Introduction to Scikit-Learn (sklearn)

This notebook demonstrates some of the most useful functions of the beautiful Scikit-Learn library.

What we're going to cover:

```
In [1]: # Let's listify the contents
        what_were_covering = [
             "0. An end-to-end Scikit-Learn workflow",
            "1. Getting the data ready",
            "2. Choose the right estimator/algorithm for our problems",
            "3. Fit the model/algorithm and use it to make predictions on our data",
            "4. Evaluating a model",
            "5. Improve a model",
            "6. Save and load a trained model",
            "7. Putting it all together!"]
In [2]: what were covering
Out[2]: ['0. An end-to-end Scikit-Learn workflow',
          '1. Getting the data ready',
         '2. Choose the right estimator/algorithm for our problems',
         '3. Fit the model/algorithm and use it to make predictions on our data',
         '4. Evaluating a model',
         '5. Improve a model',
         '6. Save and load a trained model',
         '7. Putting it all together!']
In [3]: # Standard imports
        import numpy as np
        import pandas as pd
        import matplotlib.pyplot as plt
        %matplotlib inline
```

0. An end-to-end Scikit-Learn workflow

```
In [4]: # 1. Get the data ready
        import pandas as pd
        heart_disease = pd.read_csv("../data/heart-disease.csv")
        heart_disease
```

Out[4]:

		age	sex	ср	trestbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	са	thal	targe
	0	63	1	3	145	233	1	0	150	0	2.3	0	0	1	
	1	37	1	2	130	250	0	1	187	0	3.5	0	0	2	
	2	41	0	1	130	204	0	0	172	0	1.4	2	0	2	
	3	56	1	1	120	236	0	1	178	0	0.8	2	0	2	
	4	57	0	0	120	354	0	1	163	1	0.6	2	0	2	
2	98	57	0	0	140	241	0	1	123	1	0.2	1	0	3	
2	99	45	1	3	110	264	0	1	132	0	1.2	1	0	3	
3	00	68	1	0	144	193	1	1	141	0	3.4	1	2	3	
3	01	57	1	0	130	131	0	1	115	1	1.2	1	1	3	
3	02	57	0	1	130	236	0	0	174	0	0.0	1	1	2	

303 rows × 14 columns

```
In [5]: # Create X (features matrix)
        X = heart_disease.drop("target", axis=1)
        # Create y (labels)
        y = heart_disease["target"]
```

```
In [6]: # 2. Choose the right model and hyperparameters
        from sklearn.ensemble import RandomForestClassifier
        clf = RandomForestClassifier(n estimators=100)
        # We'll keep the default hyperparameters
        clf.get_params()
Out[6]: {'bootstrap': True,
          'ccp alpha': 0.0,
          'class_weight': None,
          'criterion': 'gini',
          'max depth': None,
          'max_features': 'auto',
          'max_leaf_nodes': None,
          'max samples': None,
          'min_impurity_decrease': 0.0,
          'min_impurity_split': None,
          'min_samples_leaf': 1,
          'min samples split': 2,
          'min weight fraction leaf': 0.0,
          'n_estimators': 100,
          'n_jobs': None,
          'oob_score': False,
          'random_state': None,
          'verbose': 0,
          'warm_start': False}
In [7]: # 3. Fit the model to the training data
        from sklearn.model_selection import train_test_split
        X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
In [8]: clf.fit(X_train, y_train);
```

In [9]: X_train

Out[9]:

	age	sex	ср	trestbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	са	thal
170	56	1	2	130	256	1	0	142	1	0.6	1	1	1
44	39	1	2	140	321	0	0	182	0	0.0	2	0	2
145	70	1	1	156	245	0	0	143	0	0.0	2	0	2
175	40	1	0	110	167	0	0	114	1	2.0	1	0	3
61	54	1	1	108	309	0	1	156	0	0.0	2	0	3
190	51	0	0	130	305	0	1	142	1	1.2	1	0	3
194	60	1	2	140	185	0	0	155	0	3.0	1	0	2
178	43	1	0	120	177	0	0	120	1	2.5	1	0	3
75	55	0	1	135	250	0	0	161	0	1.4	1	0	2
4	57	0	0	120	354	0	1	163	1	0.6	2	0	2

242 rows × 13 columns

```
In [10]: # make a prediction
y_label = clf.predict(np.array([0, 2, 3, 4]))
```

```
ValueError
                                          Traceback (most recent call last)
<ipython-input-10-7cea9660990e> in <module>
      1 # make a prediction
\rightarrow 2 y label = clf.predict(np.array([0, 2, 3, 4]))
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/ensemble/_forest.py in predict(self, X)
    610
                    The predicted classes.
    611
--> 612
                proba = self.predict proba(X)
    613
    614
                if self.n outputs == 1:
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/ensemble/ forest.py in predict proba(self, X)
    654
                check is fitted(self)
    655
                # Check data
--> 656
                X = self. validate X predict(X)
    657
    658
                # Assign chunk of trees to jobs
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/ensemble/ forest.py in validate X predict(self, X)
                check_is_fitted(self)
    410
    411
--> 412
                return self.estimators_[0]._validate_X_predict(X, check_input
=True)
    413
    414
            @property
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/tree/ classes.py in validate X predict(self, X, check input)
    378
                """Validate X whenever one tries to predict, apply, predict_p
roba"""
    379
                if check input:
                    X = check_array(X, dtype=DTYPE, accept_sparse="csr")
--> 380
    381
                    if issparse(X) and (X.indices.dtype != np.intc or
                                        X.indptr.dtype != np.intc):
    382
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/utils/validation.py in check array(array, accept sparse, accept large spars
e, dtype, order, copy, force_all_finite, ensure_2d, allow_nd, ensure_min_samp
les, ensure_min_features, warn_on_dtype, estimator)
    554
                            "Reshape your data either using array.reshape(-1,
1) if "
                            "your data has a single feature or array.reshape
    555
(1, -1) "
--> 556
                            "if it contains a single sample.".format(array))
    557
    558
                # in the future np.flexible dtypes will be handled like objec
t dtypes
ValueError: Expected 2D array, got 1D array instead:
array=[0. 2. 3. 4.].
Reshape your data either using array.reshape(-1, 1) if your data has a single
feature or array.reshape(1, -1) if it contains a single sample.
```

```
In [11]: | y_preds = clf.predict(X_test)
         y_preds
Out[11]: array([1, 0, 0, 0, 1, 0, 1, 0, 0, 0, 0, 0, 1, 1, 1, 1, 0, 0, 0, 1, 0,
                0, 1, 1, 1, 1, 0, 0, 1, 1, 0, 0, 0, 0, 1, 1, 0, 1, 1, 1, 1, 0, 0,
                1, 0, 1, 1, 1, 1, 0, 1, 0, 1, 0, 0, 1, 0, 0, 1, 1])
In [12]: y_test
Out[12]: 50
                1
         83
                1
         295
                0
         120
                1
         40
                1
         25
                1
         246
                0
         205
                0
         3
                1
         Name: target, Length: 61, dtype: int64
In [13]: # 4. Evaluate the model on the training data and test data
         clf.score(X_train, y_train)
Out[13]: 1.0
In [14]: | clf.score(X_test, y_test)
Out[14]: 0.7540983606557377
In [15]: from sklearn.metrics import classification report, confusion matrix, accuracy
          score
          print(classification_report(y_test, y_preds))
                        precision
                                     recall f1-score
                                                        support
                                       0.79
                     0
                             0.71
                                                 0.75
                                                              28
                     1
                             0.80
                                       0.73
                                                 0.76
                                                              33
                                                 0.75
                                                              61
             accuracy
                             0.75
                                       0.76
                                                 0.75
            macro avg
                                                              61
         weighted avg
                             0.76
                                       0.75
                                                 0.75
                                                              61
In [16]: confusion_matrix(y_test, y_preds)
Out[16]: array([[22, 6],
                [ 9, 24]])
In [17]: | accuracy_score(y_test, y_preds)
Out[17]: 0.7540983606557377
```

```
In [18]: # 5. Improve a model
         # Try different amount of n estimators
         np.random.seed(42)
         for i in range(10, 100, 10):
             print(f"Trying model with {i} estimators...")
             clf = RandomForestClassifier(n_estimators=i).fit(X_train, y_train)
             print(f"Model accuracy on test set: {clf.score(X test, y test) * 100:.2f}
         %")
             print("")
         Trying model with 10 estimators...
         Model accuracy on test set: 75.41%
         Trying model with 20 estimators...
         Model accuracy on test set: 78.69%
         Trying model with 30 estimators...
         Model accuracy on test set: 77.05%
         Trying model with 40 estimators...
         Model accuracy on test set: 80.33%
         Trying model with 50 estimators...
         Model accuracy on test set: 80.33%
         Trying model with 60 estimators...
         Model accuracy on test set: 80.33%
         Trying model with 70 estimators...
         Model accuracy on test set: 81.97%
         Trying model with 80 estimators...
         Model accuracy on test set: 78.69%
         Trying model with 90 estimators...
         Model accuracy on test set: 80.33%
In [19]:
         # 6. Save a model and load it
         import pickle
         pickle.dump(clf, open("random_forst_model_1.pkl", "wb"))
In [20]: loaded_model = pickle.load(open("random_forst_model_1.pkl", "rb"))
         loaded_model.score(X_test, y_test)
```

Out[20]: 0.8032786885245902

1. Getting our data ready to be used with machine learning

Three main things we have to do:

- Split the data into features and labels (usually `X` & `y`)
- 2. Filling (also called imputing) or disregarding missing values
- 3. Converting non-numerical values to numerical values (also called feature encodin g)

```
In [21]: heart disease.head()
```

Out[21]:

	age	sex	ср	trestbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	са	thal	target
0	63	1	3	145	233	1	0	150	0	2.3	0	0	1	1
1	37	1	2	130	250	0	1	187	0	3.5	0	0	2	1
2	41	0	1	130	204	0	0	172	0	1.4	2	0	2	1
3	56	1	1	120	236	0	1	178	0	0.8	2	0	2	1
4	57	0	0	120	354	0	1	163	1	0.6	2	0	2	1

```
In [22]: X = heart_disease.drop("target", axis=1)
         X.head()
```

