CS5483 Data Warehousing and Data Mining Project

Author: Group1**Author: group1

Table of Contents

- Abstract
- Part1. Introduction
- Part2. Data Process and Description
 - 2.1 Pre-processing
 - 2.2 Preliminary data analysis
 - 2.3 Variable correlation analysis
- Part3. Model
 - 3.1 Normalization
 - 3.2 Feature Extraction
 - 3.3 Dimensionality reduction
- · Part4. Model
 - 4.1 Random Forest
 - 4.2 Naive Bayes
 - 4.3 InceptionV3 Model
 - 4.4 CNN Neural Network
 - 4.4.1 Optimizer & Loss Function
 - 4.4.2 Training our CNN Model
 - 4.4.3 Check the performance of our CNN model
 - 4.4.4 Look at the wrong images
 - 4.4.5 Hyper-parameter tuning
 - 4.4.6 Boarder Deeper Network
- Part5. Conclusion

Abstract

In data loading and data processing, we first preprocess the data to eliminate duplicate and abnormal data.
 Then, in the data description part, the variables are preliminarily described, and the correlation between variables is simply analyzed

Part 1. Introduction

• According to the CDC, heart disease is one of the leading causes of death for people of most races in the US (African Americans, American Indians and Alaska Natives, and white people). About half of all Americans (47%) have at least 1 of 3 key risk factors for heart disease: high blood pressure, high cholesterol, and smoking. Other key indicators include diabetic status, obesity (high BMI), not getting enough physical activity or drinking too much alcohol. Detecting and preventing the factors that have the greatest impact on heart disease is very important in healthcare. Computational developments, in turn, allow the application of machine learning methods to detect "patterns" from the data that can predict a patient's condition.

In [47]:

• The dataset come from the CDC and is a major part of the Behavioral Risk Factor Surveillance System (BRFSS), which conducts annual telephone surveys to gather data on the health status of U.S. residents. As the CDC describes: "Established in 1984 with 15 states, BRFSS now collects data in all 50 states as well as the District of Columbia and three U.S. territories. BRFSS completes more than 400,000 adult interviews each year, making it the largest continuously conducted health survey system in the world.". The most recent dataset (as of February 15, 2022) includes data from 2020. It consists of 401,958 rows and 279 columns. The vast majority of columns are questions asked to respondents about their health status.

Part 2.Data Process and Description

Import dependences

```
# import package
import pandas as pd
import os
import numpy as np
import matplotlib.pyplot as plt
from sklearn.preprocessing import StandardScaler
from scipy. spatial. distance import pdist
from scipy. spatial. distance import squareform
from scipy.cluster.hierarchy import dendrogram, linkage
from scipy.cluster.hierarchy import ward, fcluster, leaders
from scipy. cluster import hierarchy
from scipy.sparse import csr_matrix
import networks as nx
import seaborn as sns
import math
In [20]:
                                                                                                 Ы
%matplotlib inline
In [16]:
                                                                                                  H
import pandas as pd
import matplotlib.pyplot as plt
import seaborn as sns
import numpy as np
from sklearn.linear model import Ridge
import warnings
sns. set style ("white")
%store -r
```

2.1 Pre-processing

H

```
In [6]:

# import source data
heart_df = pd.read_csv('./data/heart_2020_cleaned.csv')

In [22]:

#De duplication, de unmaned, de Nan
#When keep = 'first', keep the sample when the label appears for the first time, and discard all rep
#When inplace = false, return the dataframe after removing duplicates, and the original dataframe wi
# dropna(axis=1, how='all'). Dropna (axis = 0, how ='Any') deletes columns that are all Nan and any
def cleanData(data):
    cleaned_data = data.drop_duplicates(keep='first', inplace=False).loc[:, ~data.columns.str.contai
    return cleaned_data

# clean data
heart_df=cleanData(heart_df)
```

2.2 Preliminary data analysis

```
In [25]:

# Dataset's information
heart_df.info()
```

<class 'pandas.core.frame.DataFrame'>
Int64Index: 301717 entries, 0 to 319794
Data columns (total 18 columns):

