IBM - Machine Learning Professional Certificate

Supervised Machine Learning: Classification

The Major Cities across the United States to Predict Rainfall

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TABLE OF CONTENTS

1. INTRODUCTION

- 1.1 Introduction
- 1.2 Objective
- 1.3 Coding Environment

2. DATA PROCESSING

- 2.1 Data Collection
- 2.2 Data Description
- 2.3 Quality Assessment
 - 2.3.1 Normal Distribution
 - 2.3.2 Missing Value
 - 2.3.3 Invalid Value
 - 2.3.4 Duplicate Value
 - 2.3.5 Outlier Value
- 2.4 Data Preprocessing
 - 2.4.1 Data Cleaning
 - 2.4.2 Missing Value Handling

- 2.4.3 Duplicate Handling
- 2.4.4 Outlier Handling
- 2.4.5 Centering and Scaling
- 2.4.6 Data Transformation
- 2.4.7 Correlation Coefficient
- 2.4.8 Data Encoding

2.5 - Exploratory Data Analysis

- 2.5.1 Data Visualization Analysis
- 2.5.2 Hypothesis Testing

3. MODELING

- 3.1 Data Splitting
- 3.2 Evaluation Metric
- 3.3 Hyperparameter Tuning
 - 3.3.1 Linear Regression
 - 3.3.2 K-Nearest Neighbors (KNN)
 - 3.3.3 Support Vector Machine (SVM)
 - 3.3.4 Decision Tree
 - 3.3.5 Random Forest

3.4 - Hyperparameter Tuning with Downsampling

- 3.4.1 Linear Regression
- 3.4.2 K-Nearest Neighbors (KNN)
- 3.4.3 Support Vector Machine (SVM)
- 3.4.4 Decision Tree
- 3.4.5 Random Forest

• 3.5 - Hyperparameter Tuning with Upsampling

- 3.5.1 Linear Regression
- 3.5.2 K-Nearest Neighbors (KNN)
- 3.5.3 Support Vector Machine (SVM)
- 3.5.4 Decision Tree
- 3.5.5 Random Forest

4. SUMMARY

- 4.1 Models Evaluation
- 4.2 Summary

5. REFERENCES

1. INTRODUCTION

1.1 - Introduction

This project focuses on analyzing detailed weather data from 20 major cities across the United States to predict rainfall. We will utilize various classifier models commonly applied to binary classification problems and, after evaluating and comparing these models, we aim to identify the one most suitable classifier model for our objectives.

A comprehensive exploration of the dataset will be conducted to assess its quality. To enhance the statistical significance and stability of our training model, we will employ a range of standard data processing techniques to clean and optimize the dataset. Once the data has been refined, we will perform visual analyses to further evaluate its statistical properties.

Once the dataset is prepared for modeling, we will implement several classification techniques, including Logistic Regression, K-Nearest Neighbors (KNN), Support Vector Machine (SVM), Random Forest, and Decision Tree, to predict rainfall. To effectively compare the performance of various classification models and address the class-imbalance in the dataset, this project will evaluate different classifiers using Hypermetrices Tuning with Downsampling and Upsampling preprocessing methods.

1.2 - Objective

- **Main Objective**: This analysis aims to determine whether rain occurs or not, specifying whether the models will focus on prediction or interpretation.
- **Dataset Overview**: The study utilizes a dataset comprising weather data from 20 major cities across the United States over the past 731 days, summarizing its key attributes and relevance.
- Data Exploration and Preparation: A brief summary of the data exploration process is provided, along with actions taken for data cleaning and feature engineering to enhance the dataset's quality and applicability.

- **Model Training and Comparison**: We summarize the training of various classifier models, exploring their differences in explainability and predictability. All models are compared using the same training and test splits, as well as a consistent cross-validation method.
- **Final Model Evaluation**: By employing this standardized approach, we identify the final classifier models that best fit the dataset in terms of both accuracy and explainability.
- **Key Findings and Insights**: A summary of key findings and insights is presented, highlighting the main drivers of the final model and the valuable insights derived from the dataset through the classifier analysis.

1.3 - Coding Environment

The following required modules are pre-installed in the Skills Network Labs environment. However if you run this notebook commands in a different Jupyter environment (e.g. Watson Studio or Ananconda) you will need to install these libraries by removing the # sign before !mamba in the code cell below.

```
In [ ]: # All Libraries required for this lab are listed below. The libraries pre-in
        # !mamba install -qy pandas==1.3.4 numpy==1.21.4 seaborn==0.9.0 matplotlib==
        # Note: If your environment doesn't support "!mamba install", use "!pip inst
In [ ]: import pandas as pd
        import numpy as np
        import seaborn as sns
        import matplotlib.pyplot as plt
        %matplotlib inline
        import warnings
        warnings.filterwarnings("ignore", category=FutureWarning)
        from sklearn.preprocessing import StandardScaler, PowerTransformer, MinMaxSc
        from sklearn.experimental import enable iterative imputer
        from sklearn.impute import IterativeImputer, KNNImputer
        from sklearn.linear model import LinearRegression, LogisticRegression
        from sklearn.model selection import train test split, StratifiedShuffleSplit
        from sklearn.metrics import accuracy score, precision score, recall score, f
        from sklearn.neighbors import KNeighborsClassifier
        from sklearn.svm import SVC
        from sklearn.tree import DecisionTreeClassifier
        from sklearn.ensemble import RandomForestClassifier
        from imblearn.under sampling import RandomUnderSampler, NearMiss, CondensedN
        from imblearn.over sampling import RandomOverSampler, SMOTE, BorderlineSMOTE
        import scipy
```

2. DATA PROCESSING

2.1 - Data Collection

The dataset (USA Rainfall Prediction Dataset (2024-2025)) provides comprehensive weather data collected from 20 major cities across the USA during the years 2024 and 2025. It contains a variety of weather attributes that are crucial for predicting whether it will rain the next day or not. With over 2 years of daily data, this dataset serves as a perfect starting point for building predictive models, analyzing weather trends, or even developing weather-related applications.

Each row represents a single day's weather conditions, including important features like temperature, humidity, wind speed, cloud cover, atmospheric pressure, and precipitation. The target feature, "Rain Tomorrow," is a binary label (1 = Yes, 0 = No) indicating whether it rained the next day.

The dataset can be used for:

- Building rainfall prediction models
- Exploring weather patterns
- Studying the relationship between various weather factors
- Improving forecast accuracy using machine learning models

```
In [ ]: # Loading the dataset from local drive
data = pd.read_csv("/content/usa_rain_prediction_dataset_2024_2025.csv")
print(data)
```

0 1 2 3 4 73095 73096 73097 73098 73099	Date 2024-01-01 2024-01-02 2024-01-03 2024-01-05 2025-12-27 2025-12-28 2025-12-30 2025-12-31	Location New York New York New York New York New York New York Outline Washington D.C. Washington D.C. Washington D.C. Washington D.C. Washington D.C. Washington D.C.	80.943050 78.097552	28.712617 64.740043 59.738984 34.766784 65.099438 30.610525 96.740232 63.900004	Wind Speed 28.379506 12.436433 14.184831 19.444029 3.689661 28.778327 12.282890 2.894762 24.632400 22.722505	\
0 1 2 3 4 73095 73096 73097 73098 73099	Precipitati 0.0000 0.5269 0.9168 0.0941 1.3612 0.0000 0.8710 1.1919 0.4834 0.1519	00 69.617966 95 41.606048 84 77.364763 34 52.541196 72 85.584000 00 54.168514 00 22.068055 56 52.336048 21 76.785280	Pressure 1026.030278 995.962065 980.796739 979.012163 1031.790859 977.083747 980.591675 1016.469174 1032.396146 974.835534	Rain Tomorr	0 0 1 0 0 0 0 0 0 1 1 1 0 0 0 0 0 0 0 0	

[73100 rows x 9 columns]

2.2 - Data Description

This project assumes that it is possible to predict rainfall in major cities of United States using various data related to weather.

The **Feature** Variables:

Features	Feature Type	Description	Data Type
Date	Object	The date from January 1, 2024, to December 31, 2025 (Only has 731 days)	Object
Location	Object	The 20 major cities in the United States	Object
Temperature	Numeric	The average degree of hotness as measured on the day	Float
Humidity	Numeric	The average concentration of water vapor present in the air on the day	Float
Wind Speed	Numeric	The average speed of air is moving over a certain city on the day	Float

Features	Feature Type	Description	Data Type
Precipitation	Numeric	The average condensation of atmospheric water vapor that falls from clouds	Float
Cloud Cover	Numeric	The amount of opaque clouds covering the sky valid for the day	Float
Pressure	Numeric	The force exerted against a surface by the weight of the air above that surface	Float

The **Target** Variables:

	Target	Feature Type	Description	Data Type
	Rain Tomorrow	Categorica	It rained on the next day $(1 = Yes, 0 = No)$	Integer
In []:	# Setting the data["Rain Ton	-	variables ta["Rain Tomorrow"].astype('category	/ˈ).cat.set_cat

In []: data.info()

<class 'pandas.core.frame.DataFrame'>
RangeIndex: 73100 entries, 0 to 73099

Data columns (total 9 columns):

#	Column	Non-Null Count	Dtype
0	Date	73100 non-null	object
1	Location	73100 non-null	object
2	Temperature	73100 non-null	float64
3	Humidity	73100 non-null	float64
4	Wind Speed	73100 non-null	float64
5	Precipitation	73100 non-null	float64
6	Cloud Cover	73100 non-null	float64
7	Pressure	73100 non-null	float64
8	Rain Tomorrow	73100 non-null	category
dtyp	es: category(1)	, float64(6), ob	ject(2)

In []: data_describe_object = data.describe(include='object')
 data_describe_object.T

 Out[]:
 count
 unique
 top
 freq

 Date
 73100
 731
 2024-01-01
 100

 Location
 73100
 20
 New York
 3655

memory usage: 4.5+ MB

```
In [ ]: data_describe_numeric = data.describe(include='number')
```

```
data describe numeric.T
Out[]:
                                               std
                                                          min
                                                                     25%
                       count
                                   mean
                               65.182270 20.205793
        Temperature 73100.0
                                                     30.000766
                                                                47.678968
                                                                             65.29
                               59.875041 23.066115
           Humidity 73100.0
                                                     20.000272
                                                                39.800732
                                                                             59.88
         Wind Speed 73100.0
                               15.017946
                                           8.668729
                                                      0.000712
                                                                 7.485182
                                                                             15.10
                                                                 0.000000
        Precipitation 73100.0
                                                                              0.19
                                0.390635
                                           0.474833
                                                      0.000000
         Cloud Cover 73100.0
                               54.942807 25.982487
                                                                             55.01
                                                     10.000856
                                                                32.318668
           Pressure 73100.0 1005.176013 20.203889 970.000919 987.697646 1005.28
In [ ]: data describe category = data.describe(include='category')
        data describe category.T
Out[]:
                       count unique top
                                           freq
```

0 56988

2.3 - Quality Assessment

2.3.1 - Normal Distribution

Rain Tomorrow 73100

In Machine Learning, data satisfying **Normal Distribution** is beneficial for model building (**Especially regression based models**).

Models like **Linear Discriminant Analysis (LDA)**, **Gaussian Naive Bayes**, **Logistic Regression**, **Linear Regression**, etc., are explicitly calculated from the assumption that the distribution is a bivariate or multivariate normal.

```
# The summary of skewness and kurtosis
# Get the data with type numeric
data_numeric = data.select_dtypes(include='number')
# Get the name of numeric column
data_numeric_column = list(data_numeric.columns)

# Get the skewness for numeric column
data_numeric_skew = data_numeric.skew()
# Get the kurtosis for numeric column
data_numeric_kurtosis = data_numeric.kurtosis()
```

Out[]:		Column	Skewness	Kurtosis
	3	Precipitation	1.241112	1.018783
	1	Humidity	0.003375	-1.200780
	4	Cloud Cover	0.000772	-1.204550
	2	Wind Speed	-0.007681	-1.205991
	5	Pressure	-0.010555	-1.201018
	0	Temperature	-0.014391	-1.198375

Based on the output above, we can conclude that most of the skewness in the numeric features is close to 0, indicating that the dataset is **approximately normally distributed**.