Out[22]:

	age	sex	ср	trestbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	са	thal
0	63	1	3	145	233	1	0	150	0	2.3	0	0	1
1	37	1	2	130	250	0	1	187	0	3.5	0	0	2
2	41	0	1	130	204	0	0	172	0	1.4	2	0	2
3	56	1	1	120	236	0	1	178	0	0.8	2	0	2
4	57	0	0	120	354	0	1	163	1	0.6	2	0	2

```
In [23]: y = heart_disease["target"]
         y.head()
```

```
Out[23]: 0
```

- 1
- 1
- 1
- 3 1

Name: target, dtype: int64

```
In [24]: # Split the data into training and test sets
         from sklearn.model selection import train test split
         X_train, X_test, y_train, y_test = train_test_split(X,
                                                              у,
                                                              test_size=0.3)
In [25]: X_train.shape, X_test.shape, y_train.shape, y_test.shape
Out[25]: ((212, 13), (91, 13), (212,), (91,))
In [26]: X.shape[0] * 0.8
Out[26]: 242.4
In [27]: 242 + 61
Out[27]: 303
In [28]: len(heart_disease)
Out[28]: 303
```

1.1 Make sure it's all numerical

```
car sales = pd.read csv("../data/car-sales-extended.csv")
In [29]:
          car sales.head()
Out[29]:
              Make Colour Odometer (KM) Doors
                                                Price
           0 Honda
                     White
                                   35431
                                               15323
              BMW
                      Blue
                                  192714
                                             5 19943
           1
           2 Honda
                     White
                                   84714
                                               28343
             Toyota
                     White
                                  154365
                                               13434
                                             3 14043
           4 Nissan
                      Blue
                                  181577
In [30]: car_sales["Doors"].value_counts()
Out[30]: 4
               856
                79
                65
          Name: Doors, dtype: int64
In [31]: len(car_sales)
Out[31]: 1000
```

```
In [32]: car_sales.dtypes
Out[32]: Make
                           object
         Colour
                           object
         Odometer (KM)
                            int64
                            int64
         Doors
         Price
                            int64
         dtype: object
In [33]: # Split into X/y
         X = car_sales.drop("Price", axis=1)
         y = car_sales["Price"]
         # Split into training and test
         X_train, X_test, y_train, y_test = train_test_split(X,
                                                             test_size=0.2)
```

```
# Build machine learning model
from sklearn.ensemble import RandomForestRegressor
model = RandomForestRegressor()
model.fit(X train, y train)
model.score(X_test, y_test)
ValueError
                                           Traceback (most recent call last)
<ipython-input-34-2eeea2d0b490> in <module>
      3
      4 model = RandomForestRegressor()
----> 5 model.fit(X_train, y_train)
      6 model.score(X_test, y_test)
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/ensemble/_forest.py in fit(self, X, y, sample_weight)
    293
    294
                # Validate or convert input data
                X = check array(X, accept sparse="csc", dtype=DTYPE)
--> 295
                y = check_array(y, accept_sparse='csc', ensure_2d=False, dtyp
    296
e=None)
                if sample weight is not None:
    297
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/utils/validation.py in check array(array, accept sparse, accept large spars
e, dtype, order, copy, force_all_finite, ensure_2d, allow_nd, ensure_min_samp
les, ensure min features, warn on dtype, estimator)
                            array = array.astype(dtype, casting="unsafe", cop
    529
y=False)
    530
                        else:
                             array = np.asarray(array, order=order, dtype=dtyp
--> 531
e)
    532
                    except ComplexWarning:
    533
                        raise ValueError("Complex data not supported\n"
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/numpy/
core/ asarray.py in asarray(a, dtype, order)
     83
     84
---> 85
            return array(a, dtype, copy=False, order=order)
     86
     87
```

ValueError: could not convert string to float: 'Toyota'

```
In [35]: X.head()
```

Out[35]:

```
Make Colour Odometer (KM) Doors
  Honda
           White
                          35431
                                      4
   BMW
            Blue
                         192714
                                     5
2 Honda
           White
                          84714
                                      4
  Toyota
           White
                         154365
                                      4
4 Nissan
            Blue
                         181577
                                     3
```

```
In [36]:
         # Turn the categories into numbers
         from sklearn.preprocessing import OneHotEncoder
         from sklearn.compose import ColumnTransformer
         categorical_features = ["Make", "Colour", "Doors"]
         one_hot = OneHotEncoder()
         transformer = ColumnTransformer([("one_hot",
                                             one hot,
                                             categorical_features)],
                                             remainder="passthrough")
         transformed_X = transformer.fit_transform(X)
         transformed X
```

```
Out[36]: array([[0.00000e+00, 1.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 3.54310e+04],
                [1.00000e+00, 0.00000e+00, 0.00000e+00, ..., 0.00000e+00,
                 1.00000e+00, 1.92714e+05],
                 [0.00000e+00, 1.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 8.47140e+04],
                [0.00000e+00, 0.00000e+00, 1.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 6.66040e+04],
                 [0.00000e+00, 1.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 2.15883e+05],
                 [0.00000e+00, 0.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 2.48360e+05]])
```

In [37]: X.head()

Out[37]:

	Make	Colour	Odometer (KM)	Doors
0	Honda	White	35431	4
1	BMW	Blue	192714	5
2	Honda	White	84714	4
3	Toyota	White	154365	4
4	Nissan	Blue	181577	3

In [38]: pd.DataFrame(transformed_X)

Out[38]:

	0	1	2	3	4	5	6	7	8	9	10	11	12
0	0.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	1.0	0.0	1.0	0.0	35431.0
1	1.0	0.0	0.0	0.0	0.0	1.0	0.0	0.0	0.0	0.0	0.0	1.0	192714.0
2	0.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	1.0	0.0	1.0	0.0	84714.0
3	0.0	0.0	0.0	1.0	0.0	0.0	0.0	0.0	1.0	0.0	1.0	0.0	154365.0
4	0.0	0.0	1.0	0.0	0.0	1.0	0.0	0.0	0.0	1.0	0.0	0.0	181577.0
995	0.0	0.0	0.0	1.0	1.0	0.0	0.0	0.0	0.0	0.0	1.0	0.0	35820.0
996	0.0	0.0	1.0	0.0	0.0	0.0	0.0	0.0	1.0	1.0	0.0	0.0	155144.0
997	0.0	0.0	1.0	0.0	0.0	1.0	0.0	0.0	0.0	0.0	1.0	0.0	66604.0
998	0.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	1.0	0.0	1.0	0.0	215883.0
999	0.0	0.0	0.0	1.0	0.0	1.0	0.0	0.0	0.0	0.0	1.0	0.0	248360.0

1000 rows × 13 columns

In [39]: # Another way to do it with pd.dummies... dummies = pd.get_dummies(car_sales[["Make", "Colour", "Doors"]]) dummies

Out[39]:

	Doors	Make_BMW	Make_Honda	Make_Nissan	Make_Toyota	Colour_Black	Colour_Blue
0	4	0	1	0	0	0	0
1	5	1	0	0	0	0	1
2	4	0	1	0	0	0	0
3	4	0	0	0	1	0	0
4	3	0	0	1	0	0	1
995	4	0	0	0	1	1	0
996	3	0	0	1	0	0	0
997	4	0	0	1	0	0	1
998	4	0	1	0	0	0	0
999	4	0	0	0	1	0	1

1000 rows × 10 columns

```
In [40]: # Let's refit the model
          np.random.seed(42)
          X_train, X_test, y_train, y_test = train_test_split(transformed_X,
                                                                test size=0.2)
         model.fit(X_train, y_train)
Out[40]: RandomForestRegressor(bootstrap=True, ccp_alpha=0.0, criterion='mse',
                                max depth=None, max features='auto', max leaf nodes=Non
         e,
                                max samples=None, min impurity decrease=0.0,
                                min impurity split=None, min samples leaf=1,
                                min_samples_split=2, min_weight_fraction_leaf=0.0,
                                n_estimators=100, n_jobs=None, oob_score=False,
                                 random state=None, verbose=0, warm start=False)
In [41]: X.head()
Out[41]:
              Make Colour Odometer (KM) Doors
            Honda
                    White
                                  35431
                                            4
              BMW
                     Blue
                                 192714
                                            5
            Honda
                     White
                                 84714
             Toyota
                     White
                                 154365
            Nissan
                     Blue
                                 181577
                                            3
In [42]: model.score(X_test, y_test)
```

1.2 What if there were missing values?

Out[42]: 0.3235867221569877

- 1. Fill them with some value (also known as imputation).
- 2. Remove the samples with missing data altogether.

```
In [43]: # Import car sales missing data
         car_sales_missing = pd.read_csv("../data/car-sales-extended-missing-data.csv")
         car_sales_missing.head()
```

Out[43]:

