4 Pre-Processing and Training Data

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4.2 Introduction

In preceding notebooks, performed preliminary assessments of data quality and refined the question to be answered. You found a small number of data values that gave clear choices about whether to replace values or drop a whole row. You determined that predicting the adult weekend ticket price was your primary aim. You threw away records with missing price data, but not before making the most of the other available data to look for any patterns between the states. You didn't see any and decided to treat all states equally; the state label didn't seem to be particularly useful.

In this notebook you'll start to build machine learning models. Before even starting with learning a machine learning model, however, start by considering how useful the mean value is as a predictor. This is more than just a pedagogical device. You never want to go to stakeholders with a machine learning model only to have the CEO point out that it performs worse than just guessing the average! Your first model is a baseline performance comparitor for any subsequent model. You then build up the process of efficiently and robustly creating and assessing models against it. The development we lay out may be little slower than in the real world, but this step of the capstone is definitely more than just instructional. It is good practice to build up an understanding that the machine learning pipelines you build work as expected. You can validate steps with your own functions for checking expected equivalence between, say, pandas and sklearn implementations.

4.3 Imports

```
In [1]:
        import pandas as pd
        import numpy as np
        import os
        import pickle
        import matplotlib.pyplot as plt
        import seaborn as sns
        from sklearn import __version__ as sklearn_version
        from sklearn.decomposition import PCA
        from sklearn.preprocessing import scale
        from sklearn.model selection import train test split, cross validate, GridSearchCV, learning curve
        from sklearn.preprocessing import StandardScaler, MinMaxScaler
        from sklearn.dummy import DummyRegressor
        from sklearn.linear model import LinearRegression
        from sklearn.ensemble import RandomForestRegressor
        from sklearn.metrics import r2 score, mean squared error, mean absolute error
        from sklearn.pipeline import make pipeline
        from sklearn.impute import SimpleImputer
        from sklearn.feature selection import SelectKBest, f regression
        import datetime
        from library.sb utils import save file
```

4.4 Load Data

```
ski_data = pd.read_csv('../data/ski_data_step3_features.csv')
          ski data.head().T
Out[2]:
                                                        0
                                                                           1
                                                                                          2
                                                                                                            3
                                                                                                                               4
                                     Name Alyeska Resort Eaglecrest Ski Area Hilltop Ski Area Arizona Snowbowl Sunrise Park Resort
                                    Region
                                                    Alaska
                                                                      Alaska
                                                                                     Alaska
                                                                                                      Arizona
                                                                                                                         Arizona
                                      state
                                                    Alaska
                                                                      Alaska
                                                                                     Alaska
                                                                                                      Arizona
                                                                                                                         Arizona
                               summit_elev
                                                     3939
                                                                        2600
                                                                                       2090
                                                                                                       11500
                                                                                                                           11100
                               vertical_drop
                                                     2500
                                                                        1540
                                                                                        294
                                                                                                        2300
                                                                                                                           1800
                                  base_elev
                                                      250
                                                                        1200
                                                                                       1796
                                                                                                        9200
                                                                                                                           9200
                                                                                          0
                                                                                                            0
                                     trams
                                                                           0
                                                                                                                               0
                                  fastSixes
                                                        0
                                                                           0
                                                                                          0
                                                                                                            1
                                                                                                                               0
                                 fastQuads
                                                                           0
                                                                                          0
                                                                                                            0
                                                        2
                                                                           0
                                                                                          0
                                                                                                            2
                                      quad
                                      triple
                                                                                                            2
                                                        0
                                                                           0
                                                                                          1
                                                                                                                               3
```

4.5 Extract Big Mountain Data

Big Mountain is your resort. Separate it from the rest of the data to use later.

```
In [3]: big_mountain = ski_data[ski_data.Name == 'Big Mountain Resort']
```

Out[4]:

124

Name	Big Mountain Resort
Region	Montana
state	Montana
summit_elev	6817
vertical_drop	2353
base_elev	4464
trams	0
fastSixes	0
fastQuads	3
quad	2
triple	6
double	0
surface	3
total_chairs	14
Runs	105
TerrainParks	4
LongestRun_mi	3.3
SkiableTerrain_ac	3000
Snow Making_ac	600
daysOpenLastYear	123
yearsOpen	72
averageSnowfall	333
AdultWeekend	81
projectedDaysOpen	123
NightSkiing_ac	600
resorts_per_state	12

	124
resorts_per_100kcapita	1.12278
resorts_per_100ksq_mile	8.16104
resort_skiable_area_ac_state_ratio	0.140121
resort_days_open_state_ratio	0.129338
resort_terrain_park_state_ratio	0.148148
resort_night_skiing_state_ratio	0.84507
total_chairs_runs_ratio	0.133333
total_chairs_skiable_ratio	0.00466667
fastQuads_runs_ratio	0.0285714
fastQuads_skiable_ratio	0.001

```
In [5]: ski_data.shape
Out[5]: (277, 36)
In [6]: ski_data = ski_data[ski_data.Name != 'Big Mountain Resort']
In [7]: ski_data.shape
Out[7]: (276, 36)
```

4.6 Train/Test Split

So far, you've treated ski resort data as a single entity. In machine learning, when you train your model on all of your data, you end up with no data set aside to evaluate model performance. You could keep making more and more complex models that fit the data better and better and not realise you were overfitting to that one set of samples. By partitioning the data into training and testing splits, without letting a model (or missing-value imputation) learn anything about the test split, you have a somewhat independent assessment of how your model might perform in the future. An often overlooked subtlety here is that people all too frequently use the test set to assess model performance and then compare multiple models to pick the best. This means their overall model selection process is fitting to one specific data set, now the test split. You could keep going, trying to get better and better performance on that one data set, but that's where cross-validation becomes especially useful. While training models, a test split is very useful as a final check on expected future performance.

What partition sizes would you have with a 70/30 train/test split?

Out[12]: ((193, 32), (83, 32))

```
In [8]: len(ski_data) * .7, len(ski_data) * .3
 Out[8]: (193.2, 82.8)
 In [9]: | X_train, X_test, y_train, y_test = train_test_split(ski_data.drop(columns='AdultWeekend'),
                                                              ski data.AdultWeekend, test size=0.3,
                                                              random state=47)
In [10]: X_train.shape, X_test.shape
Out[10]: ((193, 35), (83, 35))
In [11]: y_train.shape, y_test.shape
Out[11]: ((193,), (83,))
In [12]: #Code task 1#
         #Save the 'Name', 'state', and 'Region' columns from the train/test data into names train and names test
         #Then drop those columns from `X train` and `X test`. Use 'inplace=True'
         names_list = ['Name', 'state', 'Region']
         names train = X train[names list]
         names test = X test[names list]
         X train.drop(columns=names list, inplace=True)
         X test.drop(columns=names list, inplace=True)
         X train.shape, X test.shape
```

In [13]: #Code task 2#
 #Check the `dtypes` attribute of `X_train` to verify all features are numeric
 X_train.dtypes