The kurtosis values are predominantly negative, suggesting that the dataset is **platykurtic**.

Overall, the dataset generally adheres to a normal distribution, with the **exception of the Precipitation featur**.

2.3.2 - Missing Value

Missing Values contain in most of the real world datasets, i.e., feature entries with no data value stored. As many machine learning algorithms do not support missing values, detecting the missing values and properly handling them, can have a significant impact.

```
In []: # Quick check of missing variables
    data_missing = data.isnull()

data_missing.sum()
```

Out[]:

Date 0

Location 0

Temperature 0

Humidity 0

Wind Speed 0

Precipitation 0

Cloud Cover 0

Pressure 0

Rain Tomorrow 0

dtype: int64

Since there are no missing values in any of the columns, this indicates that the dataset is complete.

If there were missing values, the ratio of missing data in each column could be calculated using the appropriate code.

```
# The summary of missing variables from whole columns
# Get the name of columns
data column = list(data.columns)
# Get the total rows
data row count = np.array([len(data)] * len(data column))
# Count of missing variables
data missing count = data row count - np.array(data.count())
# Missing variables / Total rows
data missing rate = np.divide( data missing count, data row count, out=np.ze
data missing summary = pd.DataFrame( zip( data column,
                       data row count,
                       data_missing_count,
                       data_missing_rate ),
                    columns = [ "Column",
                            "Rows",
                            "Missing Values",
                            "Missing Rate %" ] )
data_missing_summary.sort_values(by="Missing Values", ascending=False)
```

Out[]:		Column	Rows	Missing Values	Missing Rate %
	0	Date	73100	0	0.0
	1	Location	73100	0	0.0
	2	Temperature	73100	0	0.0
	3	Humidity	73100	0	0.0
	4	Wind Speed	73100	0	0.0
	5	Precipitation	73100	0	0.0
	6	Cloud Cover	73100	0	0.0
	7	Pressure	73100	0	0.0
	8	Rain Tomorrow	73100	0	0.0

2.3.3 - Invalid Value

Invalid Values (Badly Formatted Values) refer to inconsistent entries commonly found in datasets, such as variables with different units across data points or incorrect data types. For instance, numerical variables like percentages and fractions are sometimes mistakenly stored as strings. It is essential to detect and correct these cases to ensure that machine learning algorithms can properly process and analyze the actual numerical values.

In []:	data_describe_object.T					
Out[]:		count	unique	top	freq	
	Date	73100	731	2024-01-01	100	
	Location	73100	20	New York	3655	

In []:	data_describe_numeric.T							
Out[]:		count	mean	std	min	25%		
	Temperature	73100.0	65.182270	20.205793	30.000766	47.678968	65.29	
	Humidity	73100.0	59.875041	23.066115	20.000272	39.800732	59.88	
	Wind Speed	73100.0	15.017946	8.668729	0.000712	7.485182	15.10	
	Precipitation	73100.0	0.390635	0.474833	0.000000	0.000000	0.19	
	Cloud Cover	73100.0	54.942807	25.982487	10.000856	32.318668	55.01	
	Pressure	73100.0	1005.176013	20.203889	970.000919	987.697646	1005.28	

Currently, the data types align with those described in the data documentation, and there are no invalid data types present in the dataset.

2.3.4 - Duplicate Value

Duplicate Values can appear in various forms, such as multiple entries of the same data point, repeated instances of entire columns, or duplication within an ID variable. While duplicates may be valid in some datasets, they often result from errors during data extraction or integration. Therefore, it is crucial to detect these duplicate values and determine whether they represent true duplicates or are a legitimate part of the dataset.

```
In [ ]: # Quick check of duplicate row
        data.duplicated().sum()
Out[]: 0
In [ ]: # Quick check of unique value
        data unique count = data.nunique()
        data unique count
Out[]:
                            0
                  Date
                          731
              Location
                           20
          Temperature 73100
             Humidity 73100
           Wind Speed 73100
          Precipitation 44754
           Cloud Cover 73100
              Pressure 73100
        Rain Tomorrow
                            2
```

dtype: int64

```
# "The summary of duplicate variables from whole columns
# ""

# Unique variables / Total rows
data_duplicate_rate = ( np.ones(len(data_unique_count)) - np.divide( list(data_unique_count, data_row_count, data_unique_count, data_unique_count, recolumns = [ "Column", "Rows", "Unique Values", "Duplicate Rate %" ] )

data_duplicate_summary.sort_values(by="Duplicate Rate %", ascending=False)
```

Out[]:	Column	Rows	Unique Values	Duplicate Rate %
out[].	Column	ROWS	Ullique values	Duplicate Nate 70

	Column	IXOVIS	Ollique values	Duplicate Nate /6
8	Rain Tomorrow	73100	2	99.997264
1	Location	73100	20	99.972640
0	Date	73100	731	99.000000
5	Precipitation	73100	44754	38.777018
2	Temperature	73100	73100	0.000000
3	Humidity	73100	73100	0.000000
4	Wind Speed	73100	73100	0.000000
6	Cloud Cover	73100	73100	0.000000
7	Pressure	73100	73100	0.000000

Upon reviewing the object-type features and the target feature, we found that they align with the ranges specified in the data description.

In examining the numeric features, we note that all features, except for the Precipitation feature, have a duplicate rate of 0%. This indicates that there are no duplicate values in the dataset, while the Precipitation feature contains some duplicates.

2.3.5 - Outlier Value

Outliers (Anomalies) are data points that differ substantially from the rest of data, and they may arise due to the diversity of the dataset or because of errors/mistakes. As machine learning algorithms are sensitive to the range and

distribution of attribute values, identifying the outliers and their nature is important for assessing the quality of the dataset.

```
In [ ]: #-----
        # The boxplots of outlier
        #------
        # Setting the size of subplots
        _, ax = plt.subplots(nrows=3, ncols=2, figsize=(10, 5))
        ax = ax.ravel()
        # Display the boxplot
        for index, column in enumerate(data_numeric.columns):
          sns.boxplot(data=data, ax=ax[index], x=column)
        # Do not blocked any title or label
        plt.tight_layout()
        plt.show()
                       60
                                           100
                                                                   60
                                                                       70
                                                                                    100
                       Temperature
                                                                 Humidity
                                                 0.0
                                20
                                                       0.5
                                                            1.0
                          15
                                                                  1.5
                                                                        2.0
                                                                             2.5
                                                                                   3.0
                       Wind Speed
                                                                Precipitation
                                                                    1010
            20
                            60
                                           100
                                                 970
                                                      980
                                                           990
                                                               1000
                                                                         1020
                                                                              1030
                                                                                    1040
                       Cloud Cover
In [ ]: #-----
        # The summary of outlier variables from numeric columns
        # Using IQR method to detect outlier variables
        data describe numeric q1 = data describe numeric.T["25%"] #data numeric.quar
        data describe numeric q3 = data describe numeric.T["75%"] #data numeric.quar
        data describe numeric iqr = data describe numeric q3 - data describe numeric
        # Min and Max boundary to detect outlier
        data describe numeric min iqr = data describe numeric q1 - 1.5 * data descri
        data describe numeric max iqr = data describe numeric q3 + 1.5 * data descri
        # Count the outliers
        data outlier count = ( (data numeric < data describe numeric min iqr) | (dat
        # Get the total rows
```

Out[]:		Column	Rows	Outlier Values	Outlier Rate %
	3	Precipitation	73100	1192	1.630643
	0	Temperature	73100	0	0.000000
	1	Humidity	73100	0	0.000000
	2	Wind Speed	73100	0	0.000000
	4	Cloud Cover	73100	0	0.000000
	5	Pressure	73100	0	0.000000

The plots above indicate the presence of outliers in the Precipitation feature.

2.4 - Data Processing

2.4.1 - Data Cleaning

Data Cleaning is the process of fixing or removing incorrect, corrupted, incorrectly formatted, duplicate, or incomplete data within a dataset. When combining multiple data sources, there are many opportunities for data to be duplicated or mislabeled. If data is incorrect, outcomes and algorithms are unreliable, even though they may look correct.

In **2.3 - Quality Assessment**, we examined the dataset and confirmed that there were no missing values or incorrect data formats. However, we identified outliers that need to be addressed.

The Date feature in the dataset is not critical for our analysis. Specifically, we aim to predict whether it will rain tomorrow, and data from the same date in the

previous year is not strongly correlated.

While the Date feature could be transformed into a seasonal classification, given that weather patterns often vary with the seasons (for example, spring typically experiences greater likelihood of rain), but we prefer to keep the dataset and model uncomplicated.

Therefore, we propose discarding the Date feature to simplify the model training process.

```
In [ ]: # Discard the unimportant features
           data.drop(columns=["Date"], inplace=True)
           data describe numeric = data.describe(include='number')
           data.info()
         <class 'pandas.core.frame.DataFrame'>
         RangeIndex: 73100 entries, 0 to 73099
         Data columns (total 8 columns):
             Column Non-Null Count Dtype
Location 73100 non-null object
Temperature 73100 non-null float64
Humidity 73100 non-null float64
Wind Speed 73100 non-null float64
Procipitation 73100 non-null float64
          #
         --- ----
          0
          1
          3
              Precipitation 73100 non-null float64
              Cloud Cover 73100 non-null float64
Pressure 73100 non-null float64
          5
          6
               Rain Tomorrow 73100 non-null category
         dtypes: category(1), float64(6), object(1)
         memory usage: 4.0+ MB
```

2.4.2 - Missing Value Handling

Missing Value Handling usually uses some techniques:

1. Median or Mean

- No matter use median or mean as imputation value, it has limitations. For example, imputing with the mean may not be appropriate if the data has extreme values, as it can be heavily influenced by outliers
- Similarly, imputing with the median may not be appropriate if the data is multimodal, as it may not represent the true central tendency of the data

2. Iterative

 An advanced imputation method that models each feature with missing values as a function of other features in a round-robin fashion. It uses a regression model to estimate missing values based on the observed values of other features. The imputation process is performed iteratively, with each iteration refining the imputed values until convergence or a specified maximum number of iterations is reached

- Commonly used regression models: Linear Regression, Bayesian Ridge (regularized linear regression), Decision Trees Regressor, Random Forest Regressor, and K-Neighbors Regressor, etc.
- **K-Neighbors Regressor** is different from KNN imputation, which learns from samples with missing values by using a distance metric that accounts for missing values, rather than imputing them

3. K-Nearest Neighbors (KNN)

- KNN Imputer imputes missing values based on the nearest neighbors, which
 means it preserves the underlying relationships in the data. It takes into
 account the feature similarities between data points to estimate the missing
 values, making it more contextually relevant
- Non-Parametric method, which means it does not make assumptions about the data's distribution. It is suitable for both numeric and categorical data, making it versatile in handling various types of missing values

4. Multiple Imputation by Chained Equations (MICE)

• The procedure imputes missing data in a dataset through an iterative series of predictive models. In each iteration, each specified variable in the dataset is imputed using the other variables in the dataset. These iterations should be run until it appears that convergence has been met

In **2.3.2 - Missing Value**, we examined the dataset and confirmed that there are no missing values.