	Make	Colour	Odometer (KM)	Doors	Price
0	Honda	White	35431.0	4.0	15323.0
1	BMW	Blue	192714.0	5.0	19943.0
2	Honda	White	84714.0	4.0	28343.0
3	Toyota	White	154365.0	4.0	13434.0
4	Nissan	Blue	181577.0	3.0	14043.0

```
In [44]: car_sales_missing.isna().sum()
Out[44]: Make
                           49
         Colour
                           50
         Odometer (KM)
                           50
         Doors
                           50
         Price
                           50
         dtype: int64
```

```
In [45]: # Create X & y
         X = car_sales_missing.drop("Price", axis=1)
         y = car_sales_missing["Price"]
```

```
In [46]: # Let's try and convert our data to numbers
         # Turn the categories into numbers
         from sklearn.preprocessing import OneHotEncoder
         from sklearn.compose import ColumnTransformer
         categorical_features = ["Make", "Colour", "Doors"]
         one_hot = OneHotEncoder()
         transformer = ColumnTransformer([("one_hot",
                                             one_hot,
                                             categorical_features)],
                                             remainder="passthrough")
         transformed_X = transformer.fit_transform(X)
         transformed X
```

```
Traceback (most recent call last)
ValueError
<ipython-input-46-f532939289ac> in <module>
     11
                                            remainder="passthrough")
     12
---> 13 transformed X = transformer.fit transform(X)
     14 transformed X
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/compose/ column transformer.py in fit transform(self, X, y)
                self. validate remainder(X)
    516
    517
--> 518
                result = self. fit transform(X, y, fit transform one)
    519
    520
                if not result:
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/compose/_column_transformer.py in _fit_transform(self, X, y, func, fitted)
    455
                            message=self. log message(name, idx, len(transfor
mers)))
                        for idx, (name, trans, column, weight) in enumerate(
    456
                                self. iter(fitted=fitted, replace strings=Tru
--> 457
e), 1))
    458
                except ValueError as e:
                    if "Expected 2D array, got 1D array instead" in str(e):
    459
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/jobli
b/parallel.py in __call__(self, iterable)
   1002
                    # remaining jobs.
   1003
                    self. iterating = False
                    if self.dispatch_one_batch(iterator):
-> 1004
                        self. iterating = self. original iterator is not None
   1005
   1006
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/jobli
b/parallel.py in dispatch one batch(self, iterator)
                        return False
    833
    834
                    else:
                        self. dispatch(tasks)
--> 835
    836
                        return True
    837
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/jobli
b/parallel.py in _dispatch(self, batch)
                with self. lock:
    752
   753
                    job idx = len(self. jobs)
--> 754
                    job = self._backend.apply_async(batch, callback=cb)
                    # A job can complete so quickly than its callback is
    755
                    # called before we get here, causing self._jobs to
    756
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/jobli
b/ parallel backends.py in apply async(self, func, callback)
            def apply async(self, func, callback=None):
    207
    208
                """Schedule a func to be run"""
                result = ImmediateResult(func)
--> 209
    210
                if callback:
    211
                    callback(result)
```

```
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/jobli
b/_parallel_backends.py in __init__(self, batch)
    588
                # Don't delay the application, to avoid keeping the input
    589
                # arguments in memory
--> 590
                self.results = batch()
    591
            def get(self):
    592
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/jobli
b/parallel.py in call (self)
    254
                with parallel_backend(self._backend, n_jobs=self._n_jobs):
    255
                    return [func(*args, **kwargs)
--> 256
                            for func, args, kwargs in self.items]
    257
            def __len__(self):
    258
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/jobli
b/parallel.py in <listcomp>(.0)
    254
                with parallel_backend(self._backend, n_jobs=self._n_jobs):
    255
                    return [func(*args, **kwargs)
--> 256
                            for func, args, kwargs in self.items]
    257
    258
            def len (self):
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/pipeline.py in _fit_transform_one(transformer, X, y, weight, message_clsnam
e, message, **fit params)
    726
            with print elapsed time(message clsname, message):
    727
                if hasattr(transformer, 'fit transform'):
                    res = transformer.fit_transform(X, y, **fit_params)
--> 728
    729
                else:
                    res = transformer.fit(X, y, **fit_params).transform(X)
    730
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/preprocessing/ encoders.py in fit transform(self, X, y)
    370
    371
                self. validate keywords()
                return super().fit transform(X, y)
--> 372
    373
    374
            def transform(self, X):
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/base.py in fit transform(self, X, y, **fit params)
    569
                if y is None:
    570
                    # fit method of arity 1 (unsupervised transformation)
                    return self.fit(X, **fit_params).transform(X)
--> 571
    572
                else:
                    # fit method of arity 2 (supervised transformation)
    573
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/preprocessing/_encoders.py in fit(self, X, y)
    345
    346
                self. validate keywords()
                self. fit(X, handle unknown=self.handle unknown)
--> 347
                self.drop_idx_ = self._compute_drop_idx()
    348
    349
                return self
```

```
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/preprocessing/ encoders.py in fit(self, X, handle unknown)
     72
     73
            def fit(self, X, handle unknown='error'):
---> 74
                X_list, n_samples, n_features = self._check_X(X)
     75
     76
                if self.categories != 'auto':
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/preprocessing/ encoders.py in check X(self, X)
     59
                    Xi = self._get_feature(X, feature_idx=i)
     60
                    Xi = check array(Xi, ensure 2d=False, dtype=None,
                                     force all finite=needs validation)
---> 61
     62
                    X columns.append(Xi)
     63
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/utils/validation.py in check array(array, accept sparse, accept large spars
e, dtype, order, copy, force all finite, ensure 2d, allow nd, ensure min samp
les, ensure_min_features, warn_on_dtype, estimator)
                if force all finite:
    576
    577
                    assert all finite(array,
--> 578
                                       allow nan=force all finite == 'allow-n
an')
    579
    580
            if ensure min samples > 0:
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/utils/validation.py in assert all finite(X, allow nan, msg dtype)
            elif X.dtype == np.dtype('object') and not allow_nan:
     63
     64
                if object dtype isnan(X).any():
                    raise ValueError("Input contains NaN")
---> 65
     66
     67
```

ValueError: Input contains NaN

Drico

```
car_sales_missing
```

Out[47]:

		маке	Colour	Odometer (KM)	Doors	Price
	0	Honda	White	35431.0	4.0	15323.0
	1	BMW	Blue	192714.0	5.0	19943.0
	2	Honda	White	84714.0	4.0	28343.0
	3	Toyota	White	154365.0	4.0	13434.0
	4	Nissan	Blue	181577.0	3.0	14043.0
9	95	Toyota	Black	35820.0	4.0	32042.0
9	96	NaN	White	155144.0	3.0	5716.0
9	97	Nissan	Blue	66604.0	4.0	31570.0
9	98	Honda	White	215883.0	4.0	4001.0
9	99	Toyota	Blue	248360.0	4.0	12732.0

Colour Odometer (KM) Doors

1000 rows × 5 columns

```
In [48]: | car_sales_missing["Doors"].value_counts()
Out[48]: 4.0
                 811
                  75
          5.0
          3.0
                  64
         Name: Doors, dtype: int64
```

Option 1: Fill missing data with Pandas

```
# Fill the "Make" column
In [49]:
         car_sales_missing["Make"].fillna("missing", inplace=True)
         # Fill the "Colour" column
         car_sales_missing["Colour"].fillna("missing", inplace=True)
         # Fill the "Odometer (KM)" column
         car_sales_missing["Odometer (KM)"].fillna(car_sales_missing["Odometer (KM)"].m
         ean(), inplace=True)
         # Fill the "Doors" column
         car_sales_missing["Doors"].fillna(4, inplace=True)
```

```
In [50]: # Check our dataframe again
         car_sales_missing.isna().sum()
Out[50]: Make
                            0
         Colour
                            0
         Odometer (KM)
                            0
         Doors
                            0
         Price
                           50
         dtype: int64
In [51]: # Remove rows with missing Price value
         car_sales_missing.dropna(inplace=True)
In [52]: car_sales_missing.isna().sum()
Out[52]: Make
         Colour
                           0
         Odometer (KM)
                          0
         Doors
                           0
         Price
                          0
         dtype: int64
In [53]: len(car_sales_missing)
Out[53]: 950
In [54]: X = car_sales_missing.drop("Price", axis=1)
         y = car_sales_missing["Price"]
```

```
In [55]: # Let's try and convert our data to numbers
         # Turn the categories into numbers
         from sklearn.preprocessing import OneHotEncoder
         from sklearn.compose import ColumnTransformer
         categorical_features = ["Make", "Colour", "Doors"]
         one hot = OneHotEncoder()
         transformer = ColumnTransformer([("one hot",
                                             one hot,
                                             categorical_features)],
                                             remainder="passthrough")
         transformed_X = transformer.fit_transform(car_sales_missing)
         transformed X
Out[55]: array([[0.00000e+00, 1.00000e+00, 0.00000e+00, ..., 0.00000e+00,
                 3.54310e+04, 1.53230e+04],
                [1.00000e+00, 0.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 1.92714e+05, 1.99430e+04],
                [0.00000e+00, 1.00000e+00, 0.00000e+00, ..., 0.00000e+00,
                 8.47140e+04, 2.83430e+04],
                [0.00000e+00, 0.00000e+00, 1.00000e+00, ..., 0.00000e+00,
                 6.66040e+04, 3.15700e+04],
                [0.00000e+00, 1.00000e+00, 0.00000e+00, ..., 0.00000e+00,
                 2.15883e+05, 4.00100e+03],
                [0.00000e+00, 0.00000e+00, 0.00000e+00, ..., 0.00000e+00,
                 2.48360e+05, 1.27320e+04]])
```

Option 2: Filling missing data and transforming categorical data with Scikit-Learn

Note: This section is different to the video. The video shows filling and transforming the entire dataset (X) and although the techniques are correct, it's best to fill and transform training and test sets separately (as shown in the code below).

The main takeaways:

- Split your data first (into train/test)
- Fill/transform the training set and test sets separately