Out[13]: summit elev int64 vertical drop int64 base elev int64 trams int64 fastSixes int64 fastQuads int64 int64 quad triple int64 double int64 surface int64 total chairs int64 Runs float64 float64 TerrainParks LongestRun mi float64 SkiableTerrain ac float64 Snow Making_ac float64 float64 daysOpenLastYear years0pen float64 averageSnowfall float64 projectedDaysOpen float64 NightSkiing ac float64 resorts per state int64 resorts_per_100kcapita float64 resorts per 100ksq mile float64 resort skiable area ac state ratio float64 resort days open state ratio float64 resort terrain park state ratio float64 resort night skiing state ratio float64 total chairs runs ratio float64 total chairs skiable ratio float64 fastQuads runs ratio float64 fastQuads skiable ratio float64 dtype: object

```
In [14]: #Code task 3#
         #Repeat this check for the test split in `X test`
         X test.dtypes
Out[14]: summit elev
                                                   int64
         vertical drop
                                                   int64
                                                   int64
         base elev
         trams
                                                   int64
         fastSixes
                                                   int64
         fastOuads
                                                   int64
         quad
                                                   int64
         triple
                                                   int64
         double
                                                   int64
         surface
                                                   int64
         total chairs
                                                   int64
         Runs
                                                float64
         TerrainParks
                                                float64
                                                float64
         LongestRun mi
         SkiableTerrain ac
                                                float64
         Snow Making ac
                                                float64
                                                float64
         daysOpenLastYear
                                                float64
         years0pen
                                                float64
         averageSnowfall
                                                float64
         projectedDaysOpen
         NightSkiing ac
                                                float64
         resorts_per_state
                                                   int64
         resorts per 100kcapita
                                                float64
         resorts per 100ksq mile
                                                float64
         resort skiable area ac state ratio
                                                float64
         resort days open state ratio
                                                float64
         resort terrain park state ratio
                                                float64
         resort_night_skiing_state_ratio
                                                float64
                                                float64
         total chairs runs ratio
         total chairs skiable ratio
                                                float64
         fastQuads runs ratio
                                                float64
         fastQuads skiable ratio
                                                float64
         dtype: object
```

You have only numeric features in your X now!

4.7 Initial Not-Even-A-Model

A good place to start is to see how good the mean is as a predictor. In other words, what if you simply say your best guess is the average price?

```
In [15]: #Code task 4#
#Calculate the mean of `y_train`
train_mean = y_train.mean()
train_mean
```

Out[15]: 63.811088082901556

sklearn 's DummyRegressor easily does this:

```
In [16]: #Code task 5#
#Fit the dummy regressor on the training data
#Hint, call its `.fit()` method with `X_train` and `y_train` as arguments
#Then print the object's `constant_` attribute and verify it's the same as the mean above
dumb_reg = DummyRegressor(strategy='mean')
dumb_reg.fit(X_train, y_train)
dumb_reg.constant_
```

Out[16]: array([[63.81108808]])

How good is this? How closely does this match, or explain, the actual values? There are many ways of assessing how good one set of values agrees with another, which brings us to the subject of metrics.

4.7.1 Metrics

4.7.1.1 R-squared, or coefficient of determination

One measure is \mathbb{R}^2 , the <u>coefficient of determination (https://en.wikipedia.org/wiki/Coefficient_of_determination)</u>. This is a measure of the proportion of variance in the dependent variable (our ticket price) that is predicted by our "model". The linked Wikipedia articles gives a nice explanation of how negative values can arise. This is frequently a cause of confusion for newcomers who, reasonably, ask how can a squared value be negative?

Recall the mean can be denoted by \bar{y} , where

$$\bar{y} = \frac{1}{n} \sum_{i=1}^{n} y_i$$

and where y_i are the individual values of the dependent variable.

The total sum of squares (error), can be expressed as

$$SS_{tot} = \sum_{i} (y_i - \bar{y})^2$$

The above formula should be familiar as it's simply the variance without the denominator to scale (divide) by the sample size.

The **residual sum** of squares is similarly defined to be

$$SS_{res} = \sum_{i} (y_i - \hat{y})^2$$

where \hat{y} are our predicted values for the depended variable.

The coefficient of determination, \mathbb{R}^2 , here is given by

$$R^2 = 1 - \frac{SS_{res}}{SS_{tot}}$$

Putting it into words, it's one minus the ratio of the residual variance to the original variance. Thus, the baseline model here, which always predicts \bar{y} , should give $R^2 = 0$. A model that perfectly predicts the observed values would have no residual error and so give $R^2 = 1$. Models that do worse than predicting the mean will have increased the sum of squares of residuals and so produce a negative R^2 .

```
In [17]: #Code task 6#
#Calculate the R^2 as defined above
def r_squared(y, ypred):
    """R-squared score.

    Calculate the R-squared, or coefficient of determination, of the input.

    Arguments:
    y -- the observed values
    ypred -- the predicted values
    """

    ybar = np.sum(y) / len(y) #yes, we could use np.mean(y)
    sum_sq_tot = np.sum((y - ybar)**2) #total sum of squares error
    sum_sq_res = np.sum((y - ypred)**2) #residual sum of squares error
    R2 = 1.0 - sum_sq_res / sum_sq_tot
    return R2
```

Make your predictions by creating an array of length the size of the training set with the single value of the mean.

```
In [18]: y_tr_pred_ = train_mean * np.ones(len(y_train))
y_tr_pred_[:5]
```

Out[18]: array([63.81108808, 63.81108808, 63.81108808, 63.81108808])

Remember the sklearn dummy regressor?

```
In [19]: y_tr_pred = dumb_reg.predict(X_train)
y_tr_pred[:5]
```

Out[19]: array([63.81108808, 63.81108808, 63.81108808, 63.81108808])

You can see that <code>DummyRegressor</code> produces exactly the same results and saves you having to mess about broadcasting the mean (or whichever other statistic we used - check out the documentation (https://scikit-learn.org/stable/modules/generated/sklearn.dummy.DummyRegressor.html) to see what's available) to an array of the appropriate length. It also gives you an object with <code>fit()</code> and <code>predict()</code> methods as well so you can use them as conveniently as any other <code>sklearn</code> estimator.

```
In [20]: r_squared(y_train, y_tr_pred)
```

Out[20]: 0.0

Exactly as expected, if you use the average value as your prediction, you get an \mathbb{R}^2 of zero *on our training set*. What if you use this "model" to predict unseen values from the test set? Remember, of course, that your "model" is trained on the training set; you still use the training set mean as your prediction.

Make your predictions by creating an array of length the size of the test set with the single value of the (training) mean.

```
In [21]: y_te_pred = train_mean * np.ones(len(y_test))
r_squared(y_test, y_te_pred)
```

Out[21]: -0.0031235200417913944

Generally, you can expect performance on a test set to be slightly worse than on the training set. As you are getting an \mathbb{R}^2 of zero on the training set, there's nowhere to go but negative!