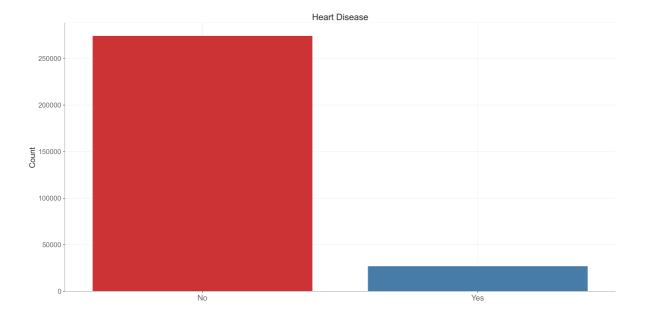
#	Column	Non-Null Count	Dtymo
#	Column	Non-Null Count	Dtype
0	HeartDisease	301717 non-null	object
1	BMI	301717 non-null	float64
2	Smoking	301717 non-null	object
3	AlcoholDrinking	301717 non-null	object
4	Stroke	301717 non-null	object
5	PhysicalHealth	301717 non-null	float64
6	MentalHealth	301717 non-null	float64
7	DiffWalking	301717 non-null	object
8	Sex	301717 non-null	object
9	AgeCategory	301717 non-null	object
10	Race	301717 non-null	object
11	Diabetic	301717 non-null	object
12	PhysicalActivity	301717 non-null	object
13	GenHealth	301717 non-null	object
14	SleepTime	301717 non-null	float64
15	Asthma	301717 non-null	object
16	KidneyDisease	301717 non-null	object
17	SkinCancer	301717 non-null	object
dtypes: float64(4), object(14)			
memory usage: 43.7+ MB			

In [28]:

```
def countp(var):
    # Plot
    fig, ax = plt. subplots (figsize=(20, 10), dpi= 200)
    sns.countplot(data = heart_df, x=var, palette='Set1')
    # Personalization
    plt.title('Heart Disease', fontsize=18)
    plt.xticks(rotation=0, fontsize=16, horizontalalignment='center', alpha=.7)
    plt.yticks(fontsize=14, alpha=.7)
    plt.grid(axis='both', alpha=.3)
    ax.set_xlabel('', fontsize=16)
    ax. set_ylabel('Count', fontsize=16)
    plt.gca().spines["top"].set alpha(0.0)
    plt.gca().spines["bottom"].set_alpha(0.3)
    plt.gca().spines["right"].set_alpha(0.0)
    plt.gca().spines["left"].set_alpha(0.3)
    return plt. show()
```

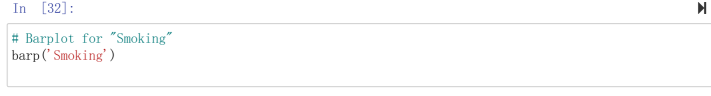
```
[30]:
```

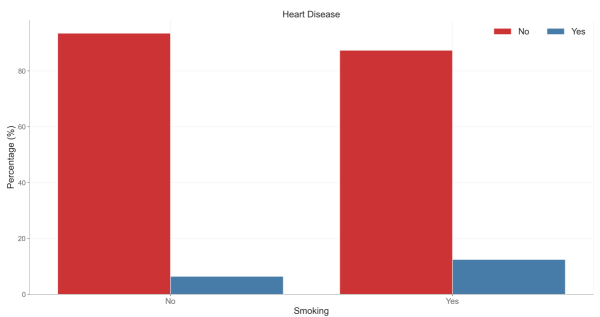
```
# Countplot for the variable "HeartDisease"
countp(var='HeartDisease')
```



The figure above shows a significant imbalance between people with and without heart disease in the data set. Therefore, the EDA of classified data will use standardized values.

```
In [31]:
def barp(var, order=None):
    # Plot
    data normalized = heart df.groupby(f' {var}')['HeartDisease'].value counts(normalize=True).renam
    fig, ax = plt. subplots (figsize=(20, 10), dpi= 200)
    sns.barplot(x=var, y='Percentage', hue='HeartDisease', data=data_normalized, order=order, palet
    # Personalization
    plt. xticks (rotation=0, fontsize=16, horizontalalignment='center', alpha=.7)
    plt.yticks(fontsize=14, alpha=.7)
    plt.grid(axis='both', alpha=.3)
    plt.title('Heart Disease', fontsize=18)
    ax. set_xlabel(f' {var}', fontsize=18)
    ax.set_ylabel('Percentage (%)', fontsize=18)
    ax.legend(frameon=False, ncol=len(heart_df.columns), fontsize='xx-large')
    plt.gca().spines["top"].set_alpha(0.0)
    plt.gca().spines["bottom"].set_alpha(0.3)
    plt.gca().spines["right"].set_alpha(0.0)
    plt.gca().spines["left"].set_alpha(0.3)
    return plt. show()
```