Thank you Robert for pointing this out (https://www.udemy.com/course/complete-machine-learning-and-datascience-zero-to-mastery/learn/#questions/9506426).

```
introduction-to-scikit-learn-video
In [56]: car sales missing = pd.read csv("../data/car-sales-extended-missing-data.csv")
          car sales missing.head()
Out[56]:
              Make Colour Odometer (KM) Doors
                                                   Price
           0 Honda
                      White
                                  35431.0
                                             4.0 15323.0
              BMW
                       Blue
                                 192714.0
                                             5.0 19943.0
                      White
           2 Honda
                                  84714.0
                                             4.0 28343.0
             Toyota
                      White
                                 154365.0
                                             4.0 13434.0
           4 Nissan
                       Blue
                                 181577.0
                                             3.0 14043.0
In [57]:
          car_sales_missing.isna().sum()
Out[57]: Make
                             49
          Colour
                             50
          Odometer (KM)
                             50
          Doors
                             50
          Price
                             50
          dtype: int64
In [58]: # Drop the rows with no labels
          car_sales_missing.dropna(subset=["Price"], inplace=True)
          car_sales_missing.isna().sum()
Out[58]: Make
                             47
```

```
Colour
                  46
Odometer (KM)
                  48
Doors
                  47
Price
                   0
dtype: int64
```

```
In [59]: # Split into X & y
         X = car_sales_missing.drop("Price", axis=1)
         y = car_sales_missing["Price"]
         # Split data into train and test
         np.random.seed(42)
         X_train, X_test, y_train, y_test = train_test_split(X,
                                                               у,
                                                               test_size=0.2)
```

```
In [60]: # Check missing values
         X.isna().sum()
```

```
Out[60]: Make
                            47
          Colour
                            46
          Odometer (KM)
                            48
                            47
          Doors
          dtype: int64
```

```
In [61]: # Fill missing values with Scikit-Learn
         from sklearn.impute import SimpleImputer
         from sklearn.compose import ColumnTransformer
         # Fill categorical values with 'missing' & numerical values with mean
         cat_imputer = SimpleImputer(strategy="constant", fill_value="missing")
         door imputer = SimpleImputer(strategy="constant", fill value=4)
         num imputer = SimpleImputer(strategy="mean")
         # Define columns
         cat_features = ["Make", "Colour"]
         door feature = ["Doors"]
         num_features = ["Odometer (KM)"]
         # Create an imputer (something that fills missing data)
         imputer = ColumnTransformer([
             ("cat_imputer", cat_imputer, cat_features),
             ("door_imputer", door_imputer, door_feature),
             ("num_imputer", num_imputer, num_features)
         1)
         # Fill train and test values separately
         filled X train = imputer.fit transform(X train)
         filled X test = imputer.transform(X test)
         # Check filled X train
         filled X train
Out[61]: array([['Honda', 'White', 4.0, 71934.0],
                ['Toyota', 'Red', 4.0, 162665.0].
                ['Honda', 'White', 4.0, 42844.0],
                ['Toyota', 'White', 4.0, 196225.0],
                ['Honda', 'Blue', 4.0, 133117.0],
                ['Honda', 'missing', 4.0, 150582.0]], dtype=object)
In [62]: # Get our transformed data array's back into DataFrame's
         car_sales_filled_train = pd.DataFrame(filled_X_train,
                                                columns=["Make", "Colour", "Doors", "Odo
         meter (KM)"])
         car sales filled test = pd.DataFrame(filled X test,
                                               columns=["Make", "Colour", "Doors", "Odom
         eter (KM)"])
         # Check missing data in training set
         car_sales_filled_train.isna().sum()
Out[62]: Make
         Colour
                          0
         Doors
                          0
         Odometer (KM)
                          0
         dtype: int64
```

```
In [63]: # Check to see the original... still missing values
         car sales missing.isna().sum()
Out[63]: Make
                          47
         Colour
                          46
         Odometer (KM)
                          48
         Doors
                          47
         Price
                           0
         dtype: int64
In [64]:
         # Now let's one hot encode the features with the same code as before
         categorical_features = ["Make", "Colour", "Doors"]
         one hot = OneHotEncoder()
         transformer = ColumnTransformer([("one_hot",
                                           one hot,
                                           categorical features)],
                                           remainder="passthrough")
         # Fill train and test values separately
         transformed X train = transformer.fit transform(car sales filled train)
         transformed_X_test = transformer.transform(car_sales_filled_test)
         # Check transformed and filled X train
         transformed_X_train.toarray()
Out[64]: array([[0.00000e+00, 1.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 7.19340e+04],
                [0.00000e+00, 0.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 1.62665e+05],
                [0.00000e+00, 1.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 4.28440e+04],
                [0.00000e+00, 0.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 1.96225e+05],
                [0.00000e+00, 1.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 1.33117e+05],
                [0.00000e+00, 1.00000e+00, 0.00000e+00, ..., 1.00000e+00,
                 0.00000e+00, 1.50582e+05]])
In [65]: # Now we've transformed X, let's see if we can fit a model
         np.random.seed(42)
         from sklearn.ensemble import RandomForestRegressor
         model = RandomForestRegressor()
         # Make sure to use transformed (filled and one-hot encoded X data)
         model.fit(transformed_X_train, y_train)
         model.score(transformed X test, y test)
Out[65]: 0.21229043336119102
```

```
In [66]: # Check Length of transformed data (filled and one-hot encoded)
         # vs. length of original data
         len(transformed_X_train.toarray())+len(transformed_X_test.toarray()), len(car_
         sales)
Out[66]: (950, 1000)
```

Note: The 50 less values in the transformed data is because we dropped the rows (50 total) with missing values in the Price column.

2. Choosing the right estimator/algorithm for our problem

Scikit-Learn uses estimator as another term for machine learning model or algorithm.

- · Classification predicting whether a sample is one thing or another
- · Regression predicting a number

Step 1 - Check the Scikit-Learn machine learning map... https://scikitlearn.org/stable/tutorial/machine learning_map/index.html (https://scikitlearn.org/stable/tutorial/machine learning map/index.html)

2.1 Picking a machine learning model for a regression problem

```
In [67]: # Import Boston housing dataset
          from sklearn.datasets import load boston
          boston = load boston()
          boston;
          boston_df = pd.DataFrame(boston["data"], columns=boston["feature_names"])
In [68]:
          boston df["target"] = pd.Series(boston["target"])
          boston_df.head()
Out[68]:
                CRIM
                       ZN INDUS CHAS
                                          NOX
                                                 RM
                                                      AGE
                                                              DIS RAD
                                                                         TAX PTRATIO
                                                                                            B LST.
           0 0.00632 18.0
                             2.31
                                     0.0 0.538 6.575
                                                      65.2 4.0900
                                                                        296.0
                                                                                  15.3 396.90
                                                                    1.0
                                                                                                 4.
             0.02731
                             7.07
                                     0.0 0.469
                                               6.421
                                                           4.9671
                                                                                       396.90
                       0.0
                                                      78.9
                                                                    2.0
                                                                        242.0
                                                                                  17.8
                                                                                                 9.
           2 0.02729
                             7.07
                       0.0
                                     0.0 0.469 7.185
                                                      61.1 4.9671
                                                                    2.0 242.0
                                                                                  17.8 392.83
                                                                                                 4.
             0.03237
                       0.0
                             2.18
                                     0.0 0.458
                                               6.998
                                                      45.8
                                                           6.0622
                                                                        222.0
                                                                                  18.7 394.63
                                     0.0 0.458 7.147
             0.06905
                       0.0
                             2.18
                                                      54.2 6.0622
                                                                    3.0 222.0
                                                                                  18.7
                                                                                       396.90
                                                                                                 5.
          # How many samples?
In [69]:
          len(boston df)
Out[69]: 506
```

```
In [70]: # Let's try the Ridge Regression model
         from sklearn.linear model import Ridge
         # Setup random seed
         np.random.seed(42)
         # Create the data
         X = boston df.drop("target", axis=1)
         y = boston_df["target"]
         # Split into train and test sets
         X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
         # Instantiate Ridge model
         model = Ridge()
         model.fit(X_train, y_train)
         # Check the score of the Ridge model on test data
         model.score(X_test, y_test)
```

Out[70]: 0.6662221670168518

How do we improve this score?

What if Ridge wasn't working?

Let's refer back to the map... https://scikit-learn.org/stable/tutorial/machine_learning_map/index.html (https://scikit-learn.org/stable/tutorial/machine_learning_map/index.html)

```
In [71]: # Let's try the Random Forst Regressor
         from sklearn.ensemble import RandomForestRegressor
         # Setup random seed
         np.random.seed(42)
         # Create the data
         X = boston_df.drop("target", axis=1)
         y = boston_df["target"]
         # Split the data
         #X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
         # Instatiate Random Forest Regressor
         rf = RandomForestRegressor(n_estimators=100)
         rf.fit(X train, y train)
         # Evaluate the Random Forest Regressor
         rf.score(X_test, y_test)
```

Out[71]: 0.8896648705127477

```
In [72]: # Check the Ridge model again
         model.score(X_test, y_test)
```

Out[72]: 0.6662221670168518

2.2 Choosing an estimator for a classification problem

Let's go to the map... https://scikit-learn.org/stable/tutorial/machine_learning_map/index.html (https://scikit-<u>learn.org/stable/tutorial/machine_learning_map/index.html)</u>

```
In [73]: heart_disease = pd.read_csv("data/heart-disease.csv")
heart_disease.head()
```

```
Traceback (most recent call last)
FileNotFoundError
<ipython-input-73-44f78f3704d4> in <module>
----> 1 heart disease = pd.read csv("data/heart-disease.csv")
      2 heart disease.head()
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/panda
s/io/parsers.py in parser_f(filepath_or_buffer, sep, delimiter, header, name
s, index_col, usecols, squeeze, prefix, mangle_dupe_cols, dtype, engine, conv
erters, true values, false values, skipinitialspace, skiprows, skipfooter, nr
ows, na values, keep default na, na filter, verbose, skip blank lines, parse
dates, infer_datetime_format, keep_date_col, date_parser, dayfirst, cache_dat
es, iterator, chunksize, compression, thousands, decimal, lineterminator, quo
techar, quoting, doublequote, escapechar, comment, encoding, dialect, error_b
ad lines, warn bad lines, delim whitespace, low memory, memory map, float pre
cision)
    674
                )
    675
--> 676
                return read(filepath or buffer, kwds)
    677
    678
            parser_f.__name__ = name
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/panda
s/io/parsers.py in read(filepath or buffer, kwds)
    446
    447
            # Create the parser.
--> 448
            parser = TextFileReader(fp or buf, **kwds)
    449
    450
            if chunksize or iterator:
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/panda
s/io/parsers.py in init (self, f, engine, **kwds)
    878
                    self.options["has_index_names"] = kwds["has_index_names"]
    879
--> 880
                self. make engine(self.engine)
    881
            def close(self):
    882
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/panda
s/io/parsers.py in make engine(self, engine)
            def make engine(self, engine="c"):
   1112
   1113
                if engine == "c":
                    self._engine = CParserWrapper(self.f, **self.options)
-> 1114
   1115
                else:
                    if engine == "python":
   1116
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/panda
s/io/parsers.py in init (self, src, **kwds)
   1889
                kwds["usecols"] = self.usecols
   1890
-> 1891
                self. reader = parsers.TextReader(src, **kwds)
                self.unnamed cols = self. reader.unnamed cols
   1892
   1893
pandas/ libs/parsers.pyx in pandas. libs.parsers.TextReader. cinit ()
pandas/ libs/parsers.pyx in pandas. libs.parsers.TextReader. setup parser sou
```

```
rce()
```

FileNotFoundError: [Errno 2] File data/heart-disease.csv does not exist: 'dat a/heart-disease.csv'

```
In [74]: len(heart disease)
Out[74]: 303
```

Consulting the map and it says to try LinearSVC.

```
In [75]: # Import the LinearSVC estimator class
         from sklearn.svm import LinearSVC
         # Setup random seed
         np.random.seed(42)
         # Make the data
         X = heart_disease.drop("target", axis=1)
         y = heart_disease["target"]
         # Split the data
         X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
         # Instantiate LinearSVC
         clf = LinearSVC(max iter=10000)
         clf.fit(X_train, y_train)
         # Evaluate the LinearSVC
         clf.score(X_test, y_test)
```

/Users/daniel/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-pac kages/sklearn/svm/ base.py:947: ConvergenceWarning: Liblinear failed to conve rge, increase the number of iterations.