Method - Median or Mean

```
# #---
# Method - Median or Mean
#-----
"''
# Median
data_missing_imputation = data_describe_numeric.T["50%"].T
# Average/Mean
data_missing_imputation = data_describe_numeric.T["mean"].T

data.fillna(value = data_missing_imputation)
"''
```

Method - Iterative

```
In [ ]: #-----
        # Method - Iterative
        #-----
        1.1.1
        # Linear Regression
        data missing iterative estimator = LinearRegression()
        # Bayesian Ridge
        data missing iterative estimator = BayesianRidge()
        # Decision Trees Regressor
        data missing iterative estimator = DecisionTreeRegressor()
        # Random Forest Regressor
        data missing iterative estimator = RandomForestRegressor()
        # K-Neighbors Regressor
        data missing iterative estimator = KNeighborsRegressor()
        # Initialization iterative imputation object
        data missing iterative imputation = IterativeImputer(estimator = data missing
        # Replace the result in the original dataset
        data[data numeric.columns] = pd.DataFrame(data missing iterative imputation.
```

Method - K-Nearest Neighbors (KNN)

Summary after Missing Values Handling

2.4.3 - Duplicate Handling

In **2.3.4 - Duplicate Value**, we examined the dataset and confirmed that there are no values with a high duplication rate that require addressing.

```
In [ ]: # Drop the duplicate rows
    data.drop_duplicates(inplace=True)
    '''
```

Summary after Duplicate Handling

2.4.4 - Outlier Handling

Outlier Handling usually uses four different techniques:

1. **Deleting Observations**

- We delete outlier values if it is due to data entry error, data processing error or outlier observations are very small in numbers. We can also use trimming at both ends to remove outliers
- **BUT** deleting the observation is not a good idea when we have small dataset

2. Transforming Values

- Transforming variables can also eliminate outliers. These transformed values reduces the variation caused by extreme values
- If dataset has to many extreme values or skewed, Log Transformation,
 Cube Root Normalization, Box-transformation, Yeo-Johnson Power
 Transformation, etc., those techniques convert values in the dataset to smaller values
- BUT these technique not always give the best results. For example, Log
 Transformation requires that each transformed value not closing to zero;
 Box-transformation requires that each transformed value is positive,
 otherwise Yeo-Johnson Power Transformation needs to be used as an alternative

3. **Imputation**

- Like imputation of missing values, we can also impute outliers. We can use
 Mean, Median, Zero value in this methods. Since we imputing there is no loss of data
- Use missing value imputation methods, such as Iterative Imputation and K-Nearest Neighbors (KNN) Imputation

4. Separately Treating

- If there are significant number of outliers and dataset is small, we should treat them separately in the statistical model. One of the approach is to treat both groups as two different groups and build individual model for both groups and then combine the output
- **BUT** this technique is tedious when the dataset is large

In **2.3.5 - Outlier Value**, we examined the dataset and identified 1,192 outlier values in the Precipitation feature. Out of a total of 73,100 observations, these outliers represent approximately 1.63% of the dataset (1,192 out of 73,100). Therefore, we will use the simplest outlier handling method: deleting the affected observations.

```
In []: # Using IQR method to detect outlier variables except Year
    data_describe_numeric_q1 = data_describe_numeric.T["25%"] #data_numeric.quar
    data_describe_numeric_q3 = data_describe_numeric.T["75%"] #data_numeric.quar
    data_describe_numeric_iqr = data_describe_numeric_q3 - data_describe_numeric
```

```
# Min and Max boundary to detect outlier
data_describe_numeric_min_iqr = data_describe_numeric_q1 - 1.5 * data_descri
data_describe_numeric_max_iqr = data_describe_numeric_q3 + 1.5 * data_descri
# Count the outliers
data_outlier_count = ( (data_numeric < data_describe_numeric_min_iqr) | (data_numeric_min_iqr) | (da
```

Method - Deleting Observations

```
In []: #-----
# Method - Deleting observations
#-----
data.drop( data[ (data["Precipitation"] < data_describe_numeric_min_iqr["Precipitation"]</pre>
```

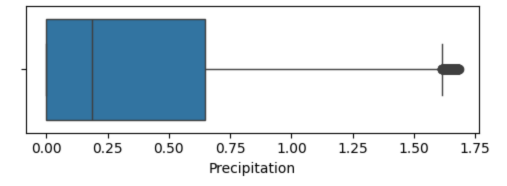
Method - Transforming Values

Method - Imputation

```
data_precipitation_outlier_imputation = data_describe_numeric["Precipitation
# Zero
data_precipitation_outlier_imputation = 0

data["Precipitation"] = np.where( (data["Precipitation"] < data_describe_num
'''</pre>
```

Visualize the Boxplot after Outlier Handling



2.4.5 - Centering and Scaling

Centering and Scaling ensures that the criterion for finding linear combinations of the predictors is based on how much variation they explain and therefore improves the numerical stability.

- **Standard Scaling** Converts features to **standard normal** variables, and it centers and scales a variable to mean 0 and standard deviation 1
- **Min-Max Scaling** Convert variables to continuous variables in the [0, 1] interval by mapping minimum values to 0 and maximum values to 1
- **Robust Scaling** Similar to min-max scaling, but instead maps the **interquartile range** (Q3 Q1) to [0, 1] interval, it means the variable itself takes values outside of the [0, 1] interval

```
In [ ]: # Get the data with type numeric that AFTER cleaning and outlier handling
data_numeric = data.select_dtypes(include='number')
```

Method - Standard Scaling

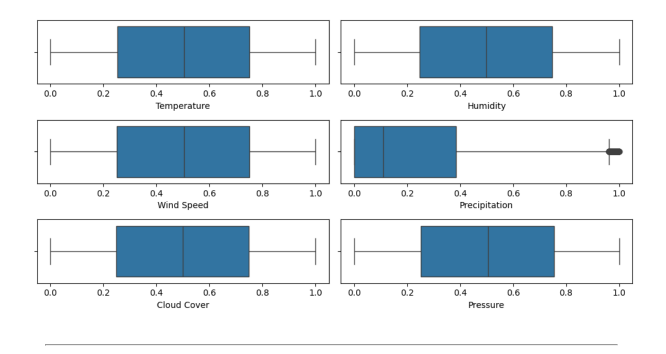
```
In []: #-----
# Method - Standard Scaling
#-----
data[data_numeric.columns] = StandardScaler().fit_transform(data_numeric)
```

Method - Min-Max Scaling

Method - Robust Scaling

```
In []: #-----
# Method - Robust Scaling
#-----
data[data_numeric.columns] = RobustScaler().fit_transform(data_numeric)
```

Visualize the Boxplot after Centering and Scaling



2.4.6 - Data Transformation

Features and predicted data are often **Skewed** (distorted away from the center), it degrades the model's ability to describe typical cases as it has to deal with rare cases on extreme values (**especially regression based models**).

Data Transformation usually can solve the skewed data. To ensure that the machine learning model capabilities is not affected, skewed data has to be transformed to approximate to a normal distribution. The method used to transform the skewed data depends on the characteristics of the data.

- Popular data transformation techniques include Log Transformation, Cube Root Normalization, Box-Transformation, Yeo-Johnson Power Transformation, etc.
- BUT these technique not always give the best results. For example, Log
 Transformation requires that each transformed value not closing to zero;
 Box-Transformation requires that each transformed value is positive, otherwise Yeo-Johnson Power Transformation needs to be used as an alternative

It is worth noting that tree-based models are not affected by these issues, as they can effectively ignore correlation concerns. Consequently, tree-based models do not require data transformation, centering, or scaling.

In **2.3.1 - Normal Distribution**, we verified that the skewness of all numerical features aligns with a normal distribution, except for the Precipitation feature, which exhibits skewness.

Even after addressing outliers in **2.4.4** - **Outlier Handling** and applying centering and scaling in **2.4.5** - **Centering and Scaling**, the boxplot still shows that the Precipitation feature remains skewed.

Method - Log Transformation

```
In []: #-----
# Method - Log Transformation
#-----
"''
# If data value closing to 0, D0 NOT use this method
data["Precipitation"] = np.log(data["Precipitation"])
"''
```

Method - Cube Root Transformation

```
In []: #-----
# Method - Cube Root Normalization
#-----
data["Precipitation"] = (data["Precipitation"]**(1/3))
'''
```

Method - Box-Transformation or Yeo-Johnson Power Transformation

Summary after Transformation

ut[]:		Column	Skewness	Kurtosis
	3	Precipitation	0.310341	-1.509251
	1	Humidity	0.004682	-1.199930
	4	Cloud Cover	0.001629	-1.204348
	2	Wind Speed	-0.008524	-1.206635
	5	Pressure	-0.010839	-1.201128
	0	Temperature	-0.014004	-1.197688

2.4.7 - Correlation Coefficient

Understanding the correlations between variables in a model is essential for several reasons:

1. Feature selection

- Process of choosing which variables or features to use in the model. Highly correlated features provide redundant information, so feature selection aims to remove uninformative features to simplify models
- By analyzing correlations, we can identify redundant features and select a minimal set of important features that best represent the target variable.
 This prevents overfitting and improves a model's ability to generalize

2. Reduce Bias

- Correlation analysis is also important for ensuring model fairness and avoiding bias. When certain features are highly correlated with sensitive attributes like gender or ethnicity, it can inadvertently encode biases into machine learning models if not properly addressed
- If a model relies too heavily on these correlated features, it risks
 discriminating against or disadvantaging certain groups. By identifying
 correlations between input features and sensitive attributes, we can evaluate
 models for potential biases, monitor feature importance, and apply
 techniques like fair representation learning to mitigate bias

3. Multicollinearity

- Another important aspect of analyzing feature correlations is detecting
 multicollinearity. Multicollinearity occurs when two or more predictor
 variables in a model are highly linearly correlated with each other. It can
 negatively impact models by increasing variance and making it difficult to
 determine the significance and effect of individual predictors
- Variables with high multicollinearity provide redundant information, similar to how correlated features do. However, multicollinearity is more problematic because it inflates standard errors and undermines reliability of estimated coefficients. By examining correlation matrices and variance inflation factors, we can identify cases of multicollinearity between input features

4. Interpretability and Debugging

- Understanding correlations also aids in interpreting machine learning models. As models become increasingly complex with many interacting variables, it can be difficult to explain why a model makes certain predictions
- By analyzing the correlation between input features and output targets, we
 gain insights into which variables have the strongest impact on the model's
 decisions. Knowing feature correlations further assists in debugging models
 that perform poorly. It allows we to identify any features that may be
 overwhelming the model or causing unintended biases

Interpreting a Correlation Coefficient

- The value of the correlation coefficient always ranges between 1 and -1, and we treat it as a general indicator of the strength of the relationship between variables
- The sign of the coefficient reflects whether the variables change in the same or opposite directions: a positive value means the variables change together in the same direction, while a negative value means they change together in opposite directions
- The absolute value of a correlation coefficient tells the magnitude of the correlation: the greater the absolute value, the stronger the correlation

Correlation Coefficient	Strength of Linearity / Monotonically	Correlation Type
-0.75 to -1	Perfectly	Negative
-0.5 to -0.75	Strong	Negative
-0.25 to -0.5	Moderate	Negative
0 to -0.25	Weak	Negative

Correlation Coefficient	Strength of Linearity / Monotonically	Correlation Type
0	None	Zero
0 to 0.25	Weak	Positive
0.25 to 0.5	Moderate	Positive
0.5 to 0.75	Strong	Positive
0.75 to 1	Perfectly	Positive

Methods of Calculate the Correlation Coefficient

Usually we use two mainstream methods to calculate the correlation coefficient:

1. Pearson's Correlation Coefficient

- The Pearson's correlation coefficient describes the linear relationship between two quantitative variables
- The assumptions for use Pearson's correlation coefficient:
 - 1. Expect a linear relationship between the two variables
 - 2. Both variables are on an interval or ratio level of measurement
 - 3. Data from both variables follow normal distributions
 - 4. Data have no outliers
- BUT it's not a good measure of correlation if variables have a nonlinear relationship, or if data have outliers, skewed distributions, or come from categorical variables

2. Spearman's Rank-Order Correlation

- Spearman's rank correlation coefficient is the most common alternative to Pearson method. It uses the rankings of data from each variable (e.g., from lowest to highest) rather than the raw data itself
- Use Spearman method when data fail to meet the assumptions of Pearson method. This happens when at least one of variables is on an ordinal level of measurement or when the data from one or both variables do not follow normal distributions
- Spearman's correlation coefficient measures the monotonicity of relationships, and monotonic relationships are less restrictive than linear relationships
 - Positive monotonic: when one variable increases, the other also increases

Negative monotonic: when one variable increases, the other decreases

In 2.3.1 - Normal Distribution, 2.4.1 - Cleaning and 2.4.2 - Outlier Handling, we confirmed that all assumptions for Pearson's correlation coefficient are met. Therefore, we will use Pearson's correlation coefficient for our analysis.

