# Machine Learning with Python: Foundations

https://www.linkedin.com/learning/machine-learning-with-python-foundations

#### Intro

- Computers receive 2 things
  - An input or data
  - A set of instructions on what to do
- ML is where computers don't get an explicit set of instructions
  - We give an input and expected output
  - Computer figures out the instructions
- Focus of ML is to predict

## **Data Collection**

#### Considerations

- Objective is to identify and gather data to be used
- There are 5 key considerations when it comes to data
- The first is accuracy
  - Accurate data is needed for a good model
  - For supervised learning, this is especially crucial
    - Bad data = bad predictions
  - Labels for data can come from 2 sources
    - Event-based (it's a label based on a fact that happened)
    - Assigned (usually by experts on the topic)
  - o It's important to have a way to validate data after it's collected and labelled
- Second is relevance
  - o Data needs to be relevant in explaining why an input resulted in a certain output
    - Ex: shoe sizes have nothing to do with loan payment rates
- Third is *quantity* 
  - o Generally speaking, most algorithms will need a lot of data for meaningful results
- Fourth is *variability* 
  - Data needs to be diverse
  - Algorithm gains a broader view of the domain
- Fifth is ethics
  - o Concerns include bias, security, privacy, and consent
  - Biased data leads to biased predictions
    - Can be unintentional or implicit from human ethics
    - Need to minimize as much as possible

import pandas

- pandas is the go-to package when it comes to data analysis
- Lots of great functions that makes it easy to use

#### **Basic Data Structures**

```
# a 1d list is called a 'series'
fruits = ["apple", "banana", "orange"]
series = pandas.Series(fruits)

# a 2d list is a 'dataframe'
myDict = {
    "letters": ['a','b','c'],
    "numbers": [1,2,3],
    "decimals": [1.1, 2.2, 3.3]
}
df = pandas.DataFrame(myDict)

# custom column names
labels = ["label1", "label2", "label3"]
df2 = pandas.DataFrame(myDict, labels)
```

- Data structures are heterogenous
  - Can store different kinds of data
  - 1D is a series and 2D is a dataframe
- Dataframes can either infer the column labels or you can pass in a custom list for the labels

#### **Reading from Files**

```
csv = pandas.read_csv("path/to/file.csv")

# if an Excel file has multiple sheets, it'll read the first one by default
excel = pandas.read_excel("path/to/file.xlsx")

# specify sheet for Excel
excel2 = pandas.read_excel("path/to/file.xlsx", sheet_name="sheet2")
```

## **Data Exploration**

## **Describing Data**

- The idea of data exploration is to be able to describe our data
  - What it's about, how much data, and what's the quality
- A single "row" of data is called an <u>instance</u>
  - Also called a record or observation
  - This is a single example of the general concept behind the data collection
- Features describe important characteristics of the instance
  - o Categorical: holds discrete data, limited to a set of possible values
  - Continuous: usually integers or real numbers, infinite possibilities
- Sometimes we want to predict features of a dataset
- Dimensionality is the number of features in a dataset
  - More dimensions means more details
  - Also means higher computational complexity
- Sparsity and density describe how much data we have
  - Complementing concepts
  - Ex: missing 20% data = 20% sparse, 80% dense

## Describing Data in Python

#### **Basics**

```
data = pandas.read_csv("dataFile.csv")
data.info()
print(data.head())
```

- The info() command will display a bunch of useful info describing our dataset
  - Dimensions
  - Types of data
- head() gives a sneak-peek of what our data actually looks like
  - Returns first 5 rows

#### **Aggregations**

```
print(data[["ColumnName"]].describe())
```

- This allows us to get info regarding a specific attribute
- It'll return 4 basic info for non-numbers
  - o count: number of (nonempty) entries
  - o unique: number of unique entries
  - top: the most frequently occurring entry
  - freq: how often it occurs
- In columns that are statistical, it'll return

```
count: number of entries
```

- o mean
- o std
- 0 25%
- 50% (aka median)
- 0 75%
- o max

#### **Getting Value Counts**

```
# gives a count of each unique value
data[["ColumnName"]].value_counts()

# gives percent of each unique value
data[["ColumnName"]].value_counts(normalize=True)
```

#### **Other Functions**

```
# get the mean of values in a column
data[["ColumnName"]].mean()

# group instances, and then get the mean values of each group
data.groupby("ColumnName1")[["ColumnName2"]].mean()

# sort based on a column (default is ascending)
data.groupby("ColumnName1")[["ColumnName2"]].mean().sort_values(by="ColumnName3")
```

#### **Multiple Aggregations**

```
# 'agg' gives us multiple aggregations
data.groupby("ColumnName1")[["ColumnName2"]].agg(["mean", "median", "max"])
```

### **Data Visualizations**

- It's sometimes easier to show patterns with visuals
- <u>Comparison</u> visualizations show the difference between 2 or more items
  - Ex: a box plot
- Relationship visualizations shows how 2+ variables can affect each other
  - Ex: line charts and scatter plots
- <u>Distribution</u> visualizations shows stat distribution of a single feature