"the number of iterations.", ConvergenceWarning)

```
Out[75]: 0.47540983606557374
```

```
In [76]: heart_disease["target"].value_counts()
```

Out[76]: 1 165 138

Name: target, dtype: int64

```
In [77]: # Import the RandomForestClassifier estimator class
         from sklearn.ensemble import RandomForestClassifier
         # Setup random seed
         np.random.seed(42)
         # Make the data
         X = heart_disease.drop("target", axis=1)
         y = heart_disease["target"]
         # Split the data
         X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
         # Instantiate Random Forest Classifier
         clf = RandomForestClassifier(n_estimators=100)
         clf.fit(X_train, y_train)
         # Evaluate the Random Forest Classifier
         clf.score(X_test, y_test)
```

Out[77]: 0.8524590163934426

Tidbit:

- 1. If you have structured data, used ensemble methods
- 2. If you have unstructured data, use deep learning or transfer learning

```
In [78]: heart_disease
```

Out[78]:

_		age	sex	ср	trestbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	са	thal	targe
	0	63	1	3	145	233	1	0	150	0	2.3	0	0	1	
	1	37	1	2	130	250	0	1	187	0	3.5	0	0	2	
	2	41	0	1	130	204	0	0	172	0	1.4	2	0	2	
	3	56	1	1	120	236	0	1	178	0	0.8	2	0	2	
	4	57	0	0	120	354	0	1	163	1	0.6	2	0	2	
	298	57	0	0	140	241	0	1	123	1	0.2	1	0	3	
	299	45	1	3	110	264	0	1	132	0	1.2	1	0	3	
	300	68	1	0	144	193	1	1	141	0	3.4	1	2	3	
	301	57	1	0	130	131	0	1	115	1	1.2	1	1	3	
	302	57	0	1	130	236	0	0	174	0	0.0	1	1	2	

303 rows × 14 columns

3. Fit the model/algorithm on our data and use it to make predictions

3.1 Fitting the model to the data

Different names for:

- X = features, features variables, data
- y = labels, targets, target variables

```
In [79]: # Import the RandomForestClassifier estimator class
         from sklearn.ensemble import RandomForestClassifier
         # Setup random seed
         np.random.seed(42)
         # Make the data
         X = heart disease.drop("target", axis=1)
         y = heart_disease["target"]
         # Split the data
         X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
         # Instantiate Random Forest Classifier
         clf = RandomForestClassifier(n estimators=100)
         # Fit the model to the data (training the machine learning model)
         clf.fit(X_train, y_train)
         # Evaluate the Random Forest Classifier (use the patterns the model has learne
         clf.score(X_test, y_test)
```

Out[79]: 0.8524590163934426

```
In [80]: X.head()
```

Out[80]:

	age	sex	ср	trestbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	са	thal
0	63	1	3	145	233	1	0	150	0	2.3	0	0	1
1	37	1	2	130	250	0	1	187	0	3.5	0	0	2
2	41	0	1	130	204	0	0	172	0	1.4	2	0	2
3	56	1	1	120	236	0	1	178	0	8.0	2	0	2
4	57	0	0	120	354	0	1	163	1	0.6	2	0	2

```
In [81]: y.tail()
Out[81]: 298
         299
                 0
         300
                 0
         301
                 0
         302
         Name: target, dtype: int64
```

Random Forest model deep dive

These resources will help you understand what's happening inside the Random Forest models we've been using.

- Random Forest Wikipedia (https://en.wikipedia.org/wiki/Random_forest)
- Random Forest Wikipedia (simple version) (https://simple.wikipedia.org/wiki/Random_forest)
- Random Forests in Python (http://blog.yhat.com/posts/random-forests-in-python.html) by yhat
- An Implementation and Explanation of the Random Forest in Python (https://towardsdatascience.com/animplementation-and-explanation-of-the-random-forest-in-python-77bf308a9b76) by Will Koehrsen

3.2 Make predictions using a machine learning model

2 ways to make predictions:

```
1. predict()
2. predict_proba()
```

```
In [82]: # Use a trained model to make predictions
clf.predict(np.array([1, 7, 8, 3, 4])) # this doesn't work...
```

```
ValueError
                                          Traceback (most recent call last)
<ipython-input-82-5908053f578c> in <module>
      1 # Use a trained model to make predictions
----> 2 clf.predict(np.array([1, 7, 8, 3, 4])) # this doesn't work...
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/ensemble/_forest.py in predict(self, X)
    610
                    The predicted classes.
    611
--> 612
                proba = self.predict proba(X)
    613
    614
                if self.n outputs == 1:
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/ensemble/ forest.py in predict proba(self, X)
    654
                check is fitted(self)
    655
                # Check data
--> 656
                X = self. validate X predict(X)
    657
    658
                # Assign chunk of trees to jobs
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/ensemble/ forest.py in validate X predict(self, X)
                check_is_fitted(self)
    410
    411
--> 412
                return self.estimators_[0]._validate_X_predict(X, check_input
=True)
    413
    414
            @property
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/tree/ classes.py in validate X predict(self, X, check input)
    378
                """Validate X whenever one tries to predict, apply, predict_p
roba"""
    379
                if check input:
                    X = check_array(X, dtype=DTYPE, accept_sparse="csr")
--> 380
    381
                    if issparse(X) and (X.indices.dtype != np.intc or
                                        X.indptr.dtype != np.intc):
    382
~/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-packages/sklear
n/utils/validation.py in check array(array, accept sparse, accept large spars
e, dtype, order, copy, force_all_finite, ensure_2d, allow_nd, ensure_min_samp
les, ensure_min_features, warn_on_dtype, estimator)
    554
                            "Reshape your data either using array.reshape(-1,
1) if "
                            "your data has a single feature or array.reshape
    555
(1, -1) "
--> 556
                            "if it contains a single sample.".format(array))
    557
    558
                # in the future np.flexible dtypes will be handled like objec
t dtypes
ValueError: Expected 2D array, got 1D array instead:
array=[1. 7. 8. 3. 4.].
Reshape your data either using array.reshape(-1, 1) if your data has a single
feature or array.reshape(1, -1) if it contains a single sample.
```

```
In [83]:
         X test.head()
Out[83]:
                        cp trestbps
                                    chol fbs restecg thalach exang oldpeak slope ca
                                                                                   thal
               age
                   sex
                                                                                 1
          179
                57
                         0
                               150
                                    276
                                          0
                                                        112
                                                                      0.6
                                                                                      1
          228
                59
                     1
                         3
                               170
                                    288
                                          0
                                                  0
                                                        159
                                                                0
                                                                      0.2
                                                                              1
                                                                                 0
                                                                                      3
           111
                57
                         2
                               150
                                    126
                                                  1
                                                       173
                                                                0
                                                                      0.2
                                                                                      3
                     1
                                          1
                                                                              2
                                                                                 1
          246
                56
                     0
                         0
                               134
                                    409
                                                        150
                                                                      1.9
                                                                                      3
           60
                71
                     0
                         2
                               110
                                    265
                                          1
                                                  0
                                                       130
                                                                      0.0
                                                                              2
                                                                                 1
                                                                                      2
In [84]: | clf.predict(X test)
Out[84]: array([0, 1, 1, 0, 1, 1, 1, 0, 0, 1, 1, 0, 1, 0, 1, 1, 1, 0, 0, 0, 0, 0,
                 1, 1, 1, 1, 1, 1, 0, 1, 0, 0, 0, 0, 1, 0, 1, 1, 1, 1, 1, 1, 1, 1,
                 1, 0, 1, 1, 0, 0, 0, 0, 1, 1, 0, 0, 0, 1, 0, 0, 0
In [85]: np.array(y test)
Out[85]: array([0, 0, 1, 0, 1, 1, 1, 0, 0, 1, 1, 1, 1, 0, 1, 1, 1, 0, 0, 0, 1, 0,
                 0, 1, 1, 0, 0, 1, 0, 1, 1, 0, 0, 1, 1, 0, 1, 1, 1, 1, 1, 1, 1, 1,
                 1, 0, 0, 1, 0, 0, 0, 0, 1, 1, 0, 0, 0, 1, 0, 0, 0])
In [86]:
         # Compare predictions to truth labels to evaluate the model
          y preds = clf.predict(X test)
          np.mean(y preds == y test)
Out[86]: 0.8524590163934426
In [87]: | clf.score(X_test, y_test)
Out[87]: 0.8524590163934426
In [88]: from sklearn.metrics import accuracy score
          accuracy score(y test, y preds)
Out[88]: 0.8524590163934426
```

Make predictions with predict proba() - use this if someone asks you "what's the probability your model is assigning to each prediction?"

```
In [89]: # predict_proba() returns probabilities of a classification label
         clf.predict proba(X test[:5])
Out[89]: array([[0.89, 0.11],
                 [0.49, 0.51],
                [0.43, 0.57],
                 [0.84, 0.16],
                 [0.18, 0.82]]
```

In [90]: # Let's predict() on the same data... clf.predict(X_test[:5])

Out[90]: array([0, 1, 1, 0, 1])

In [91]: X_test[:5]

Out[91]:

	age	sex	ср	trestbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	са	thal
179	57	1	0	150	276	0	0	112	1	0.6	1	1	1
228	59	1	3	170	288	0	0	159	0	0.2	1	0	3
111	57	1	2	150	126	1	1	173	0	0.2	2	1	3
246	56	0	0	134	409	0	0	150	1	1.9	1	2	3
60	71	0	2	110	265	1	0	130	0	0.0	2	1	2

In [92]: heart_disease["target"].value_counts()

Out[92]: 1 165

138

Name: target, dtype: int64

predict() can also be used for regression models.