Method - Pearson's Correlation Coefficient

```
# Method - Pearson's Correlation Coefficient
# Computing the Pearson's correlation coefficient, BUT corr() won't have P-v
#data_numeric_correlation = data_numeric.corr(method='pearson')'''

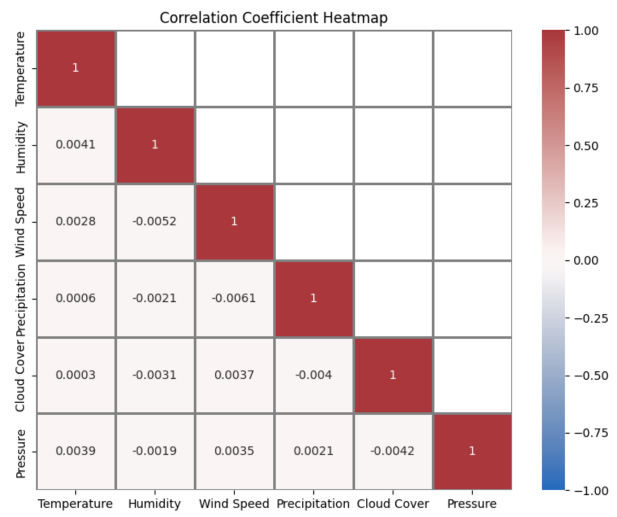
# Computing the Pearson's correlation coefficient and P-value
data_numeric_correlation = np.ones(shape=(data_numeric.shape[1], data_numeridata_numeric_correlation_p = np.zeros(shape=(data_numeric.shape[1], data_numeridata_numeric_columns):
    for index_1, column_1 in enumerate(data_numeric.columns):
        if index_2, column_2 in enumerate(data_numeric.columns):
        if index_1 != index_2:
            data_numeric_correlation[index_1, index_2], data_numeric_correlation_r
```

Method - Spearman's Correlation Coefficient

Visualize the Heatmap of Correlation Coefficient

```
annot = True,
    center = 0,
    vmin = -1,
    vmax = 1,
    cmap = 'vlag',
    linecolor = 'gray',
    linewidths = 1 )

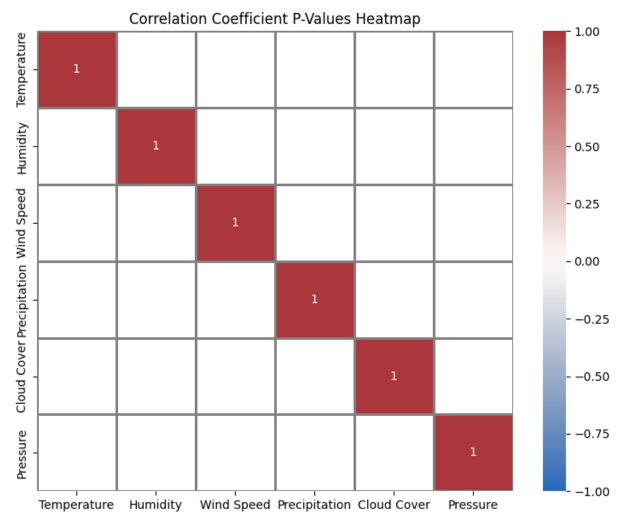
plt.title("Correlation Coefficient Heatmap")
plt.show()
```



Visualize the Heatmap of Correlation Coefficient P-Value

```
cmap = 'vlag',
  linecolor = 'gray',
  linewidths = 1 )

plt.title('Correlation Coefficient P-Values Heatmap')
plt.show()
```



From the two plots above, we observed that all numerical features exhibit no correlation with one another.

2.4.8 - Data Encoding

Data Encoding refers to the process of converting categorical or textual data into numerical format, so that it can be used as input for algorithms to process. The reason for encoding is that most machine learning algorithms work with numbers and not with text or categorical variables.

There are two types of categorical data:

- Nominal Data The categories of data do not have an inherent order. This
 means that the categories cannot be ranked or ordered. For example:
 Occupational titles for doctor, lawyer, instructor, athlete, etc.
- **Ordinal Data** The categories of data have an inherent order. This means that the categories can be ranked or ordered from highest to lowest or vice versa. For example: Grades start with A+, A, A-, B+, B, B-, etc.

Methods of Data Encoding

The choice of encoding method can have a significant impact on model performance, so it is important to choose an appropriate encoding technique based on the nature of the data and the specific requirements of the model.

1. One-Hot Encoding

- Binary column is created for each unique category in the variable. If a category is present in a sample, the corresponding column is set to 1, and all other columns are set to 0
- In the case of one-hot encoding, for N categories in a variable, it uses N binary variables
- For example, if a variable has three categories A, B and C, they can be represented as [1, 0, 0], [0, 1, 0] and [0, 0, 1], respectively

2. **Dummy Encoding**

- Dummy coding scheme is similar to one-hot encoding. This categorical data encoding method transforms the categorical variable into a set of binary variables 0/1
- The dummy encoding is a small improvement over one-hot-encoding. Dummy encoding uses N-1 features to represent N categories
- For example, if a variable has three categories A, B and C, they can be represented as [1, 0] and [0, 1], respectively

3. Binary Encoding

- Similar to one-hot encoding, but instead of creating a separate column for each category, the categories are represented as binary digits
- For example, if a variable has four categories A, B, C and D, they can be represented as 0001, 0010, 0100 and 1000, respectively

4. Label Encoding

- Each unique category is assigned a Unique Integer value
- But the assigned integers may be misinterpreted by the machine learning algorithm as having an ordered relationship when in fact they do not.

• For example, if a variable has four categories A, B, C and D, they can be represented as 0, 1, 2 and 3, respectively

4. Ordinal Encoding

- Ordinal encoding is used when the categories in a variable have an inherent ordering
- The categories are assigned a numerical value based on their order, such as 1, 2, 3, etc.
- For example, if a variable has categories Low, Medium and High, they can be assigned the values 1, 2, and 3, respectively

In sections 2.2 - Data Description and 2.4.1 - Data Cleaning, we identified two features: a textual feature, Location, and a date feature, Date. Since the Date feature does not exhibit a strong relationship with predicting whether it will rain tomorrow, we have decided to discard it. The remaining Location feature consists of the names of major cities in the United States. The goal of this project is to predict rainfall based on the provided weather features. As the Location feature will not be used in the training process, there is no need to encode it.

Method - One-Hot Encoding

Method - Dummy Encoding

```
# Method - Dummy Encoding
#-----
data_encoding_object = pd.DataFrame(data["Location"], columns=["Location"])
data_encoding_object = pd.get_dummies(data_encoding_object["Location"], drop

# Drop the old features
data.drop(columns=["Location"], inplace=True)
# Combination to the original data
data = pd.concat( [data, data_encoding_object], axis=1 )
'''
```

Method - Binary Encoding

```
In []: #-----
# Method - Binary Encoding
#----
data_encoding_object_rank = { city : index for index, city in enumerate(data
    data["Location"] = data["Location"].map(data_encoding_object_rank).apply(lam'')
```

Method - Label Encoding

```
In []: #-----
# Method - Label Encoding
#----
data_encoding_object_rank = { city : index for index, city in enumerate(data
data["Location"] = data["Location"].map(data_encoding_object_rank)
'''
```

Method - Ordinal Encoding

Summary after Data Encoding

```
In [ ]: # Summary the data after encoding
    display(data)
    data.info()
```

2.5 - Exploratory Data Analysis

2.5.1 - Data Visualization Analysis

Data Visualization is an important component of Exploratory Data Analysis (EDA), because it helps us to understand the variables and relationships between them. These variables could be dependent or independent to each other.

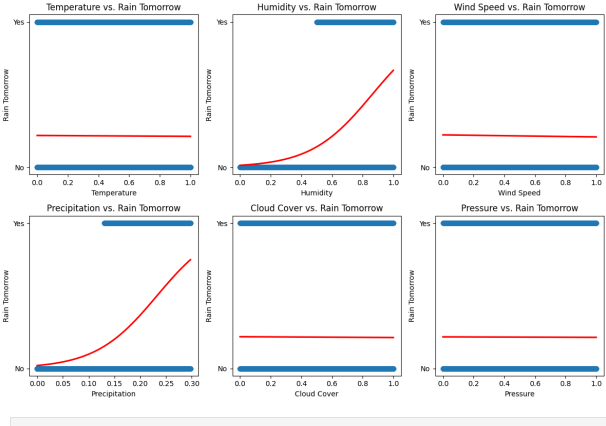
Univariate Analysis	Bivariate Analysis	Multivariate Analysis
It only summarize single variable at a time	It only summarize two variables	It only summarize more than 2 variables
It does not deal with causes and relationships	It does deal with causes and relationships and analysis is done	It does not deal with causes and relationships and analysis is done
The main purpose is to describe	The main purpose is to explain	The main purpose is to study the relationship among them

For this section, we focus on the bivariate analysis to analyzing the relationship between the two variables are positive and negative, or show no clear pattern.

```
# The scatter plots conclude linear fitting line
# Initialization the subplots
_, ax = plt.subplots(nrows=2, ncols=3, figsize=(12, 8))
ax = ax.ravel()

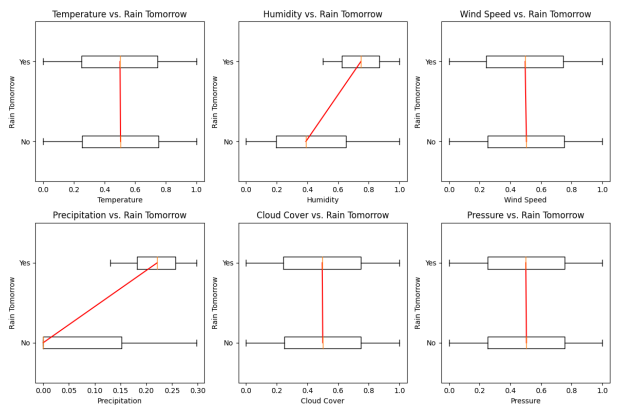
# Draw the scatter plots conclude linear fitting line
for index, column in enumerate(data_numeric.columns):
    sns.regplot(data, ax=ax[index], x=column, y="Rain Tomorrow", line_kws={"cconversed ax[index].set_xlabel(column)
    ax[index].set_ylabel("Rain Tomorrow")
    ax[index].set_yticks([0, 1], ["No", "Yes"])
    ax[index].set_title(column + " vs. Rain Tomorrow")

# Do not blocked any title or label
plt.tight_layout()
plt.show()
```



```
In [ ]: |#-----
        # The boxplots conclude linear fitting line
        # Initialization the subplots
        , ax = plt.subplots(nrows=2, ncols=3, figsize=(12, 8))
        ax = ax.ravel()
        # Get the description by split by target, which is same as pandas.describe()
        data target vs feature = data.groupby("Rain Tomorrow")
        # Draw the boxplots
        for index, column in enumerate(data numeric.columns):
          boxplot data x median, boxplot data y median, boxplot data = [], [], []
          for order, target in data target vs feature:
            boxplot data x median.append(target[column].describe()["50%"])
            boxplot data y median.append(order + 1)
            boxplot data.append(target[column])
          ax[index].boxplot( boxplot data, vert=False )
          ax[index].set xlabel(column)
          ax[index].set ylabel("Rain Tomorrow")
          ax[index].set_yticks(boxplot_data_y_median, ["No", "Yes"])
          ax[index].set title(column + " vs. Rain Tomorrow")
          # linear fitting line
          slope, intercept = np.polyfit(boxplot data x median, boxplot data y median
          ax[index].plot(boxplot_data_x_median, slope * np.array(boxplot_data_x_median)
        # Do not blocked any title or label
```

plt.tight_layout()
plt.show()



From the above plots we have the following analysis.

