt.cm.get_cmap('viridis', 10))
plt.colorbar(label='digit label', ticks=range(10))
plt.clim(-0.5, 9.5);
```

```
C:\Users\NUWAN\AppData\Local\Temp\ipykernel_996\3852404262.py:3: MatplotlibDeprecationWarning: The get_cmap function was deprecated in Matplotlib 3.7 and will be removed two minor releases later. Use ``matplotlib.colormaps[name]`` or ``matplotlib.colormaps.get_cmap(obj)`` instead.
    cmap=plt.cm.get_cmap('viridis', 10))
```



This plot gives us some good intuition into how well various numbers are separated in the larger 64-dimensional space. For example, zeros and ones have very little overlap in the parameter space. Intuitively, this makes sense: a zero is empty in the middle of the image, while a one will generally have ink in the middle. On the other hand, there seems to be a more or less continuous spectrum between ones and fours: we can understand this by realizing that some people draw ones with "hats" on them, which causes them to look similar to fours.

Overall, however, despite some mixing at the edges, the different groups appear to be fairly well localized in the parameter space: this suggests that even a very straightforward supervised classification algorithm should perform suitably on the full high-dimensional dataset. Let's give it a try.

Classification on Digits

Let's apply a classification algorithm to the digits data. As we did with the Iris data previously, we will split the data into training and testing sets and fit a Gaussian naive Bayes model:

```
In [28]: Xtrain, Xtest, ytrain, ytest = train_test_split(X, y, random_state=0)
```

```
In [29]: from sklearn.naive_bayes import GaussianNB
model = GaussianNB()
model.fit(Xtrain, ytrain)
y_model = model.predict(Xtest)
```

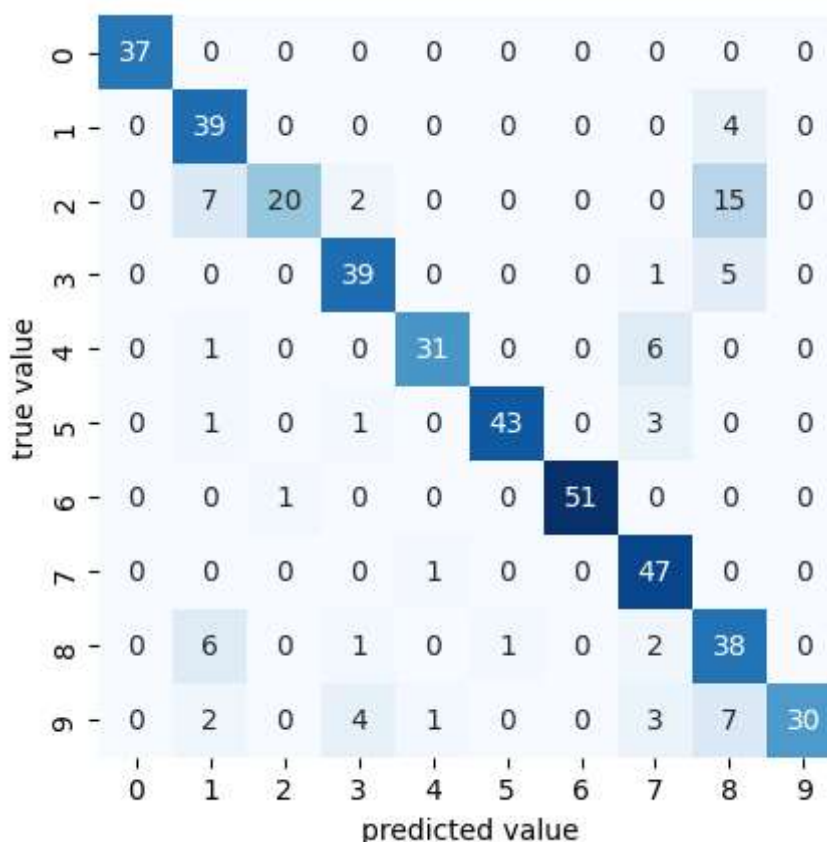
Now that we have the model's predictions, we can gauge its accuracy by comparing the true values of the test set to the predictions:

```
In [30]: from sklearn.metrics import accuracy_score  
accuracy_score(ytest, y_model)
```

```
Out[30]: 0.8333333333333334
```

With even this very simple model, we find about 83% accuracy for classification of the digits! However, this single number doesn't tell us where we've gone wrong. One nice way to do this is to use the *confusion matrix*, which we can compute with Scikit-Learn and plot with Seaborn (see the following figure):