 R^2 is a common metric, and interpretable in terms of the amount of variance explained, it's less appealing if you want an idea of how "close" your predictions are to the true values. Metrics that summarise the difference between predicted and actual values are *mean absolute error* and *mean squared error*.

4.7.1.2 Mean Absolute Error

In [22]: #Code task 7#

This is very simply the average of the absolute errors:

#Calculate the MAE as defined above

$$MAE = \frac{1}{n} \sum_{i}^{n} |y_i - \hat{y}|$$

```
def mae(y, ypred):
    """Mean absolute error.

    Calculate the mean absolute error of the arguments

    Arguments:
    y -- the observed values
    ypred -- the predicted values
    """
    abs_error = np.abs(y - ypred)
    mae = np.mean(abs_error)
    return mae

In [23]: mae(y_train, y_tr_pred)

Out[23]: 17.923463717146785

In [24]: mae(y_test, y_te_pred)
```

Mean absolute error is arguably the most intuitive of all the metrics, this essentially tells you that, on average, you might expect to be off by around \$19 if you guessed ticket price based on an average of known values.

4.7.1.3 Mean Squared Error

Out[24]: 19.136142081278486

Another common metric (and an important one internally for optimizing machine learning models) is the mean squared error. This is simply the average of the square of the errors:

$$MSE = \frac{1}{n} \sum_{i}^{n} (y_i - \hat{y})^2$$

```
In [25]: #Code task 8#
#Calculate the MSE as defined above
def mse(y, ypred):
    """Mean square error.

    Calculate the mean square error of the arguments

    Arguments:
    y -- the observed values
    ypred -- the predicted values
    """

    sq_error = (y - ypred)**2
    mse = np.mean(sq_error)
    return mse
```

```
In [26]: mse(y_train, y_tr_pred)
Out[26]: 614.1334096969057
```

```
In [27]: mse(y_test, y_te_pred)
```

Out[27]: 581.4365441953481

So here, you get a slightly better MSE on the test set than you did on the train set. And what does a squared error mean anyway? To convert this back to our measurement space, we often take the square root, to form the *root mean square error* thus:

```
In [28]: np.sqrt([mse(y_train, y_tr_pred), mse(y_test, y_te_pred)])
Out[28]: array([24.78171523, 24.11299534])
```

4.7.2 sklearn metrics

Functions are good, but you don't want to have to define functions every time we want to assess performance. sklearn.metrics provides many commonly used metrics, included the ones above.

4.7.2.0.1 R-squared

```
In [29]: r2_score(y_train, y_tr_pred), r2_score(y_test, y_te_pred)
Out[29]: (0.0, -0.0031235200417913944)
```

4.7.2.0.2 Mean absolute error

```
In [30]: mean_absolute_error(y_train, y_tr_pred), mean_absolute_error(y_test, y_te_pred)
Out[30]: (17.92346371714677, 19.136142081278486)
```

4.7.2.0.3 Mean squared error

```
In [31]: mean_squared_error(y_train, y_tr_pred), mean_squared_error(y_test, y_te_pred)
Out[31]: (614.1334096969046, 581.4365441953483)
```

4.7.3 Note On Calculating Metrics

When calling functions to calculate metrics, it is important to take care in the order of the arguments. Two of the metrics above actually don't care if the arguments are reversed; one does. Which one cares?

In a Jupyter code cell, running r2_score? will bring up the docstring for the function, and r2_score?? will bring up the actual code of the function! Try them and compare the source for sklearn 's function with yours. Feel free to explore what happens when you reverse the order of the arguments and compare behaviour of sklearn 's function and yours.

```
In [34]: # train set - using our homebrew function
    # correct order, incorrect order
    r_squared(y_train, y_tr_pred), r_squared(y_tr_pred, y_train)

Out[34]: (0.0, -3.041041349306602e+30)

In [35]: # test set - using our homebrew function
    # correct order, incorrect order
    r_squared(y_test, y_te_pred), r_squared(y_te_pred, y_test)

    <i i python-input-17-fc28c3263e5e>:15: RuntimeWarning: divide by zero encountered in double_scalars
    R2 = 1.0 - sum_sq_res / sum_sq_tot

Out[35]: (-0.0031235200417913944, -inf)
```

You can get very different results swapping the argument order. It's worth highlighting this because data scientists do this too much in the real world! Don't be one of them! Frequently the argument order doesn't matter, but it will bite you when you do it with a function that does care. It's sloppy, bad practice and if you don't make a habit of putting arguments in the right order, you will forget!

Remember:

- · argument order matters,
- · check function syntax with func? in a code cell

4.8 Initial Models

4.8.1 Imputing missing feature (predictor) values

Recall when performing EDA, you imputed (filled in) some missing values in pandas. You did this judiciously for exploratory/visualization purposes. You left many missing values in the data. You can impute missing values using scikit-learn, but note that you should learn values to impute from a train split and apply that to the test split to then assess how well your imputation worked.

4.8.1.1 Impute missing values with median

There's missing values. Recall from your data exploration that many distributions were skewed. Your first thought might be to impute missing values using the median.

4.8.1.1.1 Learn the values to impute from the train set

```
In [36]: # These are the values we'll use to fill in any missing values
         X defaults median = X train.median()
         X defaults median
Out[36]: summit elev
                                                2215.000000
         vertical drop
                                                 750.000000
         base elev
                                                1300.000000
         trams
                                                   0.000000
         fastSixes
                                                   0.000000
         fastQuads
                                                   0.000000
         quad
                                                   1.000000
         triple
                                                   1.000000
         double
                                                   1.000000
         surface
                                                   2.000000
         total chairs
                                                   7.000000
         Runs
                                                  28.000000
         TerrainParks
                                                   2.000000
         LongestRun mi
                                                   1.000000
         SkiableTerrain ac
                                                 170.000000
         Snow Making_ac
                                                  96.500000
         daysOpenLastYear
                                                 109.000000
         years0pen
                                                  57.000000
         averageSnowfall
                                                 120.000000
         projectedDaysOpen
                                                 115.000000
         NightSkiing_ac
                                                  70.000000
         resorts per state
                                                  15.000000
         resorts per 100kcapita
                                                   0.248243
         resorts per 100ksq mile
                                                  22.902162
         resort skiable area ac state ratio
                                                   0.051458
         resort days open state ratio
                                                   0.071225
         resort terrain park state ratio
                                                   0.069444
         resort night skiing state ratio
                                                   0.077081
         total chairs runs ratio
                                                   0.200000
         total chairs skiable ratio
                                                   0.040323
         fastQuads runs ratio
                                                   0.000000
         fastQuads skiable ratio
                                                   0.000000
         dtype: float64
```

4.8.1.1.2 Apply the imputation to both train and test splits

```
In [37]: #Code task 9#
#Call `X_train` and `X_test`'s `fillna()` method, passing `X_defaults_median` as the values to use
#Assign the results to `X_tr` and `X_te`, respectively
X_tr = X_train.fillna(X_defaults_median)
X_te = X_test.fillna(X_defaults_median)
```