- Increasing values for the following features are associated with the Rain
 Tomorrow Yes outcome:
 - 1. Humidity
 - 2. Precipitation
- Decreasing values for the following features are associated with the Rain
 Tomorrow No outcome:
 - None
- The values for the following features are not associated with the Rain
 Torrmow Yes or No outcome:
 - 1. Temperature
 - 2. Wind Speed
 - 3. Cloud Cover
 - 4. Pressure

2.5.2 - Hypothesis Testing

In terms of a P-value and a chosen significance level (alpha):

- If P-value <= alpha (usually 5%): significant result, reject null hypothesis (H0), dependent
- If P-value > alpha (usually 5%): not significant result, fail to reject null hypothesis (H0), independent

Hypothesis Testing

We seek to explore whether the occurrence of rain or the absence of rain leads to different weather features values over the past 731 days across 20 major cities in the United States.

To evaluate this, we formulated the following hypotheses:

- **Null (H0)**: The mean values of weather features are consistent, regardless of whether it is a rainy day or not.
- **Alternative (H1)**: The mean values of weather features differ based on the occurrence of rain.

```
In []: # Get the data with type numeric
    data_numeric = data.select_dtypes(include='number')

In []: # Split into two group by Rain Tomorrow 0 or 1
    group_0 = data_numeric[data["Rain Tomorrow"] == 0]
    group_1 = data_numeric[data["Rain Tomorrow"] == 1]

# Calculate the T-test value and P-value by Welch"s t-test
    data_t_test = {}
    for column in data_numeric.columns:
        data_t_test[column] = scipy.stats.ttest_ind(group_0[column], group_1[colum
    # Create a summary table
    data_t_test_summary = data_numeric.from_dict(data_t_test, orient="index", cc
    data_t_test_summary.sort_values(by=["P-Value"], ascending=True)
```

Out[]:		Statistic	P-Value
	Humidity	-137.020240	0.000000
	Precipitation	-176.905844	0.000000
	Wind Speed	2.639905	0.008295
	Temperature	1.115482	0.264648
	Cloud Cover	1.069446	0.284872
	Pressure	0.613900	0.539283

The P-values for the following features are less than 0.05 (5%):

1. Humidity

- 2. Precipitation
- 3. Wind Speed

We reject the null hypothesis of the T-test and conclude that there is sufficient evidence to suggest that the occurrence of rain leads to significant changes in these features.

Conversely, the P-values for the following features are greater than 0.05 (5%):

- 1. Temperature
- 2. Cloud Cover
- 3. Pressure

We do not reject the null hypothesis of the T-test and conclude that there is insufficient evidence to assert that the occurrence of rain affects their values.

3. MODELING

3.1 - Data Splitting

Data Splitting is a crucial process in machine learning, involving the partitioning of a dataset into different subsets, such as training, validation, and test sets. This is essential for training models, tuning parameters, and ultimately assessing their performance.

- **Training Set** Used to train the machine learning model, this is the core dataset where the model learns to understand patterns and relationships in the data
- **Validation Set** Assists in fine-tuning the model. It evaluates the model's performance during the training phase, helping adjust hyperparameters and prevent overfitting
- Test Set Provides a fair evaluation of the model's performance on unseen data. This is crucial for assessing the model's ability to generalize to unknown data

The following are several commonly used methods of data splitting:

- 1. **Random Splitting** Randomly divides the dataset
- 2. **Stratified Splitting** When dealing with imbalanced datasets, Stratified splitting ensures consistency in class distribution
- 3. **Time Series Splitting** Rreservation of chronological order during data partitioning, since the order of data points is crucial, as observations typically depend on previous results
- 4. **K-Fold Cross-Validation** Divides the dataset into K equally sized folds, allowing for multiple rounds of training and validation
- **Standard K-Fold Cross-Validation**: In traditional K-Fold cross-validation, the data set undergoes random partitioning into K folds of roughly equal size. In each iteration, one fold is the validation set, while the remaining K 1 folds constitute the training set. We repeat this procedure K times, ensuring that each fold is used exactly once as the validation set.
- **Stratified K-Fold Cross-Validation**: Stratified K-Fold cross-validation aims to maintain a consistent distribution of classes in each fold, aligning with the overall proportion observed in the data set.
- **Group K-Fold Cross-Validation**: Group K-Fold cross-validation is employed when working with data sets where samples exhibit interdependencies, such as time-series data or data with spatial correlations. This method guarantees that samples from the same group are either entirely within the training or validation set, preventing data leakage across folds.

We use 75% of the data for the train set and the remaining 25% for the test set.

```
In []: # Splitting the data in to X and Y, relanvet to predictor data and target da
data_X = data_numeric
data_Y = data["Rain Tomorrow"]

data_X_train, data_X_test, data_Y_train, data_Y_test = {}, {}, {}, {},
```

Mothod - Random Splitting

```
In []: #-----
# Mothod - Random Splitting
#-----
data_X_train, data_X_test, data_Y_train, data_Y_test = train_test_split(data_Y_train_test_split)
```

Mothod - Stratified Splitting

```
In [ ]: #-----
# Mothod - Stratified Splitting
#-----
```

```
data_X_train, data_X_test, data_Y_train, data_Y_test = train_test_split(data
# OR

for data_train_index, data_test_index in StratifiedShuffleSplit(n_splits=1,
    data_X_train, data_X_test, data_Y_train, data_Y_test = data_X.iloc[data_tr
```

Mothod - Time Series Splitting

Mothod - K-Fold Cross-Validation

```
In [ ]: |#-----
       # Mothod - K-Fold Cross-Validation - Standard
       kf = KFold(n splits=4, shuffle=True)
       kf.get n splits(data X)
       data kfold index = []
       for index, (data_train_index, data_test_index) in enumerate(kf.split(data_X)
        data kfold index.append(index)
        data X train[index], data X test[index], data Y train[index], data Y test[
In [ ]: #-----
       # Mothod - K-Fold Cross-Validation - Stratified
       1.1.1
       skf = StratifiedKFold(n splits=4, shuffle=True)
       skf.get n splits(data X, data Y)
       data kfold index = []
       for index, (data train index, data test index) in enumerate(skf.split(data X
        data kfold index.append(index)
        data X train[index], data X test[index], data Y train[index], data Y test[
In [ ]: | #-----
       # Mothod - K-Fold Cross-Validation - Group
       1.1.1
       # Setting the group variable
       group = np.array([1, 1, 2, 2, 3, 3, 4, 4, 5, 5, 6, 6, 7, 7, 8, 8, 9, 9, 10,
```

```
# We don't have any feature that are dependent correlations, cause we alreay
gkf = GroupKFold(n_splits=4, shuffle=True)
gkf.get_n_splits(data_X, data_Y, group)

data_kfold_index = []
for index, (data_train_index, data_test_index) in enumerate(gkf.split(data_X)
    data_kfold_index.append(index)
    data_X_train[index], data_X_test[index], data_Y_train[index], data_Y_test[index]
```

3.2 - Evaluation Metric

The following metrics are widely used in machine learning to evaluate classifier model performance:

Accuracy

- An metric that measures the proportion of correct predictions made by a model over the total number of predictions made
- Provide a good overall assessment of the model's performance when the classes are balanced
- It can be misleading in imbalanced datasets and does not differentiate between types of errors

Precision

- Precision is the proportion of true positive predictions out of all positive predictions made by the model
- It simply measures the accuracy of positive predictions
- It does not account for false negatives and can be less informative if not considered with recall

Recall

- Recall (sensitivity/true positive rate) is the proportion of true positive predictions from all actual positive samples in the dataset
- It measures the model's ability to identify all positive instances and is critical when the cost of false negatives is high
- It does not account for false positives and can be less informative if not considered with precision

F1 Score

- The F1 score is a measure of a model's accuracy that takes into account both precision and recall, where the goal is to classify instances correctly as positive or negative
- High recall score means the model has a low rate of false negatives, and it useful in imbalanced datasets

It does not account for true negative rates

3.3 - Hyperparameter Tuning

Hyperparameter Tuning is the process of selecting the optimal values for a machine learning model's hyperparameters. **Hyperparameters** are configuration variables that are set before the training process of a model begins. They control the learning process itself, rather than being learned from the data. Hyperparameters are often used to tune the performance of a model, and they can have a significant impact on the model's accuracy, generalization, and other metrics.

Grid Search can be considered as a brute force approach to hyperparameter optimization. We fit the model using all possible combinations after creating a grid of potential discrete hyperparameter values. We log each set's model performance and then choose the combination that produces the best results. This approach is called GridSearchCV, because it searches for the best set of hyperparameters from a grid of hyperparameters values.

- **Learning rate**: This hyperparameter controls the step size taken by the optimizer during each iteration of training. Too small a learning rate can result in slow convergence, while too large a learning rate can lead to instability and divergence.
- **Epochs**: This hyperparameter represents the number of times the entire training dataset is passed through the model during training. Increasing the number of epochs can improve the model's performance but may lead to overfitting if not done carefully.
- **Number of layers**: This hyperparameter determines the depth of the model, which can have a significant impact on its complexity and learning

ability.

- **Number of nodes per layer**: This hyperparameter determines the width of the model, influencing its capacity to represent complex relationships in the data.
- **Architecture**: This hyperparameter determines the overall structure of the neural network, including the number of layers, the number of neurons per layer, and the connections between layers. The optimal architecture depends on the complexity of the task and the size of the dataset.
- **Activation function**: This hyperparameter introduces non-linearity into the model, allowing it to learn complex decision boundaries. Common activation functions include sigmoid, tanh, and Rectified Linear Unit (ReLU).

An exhaustive approach that can identify the ideal hyperparameter combination is grid search. But the slowness is a disadvantage. It often takes a lot of processing power and time to fit the model with every potential combination, which might not be available.