- Ex: histograms
- Composition visualizations show the inner makeup of data
  - Ex: stacked bar charts, pie charts

## Data Visualization in Python

```
import pandas
import matplotlib.pyplot as plt

vehicles = pandas.read_csv("vehicles.csv")
```

- matplotlib is a popular package for generating data visualizations
- Note that in order to actually display the plot, we need to run plt.show() at the end

#### Scatter

```
vehicles.plot.scatter(x="citympg",y="co2emissions")
plt.show()
```

## Histogram

```
vehicles[["co2emissions"]].plot.hist()
plt.show()
```

#### Box

```
pivot = vehicles.pivot(columns="drive", values="co2emissions")
pivot.boxplot(figsize=(10,6))
plt.show()
```

- In order to use a box plot, we need to first create a pivot table
  - X-axis is column values, Y-axis is cell values

#### Stacked Bar Chart

```
pivot = vehicles.groupby("year")["drive"].value_counts().unstack()
pivot.plot.bar(stacked=True, figsize=(10,6))
```

```
plt.show()
```

- We need to first create a pivot
  - We group by year
  - Then aggregate by drive

## **Data Preparation**

## Common Data Quality Issues

- The process of ensuring the data is suitable for our ML model
- One of the most common data issues is missing data
  - Usually a few missing features in a record here and there
  - Can result from human error, bias, or lack of reliable input
  - Changes in data collection methods can also result in missing data
- Several ways to resolve this
  - Remove records with missing data
  - Use holders to represent missing data (like N/A or -1)
  - We can also try to use imputation, which is a process to reasonably guess what those values are
    - Ex: replace all missing values with the median of non-missing values
- Another issue is *outliers* 
  - Features that are unusual or very different than others
  - To fix outliers, you need to first understand what the outliers convey
- One valid reason for outliers is class imbalance
  - This is when the real world distribution of values is not uniform
  - More instances of a class label than others
  - Can lead to misleading predictions if not properly handled
- Ways to resolve
  - Undersample majority classes

## Resolving Missing Data in Python

## **Check for Missing Values**

```
import pandas
students = pandas.read_excel("students.xlsx")
mask = students['State'].isnull()

# display 'True' if attribute is missing, 'False' otherwise
print(mask)
```

```
# only display rows with missing data
print(students[mask])
```

#### **Remove Missing Values**

```
# remove all rows with any missing values
students.dropna()
```

- This is a rather extreme approach
- We usually want to remove rows that have missing values in a particular column

```
# only drop rows if both State and Zip are missing
students.dropna(subset=["State","Zip"], how="all")

# drop columns with any missing values
students.dropna(axis=1)

# specifies how many values need to be missing before it's removed
students.dropna(axis=1, thresh=10)
```

#### **Resolve Missing Values**

```
# replace all missing values with a constant value
students.fillna({'Gender':'Female'})

# replace all missing values with a value from a function
students.fillna({'Age':students['Age'].median()})

# change value of specific cell
mask = (studnets['City'] == 'Granger') & (students['State'] == 'IN')
students.loc[mask, 'Zip'] = 46530
```

## Normalizing Data

- Goal of normalization (also called standardization) is to ensure all data share a common property
  - o Typically involves scaling data to fit within a range
  - Reduces complexity and improves interpretability
- Z-Score Normalization is one approach
  - Ensures the data has a mean of 0 and std of 1

$$\circ \ v' = rac{v - ar{F}}{\sigma_F}$$

- lacksquare  $ar{F}$  is mean
- $\sigma_F$  is std
- lacksquare F is feature
- v and v' are the old and new values
- Ex: suppose we have the values 12000, 24000, 30000, 40000, 98000
  - Mean is 40800, std is 33544
  - v=40000:  $v' = \frac{40000 40800}{33544} = -0.024$
  - Final normalized values are -0.859, -0.500, -0.322, -0.024, -1.705
- Mean-max normalization gives our data a range between 2 user-defined bounds
  - Typically the bounds are 0 and 1

$$\circ \ \ v' = rac{v-min_F}{max_F-min_F}(upper_F-lower_F) + lower_F$$

 $\circ~$  Ex: bound 12000, 24000, 30000, 40000, 98000 to 0 and 1

$$v=30000$$
:  $v'=rac{30000-12000}{98000-12000}(1-0)+0=0.209$ 

- Final normalized values are 0.000, 0.140, 0.209, 0.326, 1.000
- Both these approaches are suitable for data with no significant outliers
- Log transformation works for data with lots of outliers
  - $\circ v' = \log v$
  - Base can either be 2 or 10
  - Note that this only works for positive values

## Normalize Data in Python

• We use the scikit-learn package for data transformations

#### Min-Max

```
from sklearn.preprocessing import MinMaxScaler

# transform data
c02emissions_mm = MinMaxScaler().fit_transform(vehicles[['co2emissions']])

# convert to pandas dataframe
c02emissions_mm = pandas.DataFrame(co2emissions_mm, columns=['co2emissions'])

# we can then describe and plot it
print(co2emissions_mm.describe())
c02emissions_mm.plot.hist(bins=20, figsize=(10,6))
```