In [93]: boston_df.head()

Out[93]:

	CRIM	ZN	INDUS	CHAS	NOX	RM	AGE	DIS	RAD	TAX	PTRATIO	В	LST.
0	0.00632	18.0	2.31	0.0	0.538	6.575	65.2	4.0900	1.0	296.0	15.3	396.90	4.
1	0.02731	0.0	7.07	0.0	0.469	6.421	78.9	4.9671	2.0	242.0	17.8	396.90	9.
2	0.02729	0.0	7.07	0.0	0.469	7.185	61.1	4.9671	2.0	242.0	17.8	392.83	4.
3	0.03237	0.0	2.18	0.0	0.458	6.998	45.8	6.0622	3.0	222.0	18.7	394.63	2.
4	0.06905	0.0	2.18	0.0	0.458	7.147	54.2	6.0622	3.0	222.0	18.7	396.90	5.
4													•

```
In [94]: from sklearn.ensemble import RandomForestRegressor
         np.random.seed(42)
         # Create the data
         X = boston_df.drop("target", axis=1)
         y = boston_df["target"]
         # Split into training and test sets
         X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
         # Instantiate and fit model
         model = RandomForestRegressor(n_estimators=100).fit(X_train, y_train)
         # Make predictions
         y_preds = model.predict(X_test)
In [95]: y_preds[:10]
Out[95]: array([23.002, 30.826, 16.734, 23.467, 16.853, 21.725, 19.232, 15.239,
                21.067, 20.738])
In [96]: np.array(y test[:10])
Out[96]: array([23.6, 32.4, 13.6, 22.8, 16.1, 20. , 17.8, 14. , 19.6, 16.8])
In [97]: # Compare the predictions to the truth
         from sklearn.metrics import mean_absolute_error
         mean_absolute_error(y_test, y_preds)
Out[97]: 2.1226372549019623
```

4. Evaluating a machine learning model

Three ways to evaluate Scikit-Learn models/esitmators:

- 1. Estimator score method
- 2. The scoring parameter
- 3. Problem-specific metric functions.

4.1 Evaluating a model with the score method

```
In [98]: from sklearn.ensemble import RandomForestClassifier
             np.random.seed(42)
             X = heart_disease.drop("target", axis=1)
             y = heart disease["target"]
            X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
             clf = RandomForestClassifier()
             clf.fit(X_train, y_train)
   Out[98]: RandomForestClassifier(bootstrap=True, ccp alpha=0.0, class weight=None,
                                    criterion='gini', max_depth=None, max_features='auto',
                                    max_leaf_nodes=None, max_samples=None,
                                    min_impurity_decrease=0.0, min_impurity_split=None,
                                    min samples leaf=1, min samples split=2,
                                    min weight fraction leaf=0.0, n estimators=100,
                                    n_jobs=None, oob_score=False, random_state=None,
                                    verbose=0, warm_start=False)
   In [99]: clf.score(X train, y train)
  Out[99]: 1.0
  In [100]: clf.score(X_test, y_test)
  Out[100]: 0.8524590163934426
Let's do the same but for regression...
  In [101]: from sklearn.ensemble import RandomForestRegressor
            np.random.seed(42)
             # Create the data
             X = boston_df.drop("target", axis=1)
             y = boston_df["target"]
             # Split into training and test sets
             X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
             # Instantiate and fit model
```

```
4.2 Evaluating a model using the scoring parameter
```

In [102]: | model.score(X_test, y_test)

Out[102]: 0.873969014117403

model = RandomForestRegressor(n estimators=100).fit(X train, y train)

```
In [103]: from sklearn.model selection import cross val score
          from sklearn.ensemble import RandomForestClassifier
          np.random.seed(42)
          X = heart disease.drop("target", axis=1)
          y = heart disease["target"]
          X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
          clf = RandomForestClassifier(n estimators=100)
          clf.fit(X_train, y_train);
In [104]: | clf.score(X test, y test)
Out[104]: 0.8524590163934426
In [105]: cross_val_score(clf, X, y, cv=5)
Out[105]: array([0.81967213, 0.86885246, 0.81967213, 0.78333333, 0.76666667])
In [106]: | cross_val_score(clf, X, y, cv=10)
, 0.73333333, 0.86666667, 0.733333333, 0.8
                                                                        1)
In [107]: | np.random.seed(42)
          # Single training and test split score
          clf_single_score = clf.score(X_test, y_test)
          # Take the mean of 5-fold cross-validation score
          clf_cross_val_score = np.mean(cross_val_score(clf, X, y, cv=5))
          # Compare the two
          clf_single_score, clf_cross_val_score
Out[107]: (0.8524590163934426, 0.8248087431693989)
In [108]: # Default scoring parameter of classifier = mean accuracy
          clf.score()
                                                  Traceback (most recent call last)
          TypeError
          <ipython-input-108-cca012993b3a> in <module>
               1 # Default scoring parameter of classifier = mean accuracy
          ----> 2 clf.score()
          TypeError: score() missing 2 required positional arguments: 'X' and 'y'
  In [ ]: # Scoring parameter set to None by default
          cross val score(clf, X, y, cv=5, scoring=None)
```

4.2.1 Classification model evaluation metrics

- 1. Accuracy
- 2. Area under ROC curve
- 3. Confusion matrix
- 4. Classification report

Accuracy

```
heart disease.head()
In [ ]:
In [ ]: | from sklearn.model_selection import cross_val_score
        from sklearn.ensemble import RandomForestClassifier
        np.random.seed(42)
        X = heart_disease.drop("target", axis=1)
        y = heart_disease["target"]
        clf = RandomForestClassifier(n estimators=100)
         cross_val_score = cross_val_score(clf, X, y, cv=5)
In [ ]: np.mean(cross val score)
In [ ]: print(f"Heart Disease Classifier Cross-Validated Accuracy: {np.mean(cross val
        score) *100:.2f}%")
```

Area under the receiver operating characteristic curve (AUC/ROC)

- Area under curve (AUC)
- ROC curve

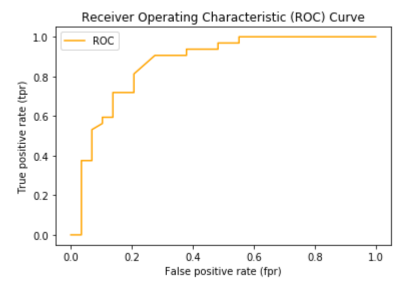
ROC curves are a comparison of a model's true postive rate (tpr) versus a models false positive rate (fpr).

- True positive = model predicts 1 when truth is 1
- False positive = model predicts 1 when truth is 0
- True negative = model predicts 0 when truth is 0
- False negative = model predicts 0 when truth is 1

```
In [109]: # Create X test... etc
          X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
```

```
In [110]: | from sklearn.metrics import roc_curve
          # Fit the classifier
          clf.fit(X train, y train)
          # Make predictions with probabilities
          y probs = clf.predict proba(X test)
          y_probs[:10], len(y_probs)
Out[110]: (array([[0.51, 0.49],
                  [0.17, 0.83],
                  [0.51, 0.49],
                  [0.72, 0.28],
                   [0.43, 0.57],
                  [0.12, 0.88],
                   [0.3, 0.7],
                  [0.97, 0.03],
                  [0.15, 0.85],
                  [0.4, 0.6]
           61)
In [111]: y probs positive = y probs[:, 1]
          y_probs_positive[:10]
Out[111]: array([0.49, 0.83, 0.49, 0.28, 0.57, 0.88, 0.7, 0.03, 0.85, 0.6])
In [112]: # Caculate fpr, tpr and thresholds
          fpr, tpr, thresholds = roc_curve(y_test, y_probs_positive)
          # Check the false positive rates
          fpr
Out[112]: array([0.
                           , 0.03448276, 0.03448276, 0.03448276, 0.03448276,
                 0.03448276, 0.03448276, 0.06896552, 0.06896552, 0.06896552,
                 0.10344828, 0.10344828, 0.13793103, 0.13793103, 0.13793103,
                 0.20689655, 0.20689655, 0.20689655, 0.27586207, 0.37931034,
                 0.37931034, 0.48275862, 0.48275862, 0.55172414, 0.55172414,
                 1.
                            1)
```

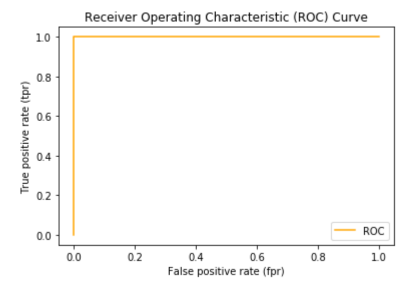
```
In [113]:
          # Create a function for plotting ROC curves
          import matplotlib.pyplot as plt
          def plot roc curve(fpr, tpr):
              Plots a ROC curve given the false positive rate (fpr)
              and true positive rate (tpr) of a model.
              # Plot roc curve
              plt.plot(fpr, tpr, color="orange", label="ROC")
              # Plot line with no predictive power (baseline)
              #plt.plot([0, 1], [0, 1], color="darkblue", linestyle="--", label="Guessin
          g")
              # Customize the plot
              plt.xlabel("False positive rate (fpr)")
              plt.ylabel("True positive rate (tpr)")
              plt.title("Receiver Operating Characteristic (ROC) Curve")
              plt.legend()
              plt.show()
          plot_roc_curve(fpr, tpr)
```



```
In [114]: from sklearn.metrics import roc auc score
          roc_auc_score(y_test, y_probs_positive)
```

Out[114]: 0.8669181034482759

```
# Plot perfect ROC curve and AUC score
fpr, tpr, thresholds = roc_curve(y_test, y_test)
plot_roc_curve(fpr, tpr)
```



```
In [116]:
          # Perfect AUC score
          roc_auc_score(y_test, y_test)
Out[116]: 1.0
```

Confusion Matrix

A confusion matrix is a quick way to compare the labels a model predicts and the actual labels it was supposed to predict.