3.3.1 - Logistic Regression

Hyperparameter:

- **C**: Inverse of regularization strength, must be a positive float, and smaller values specify stronger regularization
- **Penalty**: None means no penalty is added; l1 means add a L1 penalty term; l2 means add a L2 penalty term
- **Solver**: Algorithm to use in the optimization problem; For small datasets, liblinear is a good choice, whereas sag and saga are faster for large ones; For multiclass problems, only newton-cg, ag, saga and lbfgs handle multinomial loss

Optimal Values of Hyperparameter Tuning

```
In [ ]: for name, best in model_lr.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best
```

```
The optimal value of 'C' hyperparameter is '5.5'
The optimal value of 'penalty' hyperparameter is 'l2'
The optimal value of 'solver' hyperparameter is 'liblinear'
```

Prediction and Evaluation on Training Set

```
      Out[]:
      Metric
      Value

      0 Accuracy
      0.938254

      1 Precision
      0.869007

      2 Recall
      0.839821

      3 F1 Score
      0.854165
```

Prediction and Evaluation on Test Set

```
      Out[]:
      Metric
      Value

      0 Accuracy
      0.940201

      1 Precision
      0.876211

      2 Recall
      0.841126

      3 F1 Score
      0.858310
```

3.3.2 - K-Nearest Neighbors (KNN)

Hyperparameter:

• **N Neighbors**: Number of neighbors required for each sample

- **Algorithm**: Algorithm used to compute the nearest neighbors. ball_tree will use BallTree; kd_tree will use KDTree; auto will attempt to decide the most appropriate algorithm based on the values passed to fit method
- **Metric**: Metric to use for distance computation. minkowski means standard euclidean distance; manhattan means manhattan distance

Optimal Values of Hyperparameter Tuning

```
In [ ]: for name, best in model_knn.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best
    The optimal value of 'algorithm' hyperparameter is 'auto'
    The optimal value of 'metric' hyperparameter is 'manhattan'
    The optimal value of 'n neighbors' hyperparameter is '7'
```

Prediction and Evaluation on Training Set

```
      Out[]:
      Metric
      Value

      0 Accuracy
      0.985426

      1 Precision
      0.967525

      2 Recall
      0.964692

      3 F1 Score
      0.966106
```

Prediction and Evaluation on Test Set

```
In [ ]: #-----
# K-Nearest Neighbors (KNN) - Test Set Prediction
#-----
```

```
y_hat = model_knn.predict(data_X_test)

classifier_model_scores["Test"]["K-Nearest Neighbors"] = evaluation_classifi

classifier_model_scores["Test"]["K-Nearest Neighbors"]
```

```
      Out[]:
      Metric
      Value

      0 Accuracy
      0.976247

      1 Precision
      0.951021

      2 Recall
      0.938001

      3 F1 Score
      0.944466
```

3.3.3 - Support Vector Machine (SVM)

Hyperparameter:

- **C**: Inverse of regularization strength, must be a positive float, and smaller values specify stronger regularization
- **Kernel**: Specifies the kernel type to be used in the algorithm. linear means linear function; poly means polynomial function; rbf means radial basis function; sigmoid means hyperbolic tangent function (tanh)

Optimal Values of Hyperparameter Tuning

```
In [ ]: for name, best in model_svm.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best
    The optimal value of 'C' hyperparameter is '1.0'
    The optimal value of 'kernel' hyperparameter is 'poly'
```

Prediction and Evaluation on Training Set

```
In []: #-----
# Support Vector Machine (SVM) - Train Set Prediction
#-----
y_hat = model_svm.predict(data_X_train)
```

```
classifier_model_scores["Train"]["Support Vector Machine"] = evaluation_clas
classifier_model_scores["Train"]["Support Vector Machine"]
```

```
    Out[]: Metric Value
    O Accuracy 0.987354
    Precision 0.976045
    Recall 0.964950
    F1 Score 0.970466
```

Prediction and Evaluation on Test Set

```
    Out[]: Metric Value
    O Accuracy 0.987317
    Precision 0.978461
    Recall 0.962284
    F1 Score 0.970305
```

3.3.4 - Decision Tree

Hyperparameter:

- **Criterion**: The function to measure the quality of a split. gini for gini impurity method; entropy for entropy method
- Max Depth: The maximum depth of the tree. None means the nodes are expanded until all leaves are pure or until all leaves contain less than min samples split (default=2) samples
- **Min Samples Leaf**: The minimum number of samples required to be at a leaf node. A split point at any depth will only be considered if it leaves at least min_samples_leaf training samples in each of the left and right branches

Optimal Values of Hyperparameter Tuning

```
In [ ]: for name, best in model_dt.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'criterion' hyperparameter is 'gini'
    The optimal value of 'max_depth' hyperparameter is 'None'
    The optimal value of 'min_samples_leaf' hyperparameter is '1'
```

Prediction and Evaluation on Training Set

```
      Out[]:
      Metric Value

      0
      Accuracy
      1.0

      1
      Precision
      1.0

      2
      Recall
      1.0

      3
      F1 Score
      1.0
```

Prediction and Evaluation on Test Set

Out[]:		Metric	Value
		0	Accuracy	1.0
		1	Precision	1.0
		2	Recall	1.0
		3	F1 Score	1.0

3.3.5 - Random Forest

Hyperparameter:

- **N Estimators**: The number of trees in the forest
- **Criterion**: The function to measure the quality of a split. gini for gini impurity method; entropy for entropy method
- Max Depth: The maximum depth of the tree. None means the nodes are expanded until all leaves are pure or until all leaves contain less than min samples split (default=2) samples
- Min Samples Leaf: The minimum number of samples required to be at a leaf node. A split point at any depth will only be considered if it leaves at least min_samples_leaf training samples in each of the left and right branches
- Max Features: The number of features to consider best split. sqrt means N features square root; log2 means N features Log 2

Optimal Values of Hyperparameter Tuning

```
In [ ]: for name, best in model_rf.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best
```

```
The optimal value of 'criterion' hyperparameter is 'gini'
The optimal value of 'max_depth' hyperparameter is 'None'
The optimal value of 'max_features' hyperparameter is 'sqrt'
The optimal value of 'min_samples_leaf' hyperparameter is '1'
The optimal value of 'n estimators' hyperparameter is '100'
```

Prediction and Evaluation on Training Set

Out[]: Metric Value 0 Accuracy 1.0 1 Precision 1.0 2 Recall 1.0 3 F1 Score 1.0

Prediction and Evaluation on Test Set

```
        Out[]:
        Metric
        Value

        0
        Accuracy
        1.0

        1
        Precision
        1.0

        2
        Recall
        1.0

        3
        F1 Score
        1.0
```

3.4 - Hyperparameter Tuning with Downsampling

Downsampling decreases the number of data samples in a dataset. In doing so, it aims to correct imbalanced data and thereby improve model performance. The process of downsampling counteracts the imbalanced dataset issue. It identifies majority class points to remove based on specified criteria. These criteria can change with the chosen downsampling technique. This balances the dataset by effectively decreasing the number of samples for an overrepresented majority class until the dataset contains an equal ratio of points across all classes.

Advantages:

- 1. **Less Storage Requirement**: When storage costs money, say for cloud storage, downsampling would be preferred over upsampling to avoid raising costs.
- 2. **Faster Training**: Downsampling shrinks datasets and makes training less intensive on the CPU or GPU, which is more economically and environmentally friendly.
- 3. **Less Prone to Overfitting**: Upsampling generates new data from the old data, which can cause models to overfit to the given data. Downsampling, being the opposite (deletes data), doesn't suffer from this issue.

Disadvantages:

- 1. **Loss of Information**: Deleting points from the majority class can cause important information loss. This can be an issue if the classification of the majority class needs to be accurate. Another issue is if the dataset becomes too small for the model to train on.
- 2. **Introduced Bias**: The remaining majority class sample points can be a biased set of the original data, which negatively affects the classifier's performance.

Downsampling usually use following techniques:

Random Downsampling

- Random points in the majority class are chosen deleted from the dataset until the majority class size is equal to the minority class size
- **BUT**, this technique can cause important patterns or distributions in the majority class to disappear, negatively affecting classifier performance

Near Miss Downsampling

 Operates on the principle that data should be kept in places where the majority and minority classes are very close, as these places give us key information in distinguishing the two classes

Condensed Nearest Neighbor (CNN) Downsampling

- Find a subset of a dataset that can be used for training without loss in model performance. This is achieved by identifying a subset of the data that can be used to train a model that correctly predicts the entire dataset
- Like **Near Miss**, this process essentially removes all majority class instances far away from the decision boundary, which, again, are points that are easy to classify. It also ensures that every data in our original dataset can be correctly predicted using just the data within dataset.

Method - Random Downsampling

```
In [ ]: |#-----
      # Method - Random Downsampling
       #-----
       data X train downsampling, data Y train downsampling = RandomUnderSampler().
      # 0R
       # Calculate how many rows we would need to drop
       data ratio = data Y train.value counts(normalize = True)
       data needs to down = math.floor( round(abs(data ratio[0] - data ratio[1]), 2
       try:
        while data needs to down != 0:
          random index = random.choice(data Y train.index)
          if data Y train.iloc[random index] == 0:
            data X train.drop(random index, inplace=True)
            data Y train.drop(random index, inplace=True)
            data needs to down -= 1
       except:
        print("Please try other way.")
```

Method - Near Miss Downsampling

```
In []: #----
# Method - Near Miss Downsampling
#-----
data_X_train_downsampling, data_Y_train_downsampling = NearMiss().fit_resamp
```

Method - Condensed Nearest Neighbor (CNN) Downsampling

```
data_X_train_downsampling, data_Y_train_downsampling = CondensedNearestNeigh
'''
```

Summary after Downsampling

```
-----
In [ ]: #-----
       # The summary of downsampling
        print("Before downsampling we have total observations of data:")
        print(data Y train.value counts())
        print()
        print("Before downsampling we have normalize ratio of data:")
        print(data Y train.value counts(normalize = True))
       print()
        print("After downsampling we have total observations of data:")
        print(data Y train downsampling.value counts())
        print()
        print("After downsampling we have normalize ratio of data:")
       print(data Y train downsampling.value counts(normalize = True))
      Before downsampling we have total observations of data:
      Rain Tomorrow
           42319
      0
      1
           11612
      Name: count, dtype: int64
      Before downsampling we have normalize ratio of data:
      Rain Tomorrow
           0.784688
           0.215312
      1
      Name: proportion, dtype: float64
      After downsampling we have total observations of data:
      Rain Tomorrow
      0
           11612
      1
           11612
      Name: count, dtype: int64
      After downsampling we have normalize ratio of data:
      Rain Tomorrow
           0.5
      0
      1
           0.5
      Name: proportion, dtype: float64
```

3.4.1 - Logistic Regression

Hyperparameter:

- **C**: Inverse of regularization strength, must be a positive float, and smaller values specify stronger regularization
- **Penalty**: None means no penalty is added; l1 means add a L1 penalty term; l2 means add a L2 penalty term
- **Solver**: Algorithm to use in the optimization problem; For small datasets, liblinear is a good choice, whereas sag and saga are faster for large ones; For multiclass problems, only newton-cg, ag, saga and lbfgs handle multinomial loss

Optimal Values of Hyperparameter Tuning

```
In [ ]: for name, best in model_lr_down.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'C' hyperparameter is '1.0'
    The optimal value of 'penalty' hyperparameter is 'l2'
    The optimal value of 'solver' hyperparameter is 'liblinear'
```

Prediction and Evaluation on Training Set

```
      Out[]:
      Metric
      Value

      0 Accuracy
      0.933656

      1 Precision
      0.827597

      2 Recall
      0.873924

      3 F1 Score
      0.850130
```

Prediction and Evaluation on Test Set

3.4.2 - K-Nearest Neighbors (KNN)

Hyperparameter:

- **N Neighbors**: Number of neighbors required for each sample
- Algorithm: Algorithm used to compute the nearest neighbors. ball_tree will use BallTree; kd_tree will use KDTree; auto will attempt to decide the most appropriate algorithm based on the values passed to fit method
- **Metric**: Metric to use for distance computation. minkowski means standard euclidean distance; manhattan means manhattan distance

Optimal Values of Hyperparameter Tuning

```
In [ ]: for name, best in model_knn_down.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'algorithm' hyperparameter is 'auto'
    The optimal value of 'metric' hyperparameter is 'manhattan'
    The optimal value of 'n_neighbors' hyperparameter is '7'
```

Prediction and Evaluation on Training Set

Out[]:		Metric	Value
		0	Accuracy	0.981736
		1	Precision	0.948586
		2	Recall	0.967620
		3	F1 Score	0.958008

Prediction and Evaluation on Test Set

```
      Out[]:
      Metric
      Value

      0 Accuracy
      0.973466

      1 Precision
      0.925740

      2 Recall
      0.953242

      3 F1 Score
      0.939290
```

3.4.3 - Support Vector Machine (SVM)

Hyperparameter:

• **C**: Inverse of regularization strength, must be a positive float, and smaller values specify stronger regularization

• **Kernel**: Specifies the kernel type to be used in the algorithm. linear means linear function; poly means polynomial function; rbf means radial basis function; sigmoid means hyperbolic tangent function (tanh)

Optimal Values of Hyperparameter Tuning

```
In [ ]: for name, best in model_svm_down.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'C' hyperparameter is '1.0'
    The optimal value of 'kernel' hyperparameter is 'poly'
```