4.8.1.1.3 Scale the data

As you have features measured in many different units, with numbers that vary by orders of magnitude, start off by scaling them to put them all on a consistent scale. The <u>StandardScaler (https://scikit-learn.org/stable/modules/generated/sklearn.preprocessing.StandardScaler.html)</u> scales each feature to zero mean and unit variance.

```
In [38]: #Code task 10#
#Call the StandardScaler`s fit method on `X_tr` to fit the scaler
#then use it's `transform()` method to apply the scaling to both the train and test split
#data (`X_tr` and `X_te`), naming the results `X_tr_scaled` and `X_te_scaled`, respectively
scaler = StandardScaler()
scaler.fit(X_tr)
X_tr_scaled = scaler.transform(X_tr)
X_te_scaled = scaler.transform(X_te)
```

4.8.1.1.4 Train the model on the train split

```
In [39]: lm = LinearRegression().fit(X_tr_scaled, y_train)
```

4.8.1.1.5 Make predictions using the model on both train and test splits

```
In [40]: #Code task 11#
#Call the `predict()` method of the model (`lm`) on both the (scaled) train and test data
#Assign the predictions to `y_tr_pred` and `y_te_pred`, respectively
y_tr_pred = lm.predict(X_tr_scaled)
y_te_pred = lm.predict(X_te_scaled)
```

4.8.1.1.6 Assess model performance

```
In [41]: # r^2 - train, test
    median_r2 = r2_score(y_train, y_tr_pred), r2_score(y_test, y_te_pred)
    median_r2
```

Out[41]: (0.8177988515690604, 0.7209725843435142)

Recall that you estimated ticket price by simply using a known average. As expected, this produced an \mathbb{R}^2 of zero for both the training and test set, because \mathbb{R}^2 tells us how much of the variance you're explaining beyond that of using just the mean, and you were using just the mean. Here we see that our simple linear regression model explains over 80% of the variance on the train set and over 70% on the test set. Clearly you are onto something, although **the much lower value for the test set suggests you're overfitting somewhat**. This isn't a surprise as you've made no effort to select a parsimonious set of features or deal with multicollinearity in our data.

```
In [42]: #Code task 12#
#Now calculate the mean absolute error scores using `sklearn`'s `mean_absolute_error` function
# as we did above for R^2
# MAE - train, test
median_mae = mean_absolute_error(y_train, y_tr_pred), mean_absolute_error(y_test, y_te_pred)
median_mae
```

Out[42]: (8.547850301825427, 9.40702011858132)

Using this model, then, on average you'd expect to estimate a ticket price within \$9 or so of the real price. This is much, much better than the \$19 from just guessing using the average. There may be something to this machine learning lark after all!

```
In [43]: #Code task 13#
#And also do the same using `sklearn`'s `mean_squared_error`
# MSE - train, test
median_mse = mean_squared_error(y_train, y_tr_pred), mean_squared_error(y_test, y_te_pred)
median_mse
```

Out[43]: (111.89581253658478, 161.73156451192284)

4.8.1.2 Impute missing values with the mean

You chose to use the median for filling missing values because of the skew of many of our predictor feature distributions. What if you wanted to try something else, such as the mean?

4.8.1.2.1 Learn the values to impute from the train set

```
In [44]:
         #Code task 14#
         #As we did for the median above, calculate mean values for imputing missing values
         # These are the values we'll use to fill in any missing values
         X defaults mean = X train.mean()
         X defaults mean
Out[44]: summit elev
                                                4074.554404
         vertical drop
                                                1043.196891
         base elev
                                                3020.512953
         trams
                                                   0.103627
         fastSixes
                                                   0.072539
         fastOuads
                                                   0.673575
                                                   1.010363
         quad
         triple
                                                   1.440415
         double
                                                   1.813472
         surface
                                                   2.497409
                                                   7.611399
         total chairs
                                                  41.188482
         Runs
         TerrainParks
                                                   2.434783
                                                   1.293122
         LongestRun mi
         SkiableTerrain ac
                                                 448.785340
         Snow Making ac
                                                 129.601190
         daysOpenLastYear
                                                 110.100629
         years0pen
                                                  56.559585
         averageSnowfall
                                                 162.310160
         projectedDaysOpen
                                                 115.920245
         NightSkiing ac
                                                  86.384615
         resorts_per_state
                                                  16.264249
         resorts_per_100kcapita
                                                   0.424802
         resorts per 100ksq mile
                                                  40.957785
         resort skiable area ac state ratio
                                                   0.097205
         resort days open state ratio
                                                   0.126014
         resort terrain park state ratio
                                                   0.116022
         resort night skiing state ratio
                                                   0.155024
         total chairs runs ratio
                                                   0.271441
         total chairs skiable ratio
                                                   0.070483
         fastQuads runs ratio
                                                   0.010401
         fastQuads skiable ratio
                                                   0.001633
         dtype: float64
```

By eye, you can immediately tell that your replacement values are much higher than those from using the median.

4.8.1.2.2 Apply the imputation to both train and test splits

. . .

```
In [45]: X_tr = X_train.fillna(X_defaults_mean)
X_te = X_test.fillna(X_defaults_mean)
```

4.8.1.2.3 Scale the data

```
In [46]: scaler = StandardScaler()
    scaler.fit(X_tr)
    X_tr_scaled = scaler.transform(X_tr)
    X_te_scaled = scaler.transform(X_te)
```

4.8.1.2.4 Train the model on the train split

```
In [47]: lm = LinearRegression().fit(X_tr_scaled, y_train)
```

4.8.1.2.5 Make predictions using the model on both train and test splits

```
In [48]: y_tr_pred = lm.predict(X_tr_scaled)
y_te_pred = lm.predict(X_te_scaled)
```

4.8.1.2.6 Assess model performance

Out[51]: (112.37695054778276, 164.3926930952436)

```
In [49]: r2_score(y_train, y_tr_pred), r2_score(y_test, y_te_pred)
Out[49]: (0.8170154093990025, 0.716381471695996)
In [50]: mean_absolute_error(y_train, y_tr_pred), mean_absolute_error(y_test, y_te_pred)
Out[50]: (8.536884040670973, 9.416375625789271)
In [51]: mean_squared_error(y_train, y_tr_pred), mean_squared_error(y_test, y_te_pred)
```

These results don't seem very different to when you used the median for imputing missing values. Perhaps it doesn't make much difference here. Maybe your overtraining dominates. Maybe other feature transformations, such as taking the log, would help. You could try with just a subset of features rather than using all of them as inputs.

To perform the median/mean comparison, you copied and pasted a lot of code just to change the function for imputing missing values. It would make more sense to write a function that performed the sequence of steps:

- 1. impute missing values
- 2. scale the features
- 3. train a model
- 4. calculate model performance

But these are common steps and sklearn provides something much better than writing custom functions.