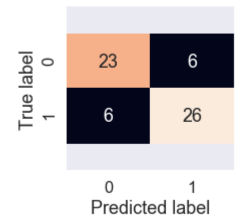
In essence, giving you an idea of where the model is getting confused.

```
In [117]:
          from sklearn.metrics import confusion matrix
          y_preds = clf.predict(X_test)
          confusion_matrix(y_test, y_preds)
Out[117]: array([[23, 6],
                 [ 6, 26]])
```

```
In [118]: # Visualize confusion matrix with pd.crosstab()
           pd.crosstab(y_test,
                       y_preds,
                       rownames=["Actual Labels"],
                       colnames=["Predicted Labels"])
Out[118]:
           Predicted Labels
                              1
              Actual Labels
                       0 23
                              6
                           6 26
In [119]: 22 + 7 + 8 + 24
Out[119]: 61
In [120]: len(X_test)
Out[120]: 61
In [121]: # # How install a conda package into the current envrionment from a Jupyter No
          tebook
           # import sys
           # !conda install --yes --prefix {sys.prefix} seaborn
In [122]: # Make our confusion matrix more visual with Seaborn's heatmap()
           import seaborn as sns
           # Set the font scale
           sns.set(font_scale=1.5)
           # Create a confusion matrix
           conf_mat = confusion_matrix(y_test, y_preds)
           # Plot it using Seaborn
           sns.heatmap(conf mat);
                                                       - 25
           0
                                                       20
                                                       - 15
                                                       - 10
                       0
                                         1
```

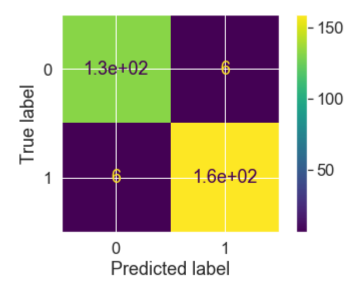
Note: In the original notebook, the function below had the "True label" as the x-axis label and the "Predicted label" as the y-axis label. But due to the way confusion_matrix() (https://scikitlearn.org/stable/modules/generated/sklearn.metrics.confusion matrix.html) outputs values, these should be swapped around. The code below has been corrected.

```
In [123]:
          def plot_conf_mat(conf_mat):
              Plots a confusion matrix using Seaborn's heatmap().
              fig, ax = plt.subplots(figsize=(3,3))
              ax = sns.heatmap(conf_mat,
                                annot=True, # Annotate the boxes with conf_mat info
                                cbar=False)
              plt.xlabel("Predicted label")
              plt.ylabel("True label")
              # Fix the broken annotations (this happened in Matplotlib 3.1.1)
              bottom, top = ax.get_ylim()
              ax.set ylim(bottom + 0.5, top-0.5);
          plot_conf_mat(conf_mat)
```



In [124]: from sklearn.metrics import plot_confusion_matrix plot_confusion_matrix(clf, X, y)

Out[124]: <sklearn.metrics._plot.confusion_matrix.ConfusionMatrixDisplay at 0x7ff630bc2</pre> d10>



Classification Report

In [125]: from sklearn.metrics import classification_report print(classification_report(y_test, y_preds))

	precision	recall	f1-score	support
0	0.79	0.79	0.79	29
1	0.81	0.81	0.81	32
accuracy			0.80	61
macro avg	0.80	0.80	0.80	61
weighted avg	0.80	0.80	0.80	61

```
In [126]: # Where precision and recall become valuable
          disease true = np.zeros(10000)
          disease true[0] = 1 # only one positive case
          disease preds = np.zeros(10000) # model predicts every case as 0
          pd.DataFrame(classification report(disease true,
                                              disease preds,
                                              output dict=True))
```

/Users/daniel/Desktop/ml-course/zero-to-mastery-ml/env/lib/python3.7/site-pac kages/sklearn/metrics/ classification.py:1272: UndefinedMetricWarning: Precis ion and F-score are ill-defined and being set to 0.0 in labels with no predic ted samples. Use `zero division` parameter to control this behavior. warn prf(average, modifier, msg start, len(result))

Out[126]:

	0.0	1.0	accuracy	macro avg	weighted avg
precision	0.99990	0.0	0.9999	0.499950	0.99980
recall	1.00000	0.0	0.9999	0.500000	0.99990
f1-score	0.99995	0.0	0.9999	0.499975	0.99985
support	9999.00000	1.0	0.9999	10000.000000	10000.00000

To summarize classification metrics:

- Accuracy is a good measure to start with if all classes are balanced (e.g. same amount of samples which are labelled with 0 or 1).
- Precision and recall become more important when classes are imbalanced.
- If false positive predictions are worse than false negatives, aim for higher precision.
- If false negative predictions are worse than false positives, aim for higher recall.
- F1-score is a combination of precision and recall.

4.2.2 Regression model evaluation metrics

Model evaluation metrics documentation - https://scikit-learn.org/stable/modules/model evaluation.html (https://scikit-learn.org/stable/modules/model_evaluation.html)

- 1. R² (pronounced r-squared) or coefficient of determination.
- 2. Mean absolute error (MAE)
- 3. Mean squared error (MSE)

R^2

What R-squared does: Compares your models predictions to the mean of the targets. Values can range from negative infinity (a very poor model) to 1. For example, if all your model does is predict the mean of the targets, it's R^2 value would be 0. And if your model perfectly predicts a range of numbers it's R^2 value would be 1.

```
In [127]: from sklearn.ensemble import RandomForestRegressor
          np.random.seed(42)
          X = boston_df.drop("target", axis=1)
          y = boston_df["target"]
          X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
          model = RandomForestRegressor(n_estimators=100)
          model.fit(X_train, y_train);
In [128]: model.score(X_test, y_test)
Out[128]: 0.873969014117403
In [129]: | from sklearn.metrics import r2_score
          # Fill an array with y test mean
          y_test_mean = np.full(len(y_test), y_test.mean())
In [130]: y_test.mean()
Out[130]: 21.488235294117644
In [131]: # Model only predicting the mean gets an R^2 score of 0
          r2_score(y_test, y_test_mean)
Out[131]: 0.0
In [132]: # Model predicting perfectly the correct values gets an R^2 score of 1
          r2_score(y_test, y_test)
Out[132]: 1.0
```

Mean absolue error (MAE)

MAE is the average of the aboslute differences between predictions and actual values. It gives you an idea of how wrong your models predictions are.

```
In [133]: # Mean absolute error
          from sklearn.metrics import mean absolute error
          y_preds = model.predict(X_test)
          mae = mean_absolute_error(y_test, y_preds)
          mae
Out[133]: 2.1226372549019623
```

```
In [134]: | df = pd.DataFrame(data={"actual values": y_test,
                                   "predicted values": y_preds})
          df["differences"] = df["predicted values"] - df["actual values"]
```

Out[134]:

	actual values	predicted values	differences
173	23.6	23.002	-0.598
274	32.4	30.826	-1.574
491	13.6	16.734	3.134
72	22.8	23.467	0.667
452	16.1	16.853	0.753
412	17.9	13.030	-4.870
436	9.6	12.490	2.890
411	17.2	13.406	-3.794
86	22.5	20.219	-2.281
75	21.4	23.898	2.498

102 rows × 3 columns

Mean squared error (MSE)

```
In [135]: # Mean squared error
          from sklearn.metrics import mean_squared_error
          y_preds = model.predict(X_test)
          mse = mean_squared_error(y_test, y_preds)
Out[135]: 9.242328990196082
In [136]: # Calculate MSE by hand
          squared = np.square(df["differences"])
          squared.mean()
Out[136]: 9.242328990196082
```

4.2.3 Finally using the scoring parameter

```
In [137]: from sklearn.model selection import cross val score
          from sklearn.ensemble import RandomForestClassifier
          np.random.seed(42)
          X = heart_disease.drop("target", axis=1)
          y = heart_disease["target"]
          clf = RandomForestClassifier(n estimators=100)
In [138]: | np.random.seed(42)
          cv acc = cross val score(clf, X, y, cv=5, scoring=None)
          cv acc
Out[138]: array([0.81967213, 0.90163934, 0.83606557, 0.783333333, 0.78333333])
In [139]: # Cross-validated accuracy
          print(f'The cross-validated accuracy is: {np.mean(cv_acc)*100:.2f}%')
          The cross-validated accuracy is: 82.48%
In [140]: | np.random.seed(42)
          cv_acc = cross_val_score(clf, X, y, cv=5, scoring="accuracy")
          print(f'The cross-validated accuracy is: {np.mean(cv_acc)*100:.2f}%')
          The cross-validated accuracy is: 82.48%
In [141]: # Precision
          cv_precision = cross_val_score(clf, X, y, cv=5, scoring="precision")
          np.mean(cv_precision)
Out[141]: 0.8085601538512754
In [142]: | # Recall
          cv recall = cross val score(clf, X, y, cv=5, scoring="recall")
          np.mean(cv recall)
Out[142]: 0.8424242424242424
In [143]: | cv f1 = cross val score(clf, X, y, cv=5, scoring="f1")
          np.mean(cv f1)
Out[143]: 0.841476533416832
```

How about our regression model?

```
from sklearn.model selection import cross val score
          from sklearn.ensemble import RandomForestRegressor
          np.random.seed(42)
          X = boston_df.drop("target", axis=1)
          y = boston_df["target"]
          model = RandomForestRegressor(n estimators=100)
In [145]: np.random.seed(42)
          cv r2 = cross val score(model, X, y, cv=5, scoring=None)
          np.mean(cv r2)
Out[145]: 0.622375083951403
In [146]: | np.random.seed(42)
          cv_r2 = cross_val_score(model, X, y, cv=5, scoring="r2")
Out[146]: array([0.76861165, 0.85851765, 0.74941131, 0.47891315, 0.25642166])
In [147]: # Mean absolute error
          cv mae = cross val score(model, X, y, cv=5, scoring="neg mean absolute error")
          cv mae
Out[147]: array([-2.12751961, -2.53956436, -3.42026733, -3.82432673, -3.06893069])
In [148]: # Mean squared error
          cv_mse = cross_val_score(model, X, y, cv=5, scoring="neg_mean_squared_error")
          np.mean(cv_mse)
Out[148]: -21.02253826604542
```

4.3 Using different evalution metrics as Scikit-Learn functions

Classification evaluation functions

```
In [149]:
          from sklearn.metrics import accuracy score, precision score, recall score, f1
          from sklearn.ensemble import RandomForestClassifier
          from sklearn.model selection import train test split
          np.random.seed(42)
          X = heart disease.drop("target", axis=1)
          y = heart disease["target"]
          X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
          clf = RandomForestClassifier(n_estimators=100)
          clf.fit(X train, y train)
          # Make some predictions
          y_preds = clf.predict(X_test)
          # Evaluate the classifier
          print("Classifier metrics on the test set")
          print(f"Accuracy: {accuracy_score(y_test, y_preds)*100:.2f}%")
          print(f"Precision: {precision_score(y_test, y_preds)}")
          print(f"Recall: {recall_score(y_test, y_preds)}")
          print(f"F1: {f1_score(y_test, y_preds)}")
```

Classifier metrics on the test set

Accuracy: 85.25%

Precision: 0.84848484848485

Recall: 0.875

F1: 0.8615384615384615

Regression evaluation functions

```
In [150]:
          from sklearn.metrics import r2 score, mean absolute error, mean squared error
          from sklearn.ensemble import RandomForestRegressor
          from sklearn.model selection import train test split
          np.random.seed(42)
          X = boston df.drop("target", axis=1)
          y = boston df["target"]
          X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
          model = RandomForestRegressor(n_estimators=100)
          model.fit(X_train, y_train)
          # Make predictions using our regression model
          y_preds = model.predict(X_test)
          # Evaluate the regression model
          print("Regression model metrics on the test set")
          print(f"R^2: {r2 score(y test, y preds)}")
          print(f"MAE: {mean_absolute_error(y_test, y_preds)}")
          print(f"MSE: {mean_squared_error(y_test, y_preds)}")
```

Regression model metrics on the test set

R^2: 0.8739690141174031 MAE: 2.1226372549019623 MSE: 9.242328990196082

5. Improving a model

First predictions = baseline predictions. First model = baseline model.