- The histogram for the original and transformed data will look the same
  - What changed is the scale of the X-axis

#### **Z-Score**

```
from sklearn.preprocessing import StandardScaler

c02emissions_zm = StandardScaler().fit_transform(vehicles[['co2emissions']])
c02emissions_zm = pandas.DataFrame(c02emissions_zm, columns=['co2emissions'])
print(c02emissions_zm.describe())
c02emissions_zm.plot.hist(bins=20, figsize=(10,6))
```

Same process, just different Scaler object

## Sampling

- Sometimes we need to split our current dataset
  - Current data could be too big
  - We might want to hold on to some of the data for later use
- This process is called sampling
  - The overall dataset is called a <u>population</u>
  - The subset we chose is called the <u>sample</u>
- Sampling methods
  - Random sampling without replacement
  - Random sampling with replacement
    - Also called bootstrapping
    - Good for datasets with little data
  - Stratified random sampling
    - Ensures the sample's distribution matches the population's

#### Sample Data in Python

```
response = 'co2emissions'
y = vehicles[[response]]

predictors = list(vehicles.columns)
predictors.remove(response)
x = vehicles[predictors]
```

- Suppose in vehicles.csv, we want co2emissions to be the output based on the other variables
  - co2emissions is the response
  - The others are the predictors

## **Split Data using Simple Random Sampling**

```
from sklearn.model_selection import train_test_split
```

```
x_train, x_test, y_train, y_test = train_test_split(x, y)
```

- x\_train holds independent variables of training set
- y\_train holds dependent variables of training set
- x\_test holds independent variables of testing set
- y\_test holds dependent variables of testing set

```
x_train, x_test, y_train, y_test = train_test_split(x, y, test_size=0.4)
```

- By default, 75% of dataset goes to training and 25% goes to test
- We can change it with test\_size

#### **Split Data using Stratified Random Sampling**

```
x_train, x_test, y_train, y_test = train_test_split(x, y, test_size=0.4,
stratify=x['drive'])
```

- We use stratify to specify which feature (column) to stratify by
- This ensures the dataset's distribution matches the distribution of population more closely

## Reducing Data Dimensionality

- Idea is to reduce number of features before modeling
- The "curse of dimensionality" is the idea that increasing features will eventually decrease the performance of the model
  - We need exponentially more data instances per feature
  - Unless we can provide more data instances, our model will decrease
  - There's a limit to how much data was can collect, leading to a limit to the number of features we should have for optimal performance
- Feature selection is one way to reduce dimensionality
  - Identify min number of features needed for a good performance
  - Remove features that have minimal impact on the model performance
- Feature extraction is another method
  - Use math to reduce it
  - Results in completely new features
  - While still reliable, the disadvantage is that the new feature values are harder to interpret as a user

## Modeling & Evaluation

## Modeling

- The most well known phase of ML
- Objective is to identify the best ML modeling approach to solve the problem
- For supervised learning, there's 2 categories
  - o Classification: use features to produce a label
  - o Regression: use features to predict a continuous value
- Lots of different ML techniques can be used for both classification and regression
- There are also some techniques that are specifically for regression problems

#### **Evaluation**

- To evaluate a model, we need to run it on data it's never seen before
  - This ensures we have a reliable assessment of the model
- We usually grade the model with an accuracy metric
  - For classification, we use % correct
  - $\circ$  For regression, we use Mean Absolute Error:  $MAE = rac{\sum |Predicted-Actual|}{NumberTestInstances}$ 
    - This gives us an average margin of error

## **Building Model in Python**

#### **Build Model**

```
import pandas
bikes = pandas.read_csv("bikes.csv")

response = "rentals"
y = bikes[[response]]

predictors = list(bikes.columns)
predictors.remove(response)
x = bikes[predictors]

from sklearn.model_selection import train_test_split
x_train, x_test, y_train, y_test = train_test_split(x, y)
```

#### **Train Model**

```
from sklearn.linear_model import LinearRegression
model = LinearRegressions().fit(x_train, y_train)
```

• Linear regression assumes there's a linear relationship between each feature and the output

```
# useful variable values
model.intercept_
model.coef_
```

- intercept\_ gives us the intercept for the equation
  - o It's a constant that gets added to the end of calculations
- coef\_ lists corresponding weights for each feature in the respective order
- These 2 variables basically give us the equation in which we can make predictions manually

#### **Evaluate Model**

```
model.score(x_test, y_test)
```

- This gives us the R-squared value
  - R-square value is also called the *coefficient of determination*
  - Common measurement for linear regression models
  - The close it is to 1, the better it is
- A R-square value of 0.98 means it explains 98% of the variability in response values for test data

```
from sklearn.metrics import mean_absolute_error
y_pred = model.predict(x_test)
mean_absolute_error(y_test, y_pred)
```

• This gives us the MAE, which is an average margin of error