4.8.2 Pipelines

One of the most important and useful components of sklearn is the <u>pipeline (https://scikit-learn.org/stable/modules/generated/sklearn.pipeline.Pipeline.html)</u>. In place of panda 's fillna DataFrame method, there is sklearn 's SimpleImputer . Remember the first linear model above performed the steps:

- 1. replace missing values with the median for each feature
- 2. scale the data to zero mean and unit variance
- 3. train a linear regression model

and all these steps were trained on the train split and then applied to the test split for assessment.

The pipeline below defines exactly those same steps. Crucially, the resultant Pipeline object has a fit() method and a predict() method, just like the LinearRegression() object itself. Just as you might create a linear regression model and train it with .fit() and predict with .predict(), you can wrap the entire process of imputing and feature scaling and regression in a single object you can train with .fit() and predict with .predict(). And that's basically a pipeline: a model on steroids.

4.8.2.1 Define the pipeline

Out[53]: sklearn.pipeline.Pipeline

```
In [54]: hasattr(pipe, 'fit'), hasattr(pipe, 'predict')
Out[54]: (True, True)
```

4.8.2.2 Fit the pipeline

Here, a single call to the pipeline's fit() method combines the steps of learning the imputation (determining what values to use to fill the missing ones), the scaling (determining the mean to subtract and the variance to divide by), and then training the model. It does this all in the one call with the training data as arguments.

4.8.2.3 Make predictions on the train and test sets

('linearregression', LinearRegression())])

```
In [56]: y_tr_pred = pipe.predict(X_train)
y_te_pred = pipe.predict(X_test)
```

4.8.2.4 Assess performance

```
In [57]: r2_score(y_train, y_tr_pred), r2_score(y_test, y_te_pred)
```

Out[57]: (0.8177988515690604, 0.7209725843435142)

And compare with your earlier (non-pipeline) result:

```
In [58]: median_r2
Out[58]: (0.8177988515690604, 0.7209725843435142)
```

```
In [59]: mean_absolute_error(y_train, y_tr_pred), mean_absolute_error(y_test, y_te_pred)
Out[59]: (8.547850301825427, 9.40702011858132)

Compare with your earlier result:

In [60]: median_mae
Out[60]: (8.547850301825427, 9.40702011858132)
In [61]: mean_squared_error(y_train, y_tr_pred), mean_squared_error(y_test, y_te_pred)
Out[61]: (111.89581253658478, 161.73156451192284)

Compare with your earlier result:

In [62]: median_mse
Out[62]: (111.89581253658478, 161.73156451192284)
```

These results confirm the pipeline is doing exactly what's expected, and results are identical to your earlier steps. This allows you to move faster but with confidence.

4.9 Refining The Linear Model

You suspected the model was overfitting. This is no real surprise given the number of features you blindly used. It's likely a judicious subset of features would generalize better. sklearn has a number of feature selection functions available. The one you'll use here is SelectKBest which, as you might guess, selects the k best features. You can read about SelectKBest <a href="https://scikit-learn.org/stable/modules/generated/sklearn.feature-selection.SelectKBest.html#sklearn.feature-selection.SelectKBest). f regression is just the

score function (https://scikit-learn.org/stable/modules/generated/sklearn.feature_selection.f_regression.html#sklearn.feature_selection.html#sklearn.featu

4.9.1 Define the pipeline

Redefine your pipeline to include this feature selection step:

4.9.2 Fit the pipeline

4.9.3 Assess performance on the train and test set

```
In [65]: y_tr_pred = pipe.predict(X_train)
y_te_pred = pipe.predict(X_test)

In [66]: r2_score(y_train, y_tr_pred), r2_score(y_test, y_te_pred)

Out[66]: (0.7674914326052744, 0.6259877354190833)

In [67]: mean_absolute_error(y_train, y_tr_pred), mean_absolute_error(y_test, y_te_pred)

Out[67]: (9.501495079727484, 11.201830190332057)
```

This has made things worse! Clearly selecting a subset of features has an impact on performance. SelectKBest defaults to k=10. You've just seen that 10 is worse than using all features. What is the best k? You could create a new pipeline with a different value of k:

4.9.4 Define a new pipeline to select a different number of features

```
In [68]: #Code task 17#
    #Modify the `SelectKBest` step to use a value of 15 for k
    pipe15 = make_pipeline(
        SimpleImputer(strategy='median'),
        StandardScaler(),
        SelectKBest(f_regression, k=15),
        LinearRegression()
)
```

4.9.5 Fit the pipeline

4.9.6 Assess performance on train and test data

```
In [70]: y_tr_pred = pipe15.predict(X_train)
y_te_pred = pipe15.predict(X_test)

In [71]: r2_score(y_train, y_tr_pred), r2_score(y_test, y_te_pred)

Out[71]: (0.7924096060483825, 0.6376199973170795)

In [72]: mean_absolute_error(y_train, y_tr_pred), mean_absolute_error(y_test, y_te_pred)

Out[72]: (9.211767769307116, 10.488246867294356)
```

You could keep going, trying different values of k, training a model, measuring performance on the test set, and then picking the model with the best test set performance. There's a fundamental problem with this approach: *you're tuning the model to the arbitrary test set*! If you continue this way you'll end up with a model works well on the particular quirks of our test set *but fails to generalize to new data*. The whole point of keeping a test set is for it to be a set of that new data, to check how well our model might perform on data it hasn't seen.

The way around this is a technique called *cross-validation*. You partition the training set into k folds, train our model on k-1 of those folds, and calculate performance on the fold not used in training. This procedure then cycles through k times with a different fold held back each time. Thus you end up building k models on k sets of data with k estimates of how the model performs on unseen data but without having to touch the test set.

4.9.7 Assessing performance using cross-validation

```
In [73]: cv_results = cross_validate(pipe15, X_train, y_train, cv=5)
In [74]: cv_scores = cv_results['test_score']
cv_scores
Out[74]: array([0.63760862, 0.72831381, 0.74443537, 0.5487915, 0.50441472])
```

Without using the same random state for initializing the CV folds, your actual numbers will be different.

```
In [75]: np.mean(cv_scores), np.std(cv_scores)
Out[75]: (0.6327128053007867, 0.09502487849877672)
```

These results highlight that assessing model performance in inherently open to variability. You'll get different results depending on the quirks of which points are in which fold. An advantage of this is that you can also obtain an estimate of the variability, or uncertainty, in your performance estimate.

```
In [76]: np.round((np.mean(cv_scores) - 2 * np.std(cv_scores), np.mean(cv_scores) + 2 * np.std(cv_scores)), 2)
Out[76]: array([0.44, 0.82])
```

4.9.8 Hyperparameter search using GridSearchCV

Pulling the above together, we have:

- a pipeline that
 - imputes missing values
 - scales the data
 - selects the k best features
 - trains a linear regression model
- a technique (cross-validation) for estimating model performance

Now you want to use cross-validation for multiple values of k and use cross-validation to pick the value of k that gives the best performance.

make_pipeline automatically names each step as the lowercase name of the step and the parameters of the step are then accessed by appending a double underscore followed by the parameter name. You know the name of the step will be 'selectkbest' and you know the parameter is 'k'.