```
In [31]: from sklearn.metrics import confusion_matrix  
  
mat = confusion_matrix(ytest, y_model)  
  
sns.heatmap(mat, square=True, annot=True, cbar=False, cmap='Blues')  
plt.xlabel('predicted value')  
plt.ylabel('true value');
```



This shows us where the mislabeled points tend to be: for example, many of the twos here are misclassified as either ones or eights.

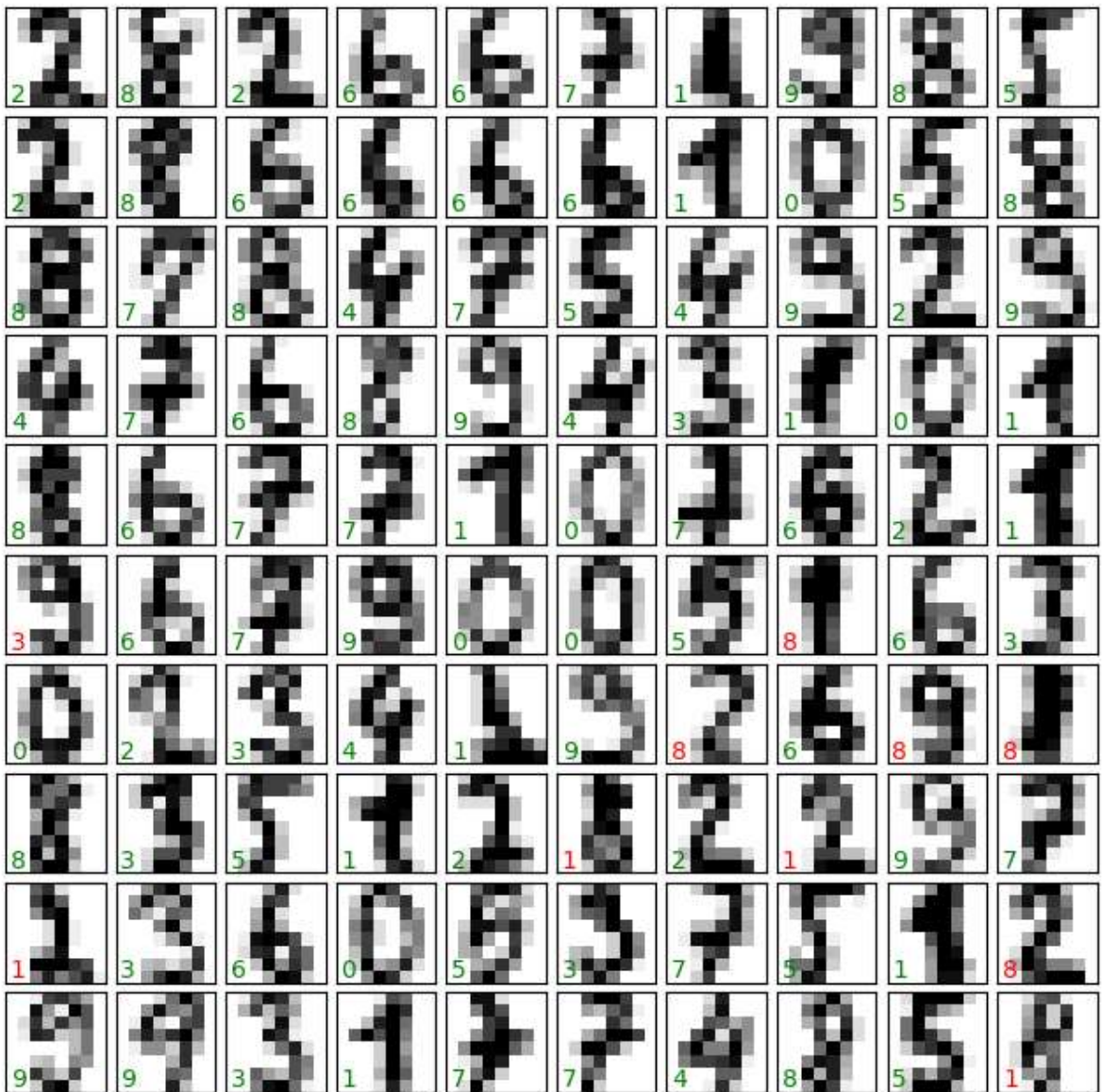
Another way to gain intuition into the characteristics of the model is to plot the inputs again, with their predicted labels. We'll use green for correct labels and red for incorrect labels; see the following figure:

```
In [32]: fig, axes = plt.subplots(10, 10, figsize=(8, 8),  
                           subplot_kw={'xticks':[], 'yticks':[]},  
                           gridspec_kw=dict(hspace=0.1, wspace=0.1))  
  
test_images = Xtest.reshape(-1, 8, 8)  
  
for i, ax in enumerate(axes.flat):
```

```

    ax.imshow(test_images[i], cmap='binary', interpolation='nearest')
    ax.text(0.05, 0.05, str(y_model[i])),
        transform=ax.transAxes,
        color='green' if (ytest[i] == y_model[i]) else 'red')

```



Examining this subset of the data can give us some insight into where the algorithm might be not performing optimally. To go beyond our 83% classification success rate, we might switch to a more sophisticated algorithm such as support vector machines, random forests, or another classification approach.

Summary

In this chapter we covered the essential features of the Scikit-Learn data representation and the Estimator API. Regardless of the type of estimator used, the same import/instantiate/fit/predict pattern holds. Armed with this information about the Estimator API, you can explore the Scikit-Learn documentation and begin trying out various models on your data.

In the next lab session, we will explore the most important topic in machine learning: how to select and validate your model.