Prediction and Evaluation on Training Set

```
      Out[]:
      Metric
      Value

      0 Accuracy
      0.987836

      1 Precision
      0.976182

      2 Recall
      0.967103

      3 F1 Score
      0.971621
```

Prediction and Evaluation on Test Set

3.4.4 - Decision Tree

Hyperparameter:

- **Criterion**: The function to measure the quality of a split. gini for gini impurity method; entropy for entropy method
- Max Depth: The maximum depth of the tree. None means the nodes are expanded until all leaves are pure or until all leaves contain less than min samples split (default=2) samples
- Min Samples Leaf: The minimum number of samples required to be at a leaf node. A split point at any depth will only be considered if it leaves at least min_samples_leaf training samples in each of the left and right branches

Optimal Values of Hyperparameter Tuning

```
In [ ]: for name, best in model_dt_down.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'criterion' hyperparameter is 'gini'
    The optimal value of 'max_depth' hyperparameter is 'None'
    The optimal value of 'min samples leaf' hyperparameter is '1'
```

Prediction and Evaluation on Training Set

Prediction and Evaluation on Test Set

1.0

1.0

```
        Out[]:
        Metric
        Value

        0
        Accuracy
        1.0

        1
        Precision
        1.0

        2
        Recall
        1.0

        3
        F1 Score
        1.0
```

2

Recall

3 F1 Score

3.4.5 - Random Forest

Hyperparameter:

- **N Estimators**: The number of trees in the forest
- **Criterion**: The function to measure the quality of a split. gini for gini impurity method; entropy for entropy method
- **Max Depth**: The maximum depth of the tree. None means the nodes are expanded until all leaves are pure or until all leaves contain less than

min samples split (default=2) samples

- **Min Samples Leaf**: The minimum number of samples required to be at a leaf node. A split point at any depth will only be considered if it leaves at least min_samples_leaf training samples in each of the left and right branches
- Max Features: The number of features to consider best split. sqrt means N features square root; log2 means N features Log 2

Optimal Values of Hyperparameter Tuning

```
In []: for name, best in model_rf_down.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'criterion' hyperparameter is 'gini'
    The optimal value of 'max_depth' hyperparameter is 'None'
    The optimal value of 'max_features' hyperparameter is 'sqrt'
    The optimal value of 'min_samples_leaf' hyperparameter is '1'
    The optimal value of 'n_estimators' hyperparameter is '150'
```

Prediction and Evaluation on Training Set

```
      Out[]:
      Metric
      Value

      0
      Accuracy
      1.0

      1
      Precision
      1.0

      2
      Recall
      1.0

      3
      F1 Score
      1.0
```

Prediction and Evaluation on Test Set

Out[]:		Metric	Value
		0	Accuracy	1.0
		1	Precision	1.0
		2	Recall	1.0
		3	F1 Score	1.0

3.5 - Hyperparameter Tuning with Upsampling

Upsampling increases the number of data samples in a dataset. In doing so, it aims to correct imbalanced data and thereby improve model performance. The process of upsampling counteracts the imbalanced dataset issue. It populates the dataset with points synthesized from characteristics of the original dataset's minority class. This balances the dataset by effectively increasing the number of samples for an underrepresented minority class until the dataset contains an equal ratio of points across all classes.

Advantages:

• **No Information Loss**: Unlike downsampling, which removes data points from the majority class, upsampling generates new data points, avoiding any information loss.

• Increase Data at Low Costs: Upsampling is especially effective, and is often the only way, to increase dataset size on demand in cases where data can only be acquired through observation. For instance, certain medical conditions are simply too rare to allow for more data to be collected.

Disadvantages:

- **Overfitting**: Because upsampling creates new data based on the existing minority class data, the classifier can be overfitted to the data. Upsampling assumes that the existing data adequately captures reality; if that is not the case, the classifier may not be able to generalize very well.
- **Data Noise**: Upsampling can increase the amount of noise in the data, reducing the classifier's reliability and performance.2
- **Computational Complexity**: By increasing the amount of data, training the classifier will be more computational expensive, which can be an issue when using cloud computing.2

Upsampling usually use following techniques:

Random Upsampling

- Process of duplicating random data points in the minority class until the size of the minority class is equal to the majority class
- Its ease of implementation, lack of stretching assumptions about the data, and low time complexity due to a simple algorithm
- It has limitations, however. Because random oversampling solely adds duplicate datapoints, it can lead to overfitting

Synthetic Minority Oversampling Technique (SMOTE)

- A statistical technique for increasing the number of cases in dataset in a balanced way. The component works by generating new instances from existing minority cases that you supply as input
- The new instances are not just copies of existing minority cases. Instead, the algorithm takes samples of the feature space for each target class and its nearest neighbors. The algorithm then generates new examples that combine features of the target case with features of its neighbors

Borderline SMOTE

- Used to combat the issue of artificial dataset noise and to create 'harder' data points. 'Harder' data points are data points close to the decision boundary, and therefore harder to classify
- Identifies the minority class points that are close to many majority class points and puts them into a DANGER set. DANGER points are the 'hard' data points to learn, which again is because they're harder to classify compared to points that are surrounded by minority class points

Method - Random Upsampling

```
# # Method - Random Upsampling
# 
data_X_train_upsampling, data_Y_train_upsampling = RandomOverSampler().fit_r
# OR

# Calculate how many rows we would need to drop
data_ratio = data_Y_train.value_counts(normalize = True)
data_needs_to_up = math.floor( round(abs(data_ratio[0] - data_ratio[1]), 2)

try:
    while data_needs_to_up != 0:
        random_index = random.choice(data_Y_train.index)
        if data_Y_train.iloc[random_index] == 1:
            data_X_train.loc[len(data_X_train.index)] = data_X_train.iloc[random_index]
```

Method - Synthetic Minority Oversampling Technique (SMOTE)

Method - Borderline SMOTE

```
In []: #-----
# Method - Borderline SMOTE
#----
data_X_train_upsampling, data_Y_train_upsampling = BorderlineSMOTE().fit_res
'''
```

Summary after Upsampling

```
Before upsampling we have total observations of data:
Rain Tomorrow
0
  42319
1
    11612
Name: count, dtype: int64
Before upsampling we have normalize ratio of data:
Rain Tomorrow
    0.784688
1 0.215312
Name: proportion, dtype: float64
After upsampling we have total observations of data:
Rain Tomorrow
    42319
1
    42319
Name: count, dtype: int64
After upsampling we have normalize ratio of data:
Rain Tomorrow
    0.5
0
     0.5
1
Name: proportion, dtype: float64
```

3.5.1 - Logistic Regression

Hyperparameter:

- **C**: Inverse of regularization strength, must be a positive float, and smaller values specify stronger regularization
- **Penalty**: None means no penalty is added; l1 means add a L1 penalty term; l2 means add a L2 penalty term
- **Solver**: Algorithm to use in the optimization problem; For small datasets, liblinear is a good choice, whereas sag and saga are faster for large ones; For multiclass problems, only newton-cg, ag, saga and lbfgs handle multinomial loss

Optimal Values of Hyperparameter Tuning

```
In []: for name, best in model_lr_up.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'C' hyperparameter is '1.0'
    The optimal value of 'penalty' hyperparameter is 'l2'
    The optimal value of 'solver' hyperparameter is 'saga'
```

Prediction and Evaluation on Training Set

```
      Out[]:
      Metric
      Value

      0 Accuracy
      0.925368

      1 Precision
      0.771955

      2 Recall
      0.927317

      3 F1 Score
      0.842534
```

Prediction and Evaluation on Test Set

3.5.2 - K-Nearest Neighbors (KNN)

Hyperparameter:

- **N Neighbors**: Number of neighbors required for each sample
- **Algorithm**: Algorithm used to compute the nearest neighbors. ball_tree will use BallTree; kd_tree will use KDTree; auto will attempt to decide the most appropriate algorithm based on the values passed to fit method
- **Metric**: Metric to use for distance computation. minkowski means standard euclidean distance; manhattan means manhattan distance

Optimal Values of Hyperparameter Tuning

```
In []: for name, best in model_knn_up.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'algorithm' hyperparameter is 'auto'
    The optimal value of 'metric' hyperparameter is 'manhattan'
    The optimal value of 'n_neighbors' hyperparameter is '3'
```

Prediction and Evaluation on Training Set

```
      Out[]:
      Metric
      Value

      0 Accuracy
      0.980030

      1 Precision
      0.916041

      2 Recall
      0.998794

      3 F1 Score
      0.955630
```

Prediction and Evaluation on Test Set

3.5.3 - Support Vector Machine (SVM)

Hyperparameter:

3 F1 Score 0.908762

- **C**: Inverse of regularization strength, must be a positive float, and smaller values specify stronger regularization
- **Kernel**: Specifies the kernel type to be used in the algorithm. linear means linear function; poly means polynomial function; rbf means radial basis function; sigmoid means hyperbolic tangent function (tanh)

Optimal Values of Hyperparameter Tuning

```
In [ ]: for name, best in model_svm_up.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'C' hyperparameter is '1.0'
    The optimal value of 'kernel' hyperparameter is 'poly'
```

Prediction and Evaluation on Training Set

Prediction and Evaluation on Test Set

3.5.4 - Decision Tree

Hyperparameter:

- **Criterion**: The function to measure the quality of a split. gini for gini impurity method; entropy for entropy method
- Max Depth: The maximum depth of the tree. None means the nodes are expanded until all leaves are pure or until all leaves contain less than min_samples_split (default=2) samples

 Min Samples Leaf: The minimum number of samples required to be at a leaf node. A split point at any depth will only be considered if it leaves at least min_samples_leaf training samples in each of the left and right branches

Optimal Values of Hyperparameter Tuning

```
In []: for name, best in model_dt_up.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'criterion' hyperparameter is 'gini'
    The optimal value of 'max_depth' hyperparameter is 'None'
    The optimal value of 'min samples leaf' hyperparameter is '1'
```

Prediction and Evaluation on Training Set

```
      Out[]:
      Metric
      Value

      0 Accuracy
      1.0

      1 Precision
      1.0

      2 Recall
      1.0

      3 F1 Score
      1.0
```

Prediction and Evaluation on Test Set

```
In [ ]: #-----
# Decision Tree - Test Set Prediction
#-----
```

```
y_hat = model_dt_up.predict(data_X_test)

classifier_model_scores["Test"]["Upsampling: Decision Tree"] = evaluation_cl

classifier_model_scores["Test"]["Upsampling: Decision Tree"]
```

```
        Out[]:
        Metric
        Value

        0 Accuracy
        0.999944

        1 Precision
        1.000000

        2 Recall
        0.999742

        3 F1 Score
        0.999871
```

3.5.5 - Random Forest

Hyperparameter:

- **N Estimators**: The number of trees in the forest
- **Criterion**: The function to measure the quality of a split. gini for gini impurity method; entropy for entropy method
- Max Depth: The maximum depth of the tree. None means the nodes are expanded until all leaves are pure or until all leaves contain less than min samples split (default=2) samples
- Min Samples Leaf: The minimum number of samples required to be at a leaf node. A split point at any depth will only be considered if it leaves at least min_samples_leaf training samples in each of the left and right branches
- Max Features: The number of features to consider best split. sqrt means N features square root; log2 means N features Log 2

Optimal Values of Hyperparameter Tuning

```
In []: for name, best in model_rf_up.best_params_.items():
    print("The optimal value of '" + name + "' hyperparameter is '" + str(best

The optimal value of 'criterion' hyperparameter is 'gini'
The optimal value of 'max_depth' hyperparameter is 'None'
The optimal value of 'max_features' hyperparameter is 'sqrt'
The optimal value of 'min_samples_leaf' hyperparameter is '1'
The optimal value of 'n_estimators' hyperparameter is '100'
```