You can also list the names of all the parameters in a pipeline like this:

```
In [77]:
         #Code task 18#
         #Call `pipe`'s `get params()` method to get a dict of available parameters and print their names
         #using dict's `keys()` method
         pipe.get_params('keys()')
Out[77]: {'memory': None,
           'steps': [('simpleimputer', SimpleImputer(strategy='median')),
            ('standardscaler', StandardScaler()),
            ('selectkbest',
            SelectKBest(score_func=<function f_regression at 0x0000021089ED98B0>)),
            ('linearregression', LinearRegression())],
           'verbose': False,
           'simpleimputer': SimpleImputer(strategy='median'),
           'standardscaler': StandardScaler(),
           'selectkbest': SelectKBest(score func=<function f regression at 0x0000021089ED98B0>),
           'linearregression': LinearRegression(),
           'simpleimputer__add_indicator': False,
           'simpleimputer__copy': True,
           'simpleimputer__fill_value': None,
           'simpleimputer__missing_values': nan,
           'simpleimputer strategy': 'median',
           'simpleimputer verbose': 0,
           'standardscaler__copy': True,
           'standardscaler__with_mean': True,
           'standardscaler__with_std': True,
           'selectkbest k': 10,
           'selectkbest__score_func': <function sklearn.feature_selection._univariate_selection.f_regression(X, y, *, center=True)
           'linearregression copy X': True,
           'linearregression fit intercept': True,
           'linearregression__n_jobs': None,
           'linearregression__normalize': False}
```

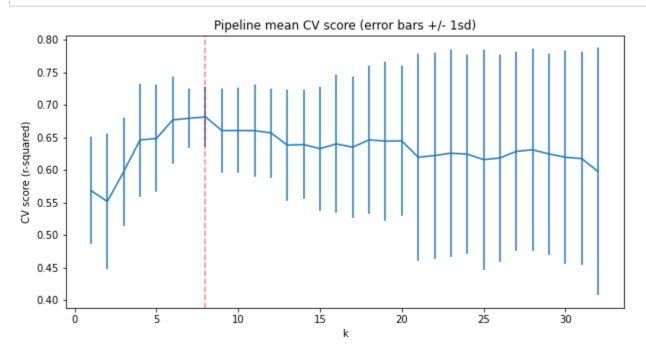
The above can be particularly useful as your pipelines becomes more complex (you can even nest pipelines within pipelines).

```
In [78]: k = [k+1 for k in range(len(X_train.columns))]
    grid_params = {'selectkbest__k': k}
```

Now you have a range of k to investigate. Is 1 feature best? 2? 3? 4? All of them? You could write a for loop and iterate over each possible value, doing all the housekeeping ourselves to track the best value of k. But this is a common task so there's a built in function in sklearn. This is GridSearchCV (https://scikit-learn.org/stable/modules/generated/sklearn.model_selection.GridSearchCV.html). This takes the pipeline object, in fact it takes anything with a fttps://scikit-learn.org/stable/modules/generated/sklearn.model_selection.GridSearchCV.html). This takes the pipeline object, in fact it takes anything with a fttps://scikit-learn.org/stable/modules/generated/sklearn.model_selection.GridSearchCV.html). This takes the pipeline object, in fact it takes anything with a fttps://scikit-learn.org/stable/modules/generated/sklearn.model_selection.GridSearchCV.html). This takes the pipeline object, in fact it takes anything with a fttps://scikit-learn.org/stable/modules/generated/sklearn.model_selection.GridSearchCV.html). This takes the pipeline object, in fact it takes anything with a fttps://scikit-learn.org/stable/modules/generated/sklearn.model_selection.GridSearchCV.html). This takes the pipeline object, in fact it takes anything a selection or regressor object itself directly passed into GridSearchCV. The other key input is the parameters and values

```
In [79]: | lr grid cv = GridSearchCV(pipe, param grid=grid params, cv=5, n jobs=-1)
In [80]: | lr grid cv.fit(X train, y train)
Out[80]: GridSearchCV(cv=5,
                       estimator=Pipeline(steps=[('simpleimputer',
                                                  SimpleImputer(strategy='median')),
                                                 ('standardscaler', StandardScaler()),
                                                 ('selectkbest',
                                                  SelectKBest(score func=<function f regression at 0x0000021089ED98B0>)),
                                                 ('linearregression',
                                                  LinearRegression())]),
                       n jobs=-1,
                       param_grid={'selectkbest__k': [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11,
                                                      12, 13, 14, 15, 16, 17, 18, 19, 20,
                                                      21, 22, 23, 24, 25, 26, 27, 28, 29,
                                                      30, ... 1})
In [81]: | score mean = lr grid cv.cv results ['mean test score']
         score std = lr grid cv.cv results ['std test score']
         cv k = [k for k in lr grid cv.cv results ['param selectkbest k']]
In [82]: #Code task 19#
         #Print the `best params ` attribute of `lr grid cv`
         lr_grid_cv.best_params_
Out[82]: {'selectkbest k': 8}
```

In [83]: #Code task 20# #Assign the value of k from the above dict of `best_params_` and assign it to `best_k` best_k = lr_grid_cv.best_params_['selectkbest_k'] plt.subplots(figsize=(10, 5)) plt.errorbar(cv_k, score_mean, yerr=score_std) plt.axvline(x=best_k, c='r', ls='--', alpha=.5) plt.xlabel('k') plt.ylabel('CV score (r-squared)') plt.title('Pipeline mean CV score (error bars +/- 1sd)');



The above suggests a good value for k is 8. There was an initial rapid increase with k, followed by a slow decline. Also noticeable is the variance of the results greatly increase above k=8. As you increasingly overfit, expect greater swings in performance as different points move in and out of the train/test folds.

Which features were most useful? Step into your best model, shown below. Starting with the fitted grid search object, you get the best estimator, then the named step 'selectkbest', for which you can its <code>get_support()</code> method for a logical mask of the features selected.

```
In [84]: selected = lr_grid_cv.best_estimator_.named_steps.selectkbest.get_support()
```

Similarly, instead of using the 'selectkbest' named step, you can access the named step for the linear regression model and, from that, grab the model coefficients via its coef_ attribute:

```
In [85]: #Code task 21#
#Get the linear model coefficients from the `coef_` attribute and store in `coefs`,
#get the matching feature names from the column names of the dataframe,
#and display the results as a pandas Series with `coefs` as the values and `features` as the index,
#sorting the values in descending order
coefs = lr_grid_cv.best_estimator_.named_steps.linearregression.coef_
features = X_train.columns[selected]
pd.Series(coefs, index=features).sort_values(ascending=False)
```

```
Out[85]: vertical_drop
                               10.767857
                                6.290074
         Snow Making ac
                                5.794156
         total chairs
         fastQuads
                                5.745626
                                5.370555
          Runs
         LongestRun_mi
                                0.181814
         trams
                               -4.142024
          SkiableTerrain ac
                               -5.249780
```

dtype: float64

These results suggest that vertical drop is your biggest positive feature. This makes intuitive sense and is consistent with what you saw during the EDA work. Also, you see the area covered by snow making equipment is a strong positive as well. People like guaranteed skiing! The skiable terrain area is negatively associated with ticket price! This seems odd. People will pay less for larger resorts? There could be all manner of reasons for this. It could be an effect whereby larger resorts can host more visitors at any one time and so can charge less per ticket. As has been mentioned previously, the data are missing information about visitor numbers. Bear in mind, the coefficient for skiable terrain is negative *for this model*. For example, if you kept the total number of chairs and fastQuads constant, but increased the skiable terrain extent, you might imagine the resort is worse off because the chairlift capacity is stretched thinner.