From a data perspective:

- Could we collect more data? (generally, the more data, the better)
- Could we improve our data?

From a model perspective:

- Is there a better model we could use?
- Could we improve the current model?

Hyperparameters vs. Parameters

- Parameters = model find these patterns in data
- Hyperparameters = settings on a model you can adjust to (potentially) improve its ability to find patterns

Three ways to adjust hyperparameters:

- 1. By hand
- 2. Randomly with RandomSearchCV
- Exhaustively with GridSearchCV

```
In [ ]: from sklearn.ensemble import RandomForestClassifier
        clf = RandomForestClassifier(n_estimator=100)
In [ ]: clf.get params()
```

5.1 Tuning hyperparameters by hand

Let's make 3 sets, training, validation and test.

```
In [ ]: clf.get params()
```

We're going to try and adjust:

- max_depth
- max features
- min samples leaf
- min samples split
- n estimators

```
In [ ]: | def evaluate_preds(y_true, y_preds):
            Performs evaluation comparison on y_true labels vs. y_pred labels
            on a classification.
            accuracy = accuracy_score(y_true, y_preds)
            precision = precision score(y true, y preds)
            recall = recall_score(y_true, y_preds)
            f1 = f1_score(y_true, y_preds)
            metric_dict = {"accuracy": round(accuracy, 2),
                            "precision": round(precision, 2),
                            "recall": round(recall, 2),
                            "f1": round(f1, 2)}
            print(f"Acc: {accuracy * 100:.2f}%")
            print(f"Precision: {precision:.2f}")
            print(f"Recall: {recall:.2f}")
            print(f"F1 score: {f1:.2f}")
            return metric dict
```

```
In [ ]: from sklearn.ensemble import RandomForestClassifier
        np.random.seed(42)
        # Shuffle the data
        heart_disease_shuffled = heart_disease.sample(frac=1)
        # Split into X & v
        X = heart disease shuffled.drop("target", axis=1)
        y = heart_disease_shuffled["target"]
        # Split the data into train, validation & test sets
        train_split = round(0.7 * len(heart_disease_shuffled)) # 70% of data
        valid_split = round(train_split + 0.15 * len(heart_disease_shuffled)) # 15% of
        data
        X_train, y_train = X[:train_split], y[:train_split]
        X valid, y valid = X[train split:valid split], y[train split:valid split]
        X_test, y_test = X[valid_split:], y[:valid_split]
        clf = RandomForestClassifier()
        clf.fit(X train, y train)
        # Make baseline predictions
        y_preds = clf.predict(X_valid)
        # Evaluate the classifier on validation set
        baseline metrics = evaluate preds(y valid, y preds)
        baseline_metrics
In [ ]: | np.random.seed(42)
        # Create a second classifier with different hyperparameters
        clf 2 = RandomForestClassifier(n estimators=100)
```

```
clf_2.fit(X_train, y_train)
# Make predictions with different hyperparameters
y_preds_2 = clf_2.predict(X_valid)
# Evalute the 2nd classsifier
clf_2_metrics = evaluate_preds(y_valid, y_preds_2)
```

5.2 Hyperparameter tuning with RandomizedSearchCV

```
In [ ]: | from sklearn.model selection import RandomizedSearchCV
        grid = {"n_estimators": [10, 100, 200, 500, 1000, 1200],
                 "max depth": [None, 5, 10, 20, 30],
                 "max_features": ["auto", "sqrt"],
                 "min_samples_split": [2, 4, 6],
                 "min_samples_leaf": [1, 2, 4]}
        np.random.seed(42)
        # Split into X & y
        X = heart_disease_shuffled.drop("target", axis=1)
        y = heart_disease_shuffled["target"]
        # Split into train and test sets
        X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
        # Instantiate RandomForestClassifier
        clf = RandomForestClassifier(n jobs=1)
        # Setup RandomizedSearchCV
        rs_clf = RandomizedSearchCV(estimator=clf,
                                     param distributions=grid,
                                     n_iter=10, # number of models to try
                                     verbose=2)
        # Fit the RandomizedSearchCV version of clf
        rs clf.fit(X train, y train);
In [ ]: | rs_clf.best_params_
In [ ]: # Make predictions with the best hyperparameters
        rs_y_preds = rs_clf.predict(X_test)
        # Evaluate the predictions
        rs_metrics = evaluate_preds(y_test, rs_y_preds)
```

5.3 Hyperparameter tuning with GridSearchCV

```
In [ ]: grid
In [ ]: grid_2 = {'n_estimators': [100, 200, 500],
                   'max depth': [None],
                   'max_features': ['auto', 'sqrt'],
                   'min_samples_split': [6],
                   'min samples leaf': [1, 2]}
```

```
In [ ]: from sklearn.model selection import GridSearchCV, train test split
        np.random.seed(42)
        # Split into X & y
        X = heart_disease_shuffled.drop("target", axis=1)
        y = heart disease shuffled["target"]
        # Split into train and test sets
        X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
        # # Instantiate RandomForestClassifier
        # clf = RandomForestClassifier(n_jobs=1)
        # # Setup GridSearchCV
        # gs_clf = GridSearchCV(estimator=clf,
                                 param grid=grid 2,
        #
                                 cv=5,
        #
                                 verbose=2)
        # Fit the GridSearchCV version of clf
        #gs_clf.fit(X_train, y_train);
In [ ]: gs_clf.best_params_
In [ ]: | gs y preds = gs clf.predict(X test)
        # evaluate the predictions
        gs_metrics = evaluate_preds(y_test, gs_y_preds)
```

Let's compare our different models metrics.

```
In [ ]: | compare_metrics = pd.DataFrame({"baseline": baseline_metrics,
                                          "clf 2": clf 2 metrics,
                                          "random search": rs metrics,
                                          "grid search": gs metrics})
        compare_metrics.plot.bar(figsize=(10, 8));
```

6. Saving and loading trained machine learning models

Two ways to save and load machine learning models:

- 1. With Python's pickle module
- 2. With the joblib module

Pickle

```
In [ ]: import pickle
        # Save an extisting model to file
        pickle.dump(gs clf, open("gs random random forest model 1.pkl", "wb"))
In [ ]: # Load a saved model
        loaded pickle model = pickle.load(open("gs random random forest model 1.pkl",
In [ ]: | # Make some predictions
        pickle y preds = loaded pickle model.predict(X test)
        evaluate_preds(y_test, pickle_y_preds)
```

Joblib

```
In [ ]: from joblib import dump, load
        # Save model to file
        dump(gs_clf, filename="gs_random_forest_model_1.joblib")
In [ ]: # Import a saved joblib model
        loaded joblib model = load(filename="gs random forest model 1.joblib")
In [ ]: # Make and evaluate joblib predictions
        joblib_y_preds = loaded_joblib_model.predict(X_test)
        evaluate_preds(y_test, joblib_y_preds)
```

7. Putting it all together!

```
In [ ]: data = pd.read_csv("data/car-sales-extended-missing-data.csv")
In [ ]: data.dtypes
In [ ]: data.isna().sum()
```

Steps we want to do (all in one cell):

- 1. Fill missing data
- 2. Convert data to numbers
- Build a model on the data

```
In [ ]: # Getting data ready
        import pandas as pd
        from sklearn.compose import ColumnTransformer
        from sklearn.pipeline import Pipeline
        from sklearn.impute import SimpleImputer
        from sklearn.preprocessing import OneHotEncoder
        # Modelling
        from sklearn.ensemble import RandomForestRegressor
        from sklearn.model_selection import train_test_split, GridSearchCV
        # Setup random seed
        import numpy as np
        np.random.seed(42)
        # Import data and drop rows with missing labels
        data = pd.read csv("data/car-sales-extended-missing-data.csv")
        data.dropna(subset=["Price"], inplace=True)
         # Define different features and transformer pipeline
        categorical_features = ["Make", "Colour"]
        categorical transformer = Pipeline(steps=[
            ("imputer", SimpleImputer(strategy="constant", fill value="missing")),
            ("onehot", OneHotEncoder(handle unknown="ignore"))])
        door feature = ["Doors"]
        door transformer = Pipeline(steps=[
            ("imputer", SimpleImputer(strategy="constant", fill_value=4))
        1)
        numeric_features = ["Odometer (KM)"]
        numeric transformer = Pipeline(steps=[
             ("imputer", SimpleImputer(strategy="mean"))
        ])
        # Setup preprocessing steps (fill missing values, then convert to numbers)
        preprocessor = ColumnTransformer(
                             transformers=[
                                 ("cat", categorical transformer, categorical features
        ),
                                 ("door", door_transformer, door_feature),
                                 ("num", numeric transformer, numeric features)
                             1)
        # Creating a preprocessing and modelling pipeline
        model = Pipeline(steps=[("preprocessor", preprocessor),
                                 ("model", RandomForestRegressor())])
        # Split data
        X = data.drop("Price", axis=1)
        y = data["Price"]
        X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
        # Fit and score the model
        model.fit(X_train, y_train)
        model.score(X_test, y_test)
```

It's also possible to use GridSearchCV or RandomizedSesrchCV with our Pipeline.

```
In [ ]: # Use GridSearchCV with our regression Pipeline
        from sklearn.model selection import GridSearchCV
        pipe_grid = {
             "preprocessor__num__imputer__strategy": ["mean", "median"],
            "model__n_estimators": [100, 1000],
            "model _max_depth": [None, 5],
            "model__max_features": ["auto"],
             "model__min_samples_split": [2, 4]
        }
        gs_model = GridSearchCV(model, pipe_grid, cv=5, verbose=2)
         gs_model.fit(X_train, y_train)
In [ ]: | gs_model.score(X_test, y_test)
In [ ]: what_were_covering
```