Prediction and Evaluation on Training Set

Out[]: Metric Value 0 Accuracy 1.0 1 Precision 1.0 2 Recall 1.0 3 F1 Score 1.0

Prediction and Evaluation on Test Set

Out[]: Metric Value 0 Accuracy 1.0 1 Precision 1.0 2 Recall 1.0 3 F1 Score 1.0

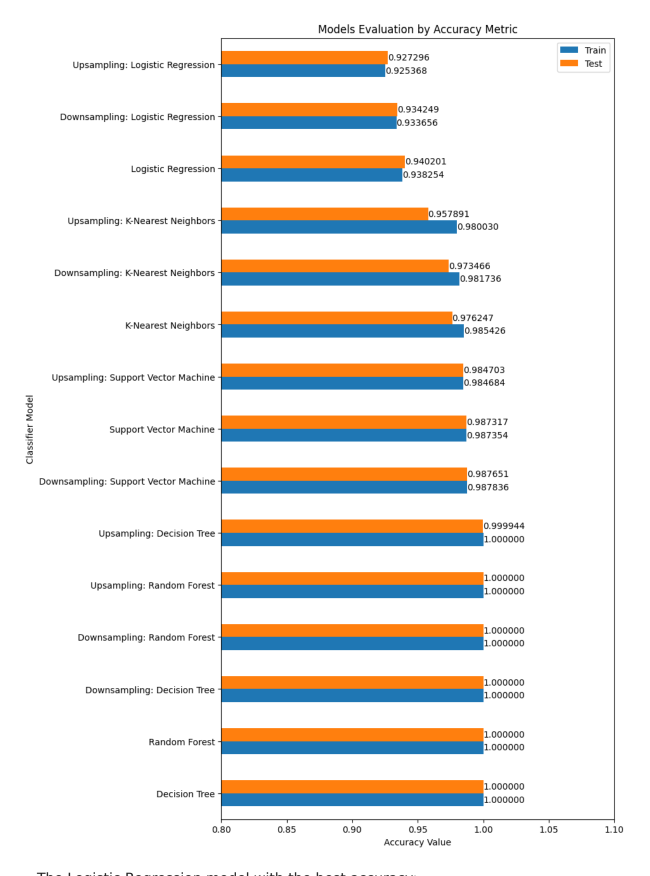
4. SUMMARY

4.1 - Models Evaluation

We will now compare the prediction results of various classifier models on both the training and test sets. The most suitable classification model for our project will be selected based on metrics such as Accuracy, Precision, Recall, and F1 Score.

```
In [ ]: # Split the metric values of train set and test set
model_scores_train = pd.concat( classifier_model_scores["Train"] )
model_scores_test = pd.concat( classifier_model_scores["Test"] )
```

Model Evaluation Based on the Accuracy Metric



The Logistic Regression model with the best accuracy:

- Logistic Regression
 - 93.8254 % on training set
 - 94.0201 % on test set

The K-Nearest Neighbors model with the best accuracy:

- K-Nearest Neighbors
 - 98.5426 % on training set
 - 97.6247 % on test set

The Support Vector Machine model with the best accuracy:

- Downsampling: Support Vector Machine
 - 98.7836 % on training set
 - 98.7651 % on test set

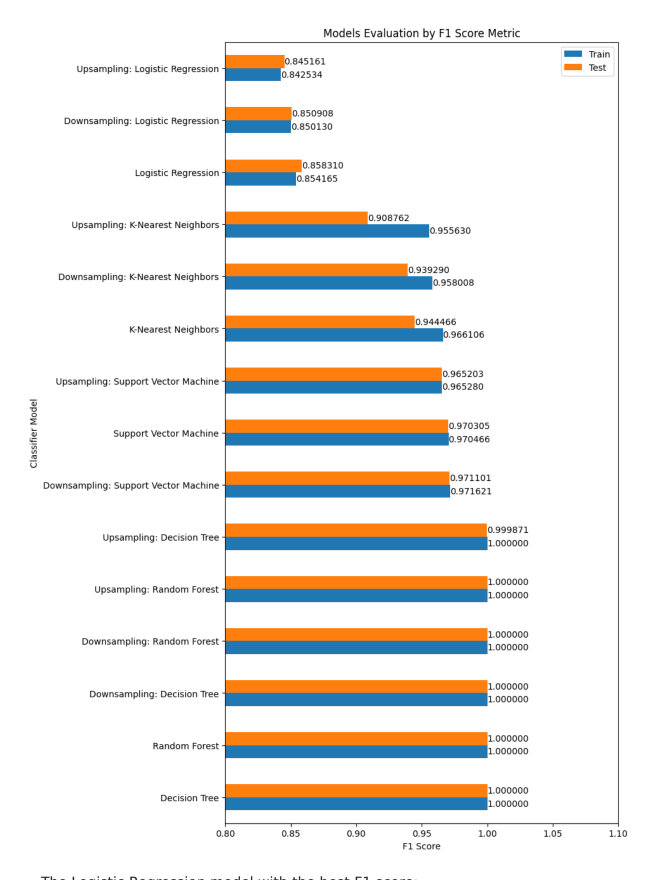
The Decision Tree model with the best accuracy:

- Decision Tree and Downsampling: Decision Tree
 - 100 % on training set
 - 100 % on test set

The Random Forest model with the best accuracy:

- Random Forest, Downsampling: Random Forest, and Upsampling: Random Forest
 - 100 % on training set
 - 100 % on test set

Model Evaluation Based on the F1 Score Metric



The Logistic Regression model with the best F1 score:

- Logistic Regression
 - 0.854165 on training set
 - 0.85831 on test set

The K-Nearest Neighbors model with the best F1 score:

- K-Nearest Neighbors
 - 0.966106 on training set
 - 0.944466 on test set

The Support Vector Machine model with the best F1 score:

- Downsampling: Support Vector Machine
 - 0.971621 on training set
 - 0.971101 on test set

The Decision Tree model with the best F1 score:

- Decision Tree and Downsampling: Decision Tree
 - 1.0 on training set
 - 1.0 on test set

The Random Forest model with the best F1 score:

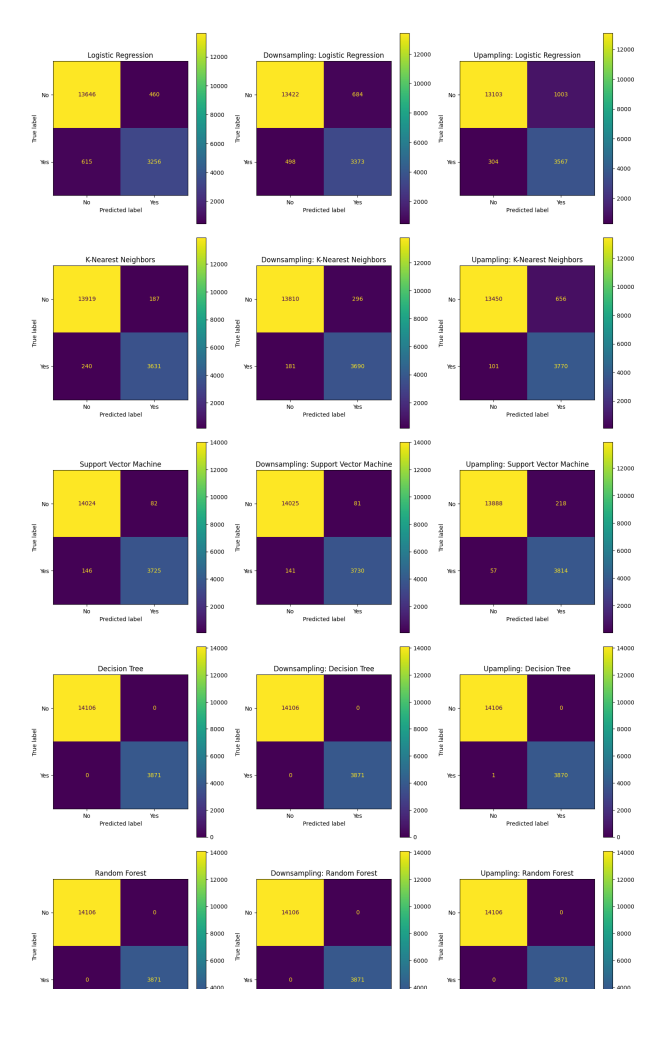
- Random Forest, Downsampling: Random Forest, and Upsampling: Random Forest
 - 1.0 on training set
 - 1.0 on test set

Model Evaluation by Confusion Matrix on the Test Set

```
In [ ]: model scores confusion matrices = { "Logistic Regression" : model lr,
                            "Downsampling: Logistic Regression" : model lr down,
                            "Upampling: Logistic Regression" : model lr up,
                            "K-Nearest Neighbors" : model knn,
                            "Downsampling: K-Nearest Neighbors" : model knn down,
                            "Upampling: K-Nearest Neighbors" : model knn up,
                            "Support Vector Machine" : model svm,
                            "Downsampling: Support Vector Machine" : model svm down,
                            "Upampling: Support Vector Machine" : model svm up,
                            "Decision Tree" : model dt,
                            "Downsampling: Decision Tree" : model dt down,
                            "Upampling: Decision Tree" : model_dt_up,
                            "Random Forest" : model rf,
                            "Downsampling: Random Forest" : model rf down,
                            "Upampling: Random Forest" : model rf up }
        _, ax = plt.subplots(nrows=5, ncols=3, figsize=(15, 25))
        ax = ax.ravel()
        for index, (name, model) in enumerate(model scores confusion matrices.items(
```

```
cmd = ConfusionMatrixDisplay(confusion_matrix(data_Y_test, model.predict(c
cmd.plot(ax=ax[index])
cmd.ax_.set_title(name)

plt.tight_layout()
plt.show()
```



From the confusion matrix we get the following information:

Model	Original	Downsampling	Upsampling
Logistic Regression	1075 Failed	1182 Failed	1307 Failed
K-Nearest Neighbors	427 Failed	477 Failed	757 Failed
Support Vector Machine	228 Failed	222 Failed	275 Failed
Decision Tree	0 Failed	0 Failed	1 Failed
Random Forest	0 Failed	0 Failed	0 Failed

The Logistic Regression model with the better performance on confusion matrix:

• Logistic Regression: 1075 Failed

The K-Nearest Neighbors model with the better performance on confusion matrix:

K-Nearest Neighbors: 427 Failed

The Support Vector Machine model with the better performance on confusion matrix:

• Downsampling: Support Vector Machine: 222 Failed

The Decision Tree model with the better performance on confusion matrix:

Decision Tree and Downsampling: Decision Tree: 0 Failed

The Random Forest model with the better performance on confusion matrix:

 Random Forest, Downsampling: Random Forest, and Upsampling: Random Forest: 0 Failed

4.2 - Summary

Based on the analysis in the preceding sections, the most suitable model for our project is the random forest model. Decision trees and random forests performed better than other classifier models in this project, both on imbalanced data and on balanced data after upsampling and downsampling.

However, two important considerations arise:

- Given the characteristics of decision trees and random forests, both models
 may be susceptible to overfitting. When the data is concentrated and the
 dataset is split using stratified sampling, the training and test sets can
 display similar distributions. This similarity can exacerbate overfitting in treebased models, especially if the overall dataset lacks sufficient diversity.
- 2. Many classification models perform better with downsampling than with upsampling, demonstrating that in cases of unbalanced and non-generalized datasets, reducing repeated observations can significantly lower training costs while preserving high model performance.

In the future, we should consider adding data from other cities as test sets to improve the generalization of the dataset, thereby enhancing the validation of the classifier model outcomes. Since our data is sourced from 20 major cities in the United States, the model may have become overly familiar with the weather characteristics of these specific locations.

5. REFERENCES

Sources	Article	Author
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Statology	How to Perform t-Tests in Pandas	Zach Bobbitt
Deepchecks	Understanding F1 Score, Accuracy, ROC-AUC, and PR-AUC Metrics for Models	Community Blog

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