4.10 Random Forest Model

A model that can work very well in a lot of cases is the random forest. For regression, this is provided by sklearn 's RandomForestRegressor class.

Time to stop the bad practice of repeatedly checking performance on the test split. Instead, go straight from defining the pipeline to assessing performance using cross-validation. cross_validate will perform the fitting as part of the process. This uses the default settings for the random forest so you'll then proceed to investigate some different hyperparameters.

4.10.1 Define the pipeline

4.10.2 Fit and assess performance using cross-validation

```
In [87]: #Code task 23#
    #Call `cross_validate` to estimate the pipeline's performance.
    #Pass it the random forest pipe object, `X_train` and `y_train`,
    #and get it to use 5-fold cross-validation
    rf_default_cv_results = cross_validate(RF_pipe, X_train, y_train, cv=5)

In [88]: rf_cv_scores = rf_default_cv_results['test_score']
    rf_cv_scores

Out[88]: array([0.69249204, 0.78061953, 0.77546915, 0.62190924, 0.61742339])

In [89]: np.mean(rf_cv_scores), np.std(rf_cv_scores)

Out[89]: (0.6975826707112506, 0.07090742940774528)
```

4.10.3 Hyperparameter search using GridSearchCV

Random forest has a number of hyperparameters that can be explored, however here you'll limit yourselves to exploring some different values for the number of trees. You'll try it with and without feature scaling, and try both the mean and median as strategies for imputing missing values.

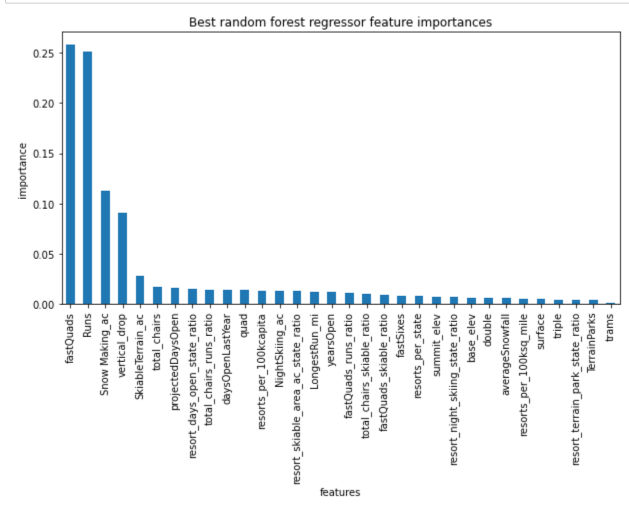
```
In [90]: n est = [int(n) for n in np.logspace(start=1, stop=3, num=20)]
         grid params = {
                  'randomforestregressor n estimators': n est,
                  'standardscaler': [StandardScaler(), None],
                  'simpleimputer strategy': ['mean', 'median']
         grid params
Out[90]: {'randomforestregressor n estimators': [10,
           12,
           16,
           20,
           26,
           33,
           42,
           54,
           69,
           88,
           112,
           143,
           183,
           233,
           297,
           379,
           483,
           615,
           784,
           1000],
           'standardscaler': [StandardScaler(), None],
           'simpleimputer__strategy': ['mean', 'median']}
In [91]:
         #Code task 24#
         #Call `GridSearchCV` with the random forest pipeline, passing in the above `grid params`
         #dict for parameters to evaluate, 5-fold cross-validation, and all available CPU cores (if desired)
         rf grid cv = GridSearchCV(RF pipe, param grid=grid params, cv=5, n jobs=-1)
```

```
In [92]: #Code task 25#
         #Now call the `GridSearchCV`'s `fit()` method with `X train` and `y train` as arguments
         #to actually start the grid search. This may take a minute or two.
         rf grid cv.fit(X train, y train)
Out[92]: GridSearchCV(cv=5,
                       estimator=Pipeline(steps=[('simpleimputer',
                                                  SimpleImputer(strategy='median')),
                                                  ('standardscaler', StandardScaler()),
                                                  ('randomforestregressor',
                                                  RandomForestRegressor(random state=47))]),
                       n jobs=-1,
                       param grid={'randomforestregressor n estimators': [10, 12, 16, 20,
                                                                            26, 33, 42, 54,
                                                                            69, 88, 112,
                                                                            143, 183, 233,
                                                                            297, 379, 483,
                                                                            615, 784,
                                                                            1000],
                                   'simpleimputer strategy': ['mean', 'median'],
                                    'standardscaler': [StandardScaler(), None]})
In [93]: #Code task 26#
         #Print the best params (`best_params_` attribute) from the grid search
         rf grid cv.best params
Out[93]: {'randomforestregressor n estimators': 69,
           'simpleimputer strategy': 'median',
           'standardscaler': None}
         It looks like imputing with the median helps, but scaling the features doesn't.
In [94]: rf best cv results = cross validate(rf grid cv.best estimator, X train, y train, cv=5)
         rf best scores = rf best cv results['test score']
         rf best scores
Out[94]: array([0.6951357 , 0.79430697, 0.77170917, 0.62254707, 0.66499334])
In [95]: np.mean(rf best scores), np.std(rf best scores)
Out[95]: (0.7097384501425082, 0.06451341966873386)
```

You've marginally improved upon the default CV results. Random forest has many more hyperparameters you could tune, but we won't dive into that

here.

```
In [96]: #Code task 27#
    #Plot a barplot of the random forest's feature importances,
    #assigning the `feature_importances_` attribute of
    #`rf_grid_cv.best_estimator_.named_steps.randomforestregressor` to the name `imps` to then
    #create a pandas Series object of the feature importances, with the index given by the
    #training data column names, sorting the values in descending order
    plt.subplots(figsize=(10, 5))
    imps = rf_grid_cv.best_estimator_.named_steps.randomforestregressor.feature_importances_
    rf_feat_imps = pd.Series(imps, index=X_train.columns).sort_values(ascending=False)
    rf_feat_imps.plot(kind='bar')
    plt.xlabel('features')
    plt.ylabel('importance')
    plt.title('Best_random_forest_regressor_feature_importances');
```



Encouragingly, the dominant top four features are in common with your linear model:

- fastQuads
- Runs
- · Snow Making ac
- · vertical drop

4.11 Final Model Selection

Time to select your final model to use for further business modeling! It would be good to revisit the above model selection; there is undoubtedly more that could be done to explore possible hyperparameters. It would also be worthwhile to investigate removing the least useful features. Gathering or calculating, and storing, features adds business cost and dependencies, so if features genuinely are not needed they should be removed. Building a simpler model with fewer features can also have the advantage of being easier to sell (and/or explain) to stakeholders. Certainly there seem to be four strong features here and so a model using only those would probably work well. However, you want to explore some different scenarios where other features vary so keep the fuller model for now. The business is waiting for this model and you have something that you have confidence in to be much better than guessing with the average price.

Or, rather, you have two "somethings". You built a **best linear model** and a **best random forest model**. You need to finally choose between them. You can calculate the mean absolute error using cross-validation. Although cross-validate defaults to the R^2 metric for scoring (https://scikit-learn.org/stable/modules/model_evaluation.html#scoring) regression, you can specify the mean absolute error as an alternative via the scoring parameter.

4.11.1 Linear regression model performance

4.11.2 Random forest regression model performance

4.11.3 Conclusion

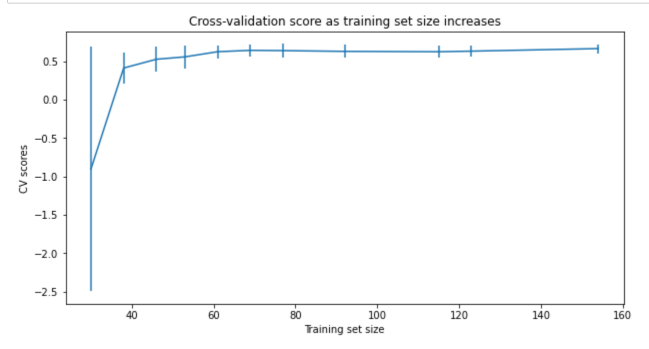
The random forest model has a lower cross-validation mean absolute error by almost \$1. It also exhibits less variability. Verifying performance on the test set produces performance consistent with the cross-validation results.

4.12 Data quantity assessment

Finally, you need to advise the business whether it needs to undertake further data collection. Would more data be useful? We're often led to believe more data is always good, but gathering data invariably has a cost associated with it. Assess this trade off by seeing how performance varies with differing data set sizes. The learning curve function does this conveniently.

```
In [103]: fractions = [.2, .25, .3, .35, .4, .45, .5, .6, .75, .8, 1.0]
    train_size, train_scores, test_scores = learning_curve(pipe, X_train, y_train, train_sizes=fractions)
    train_scores_mean = np.mean(train_scores, axis=1)
    train_scores_std = np.std(train_scores, axis=1)
    test_scores_mean = np.mean(test_scores, axis=1)
    test_scores_std = np.std(test_scores, axis=1)
```

```
In [104]: plt.subplots(figsize=(10, 5))
    plt.errorbar(train_size, test_scores_mean, yerr=test_scores_std)
    plt.xlabel('Training set size')
    plt.ylabel('CV scores')
    plt.title('Cross-validation score as training set size increases');
```



This shows that you seem to have plenty of data. There's an initial rapid improvement in model scores as one would expect, but it's essentially levelled off by around a sample size of 40-50.

4.13 Save best model object from pipeline

```
In [105]: #Code task 28#
          #This may not be "production grade ML deployment" practice, but adding some basic
          #information to your saved models can save your bacon in development.
          #Just what version model have you just loaded to reuse? What version of `sklearn`
          #created it? When did you make it?
          #Assign the pandas version number (`pd. version `) to the `pandas version` attribute,
          #the numpy version (`np. version `) to the `numpy version` attribute,
          #the sklearn version (`sklearn version`) to the `sklearn version` attribute,
          #and the current datetime (`datetime.datetime.now()`) to the `build datetime` attribute
          #Let's call this model version '1.0'
          best model = rf grid cv.best estimator
          best model.version = '1.0'
          best model.pandas version = pd. version
          best_model.numpy_version = np.__version__
          best model.sklearn version = sklearn version
          best model.X columns = [col for col in X train.columns]
          best model.build datetime = datetime.datetime.now()
In [106]: # save the model
          modelpath = '../models'
          save file(best model, 'ski resort pricing model.pkl', modelpath)
```

4.14 Summary

A file already exists with this name.

Writing file. "../models\ski_resort_pricing model.pkl"

Do you want to overwrite? (Y/N)y

Q: 1 Write a summary of the work in this notebook. Capture the fact that you gained a baseline idea of performance by simply taking the average price and how well that did. Then highlight that you built a linear model and the features that found. Comment on the estimate of its performance from cross-validation and whether its performance on the test split was consistent with this estimate. Also highlight that a random forest regressor was tried, what preprocessing steps were found to be best, and again what its estimated performance via cross-validation was and whether its performance on the test set was consistent with that. State which model you have decided to use going forwards and why. This summary should provide a quick overview for someone wanting to know quickly why the given model was chosen for the next part of the business problem to help guide important business decisions.

A: 1 I imported more than a dozen necessary libraries, loaded my dataset file from step 3, and then copied off my target resort data to save it for later. Next I used a sklearn method to divide the data rows into a training set and a smaller testing set (including some shuffling). I also saved off the non-

numeric columns, then removed these from the data to support use of several advanced functions available to handle the remaining numeric data.

I tried evaluating how good the training data's average was as a predictor, which is also fairly easily obtained from the DummyRegressor function. I reviewed definitions for **mean**, **total sum of squares**, and **residual sum of squares**. The last two lead to a derivation for **coefficient of determination**. I reviewed a python function definition for that last one, and verified that running it against the training set, and result was as expected zero. Against the test set it was slightly worse, also as expected.

Then I looked at a pair of additional metrics used to evaluate and summarize the difference between predicted and actual values. **Mean Absolute Error** was defined and implemented in Python code, and then used to evaluate the training data set and the test data set. The result suggests that using overall average price as an estimate will generally be off by about \$19. **Mean Square Error** is another metric defined and implemented in Python code. I used that similarly to evaluate the training set and then looked at a variation involving taking the square root of the result as the final step. This "RMS" value does sound familiar from something similar that I studied back during Electrical Engineering courses, related to alternating current signals.

I confirmed that all of these metrics are easily available from sklearn.metrics package, and others; and that the corresponding results matched in all cases. There was a quick word about keeping the sequence of function call arguments in mind, and a bit about learning how to properly use these functions.

I tried plugging the data set's missing values with corresponding median values via fillna(), and looked at how that worked out via the fill, scale, train, predict cycle. I used standard functions from the sklearn.metrics package. And I tried using 'median' rather than 'mean' as fill values but they worked out pretty similarly. Finally I tried using pipeline set of methods, featuring SimpleImputer (which takes on the role of fillna()).