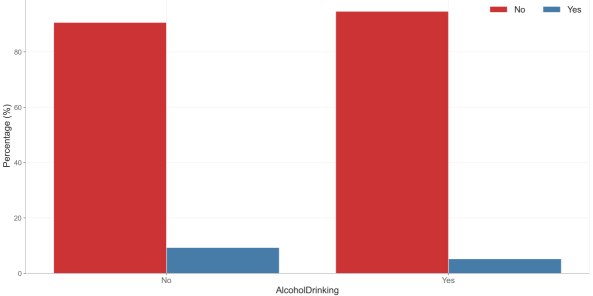


The chart above shows the significant prevalence of heart disease in people with smoking habits.

```
In [39]:
def barp(var, order=None):
    # Plot
    data_normalized = heart_df.groupby(f' {var}')['HeartDisease'].value_counts(normalize=True).renam
    fig, ax = plt. subplots (figsize=(20, 10), dpi= 200)
    sns.barplot(x=var, y='Percentage', hue='HeartDisease', data=data_normalized, order=order, palet
    # Personalization
    plt.xticks(rotation=0, fontsize=16, horizontalalignment='center', alpha=.7)
    plt.yticks(fontsize=14, alpha=.7)
    plt.grid(axis='both', alpha=.3)
    plt.title('Heart Disease', fontsize=18)
    ax. set_xlabel(f' {var}', fontsize=18)
    ax.set_ylabel('Percentage (%)', fontsize=18)
    ax.legend(frameon=False, ncol=len(heart_df.columns), fontsize='xx-large')
    plt.gca().spines["top"].set_alpha(0.0)
    plt.gca().spines["bottom"].set_alpha(0.3)
    plt.gca().spines["right"].set_alpha(0.0)
    plt.gca().spines["left"].set alpha(0.3)
    return plt. show()
```

Barplot for "AlcoholDrinking" barp('AlcoholDrinking')

Heart Disease



The chart shows that the prevalence of heart disease is very high for people who do not drink alcohol.

Description of each variable in the dataset:

Variable	Description	
HeartDisease	Respondents that have ever reported having coronary heart disease (CHD) or myocardial infarction (MI)	
ВМІ	Body Mass Index (BMI)	
Smoking	Have you smoked at least 100 cigarettes in your entire life?	

Variable Description

Heavy drinkers (adult men having more than 14 drinks per week and adult women having more than 7 drinks AlcoholDrinking per week Stroke (Ever told) (you had) a stroke? PhysicalHealth How many days during the past 30 days was your physical health not good? MentalHealth How many days during the past 30 days was your mental health not good? DiffWalking Do you have serious difficulty walking or climbing stairs? Sex Are you male or female? AgeCategory Fourteen-level age category Imputed race/ethnicity value Race Diabetic (Ever told) (you had) diabetes? PhysicalActivity Adults who reported doing physical activity or exercise during the past 30 days other than their regular job GenHealth Would you say that in general your health is... SleepTime On average, how many hours of sleep do you get in a 24-hour period? Asthma (Ever told) (you had) asthma? KidneyDisease Not including kidney stones, bladder infection or incontinence, were you ever told you had kidney disease? (Ever told) (you had) skin cancer? SkinCancer

2.3 Variable correlation analysis

return working df, converted columns

```
In [44]:

def data_convert(df):
    if type(df)!=pd.core.frame.DataFrame:
        raise ValueError('input is not a pandas dataframe')
    working_df = df.copy()
    cols = working_df.columns
```

```
working_dr dr.copy()
cols = working_df.columns
converted_columns = {}
for col in cols:
    if working_df[col].dtype == '0':
        unique_values = working_df[col].unique()
        converted_values = {v:k for k, v in enumerate(unique_values)}
        for value in unique_values:
            working_df[col] = working_df[col].replace(value, converted_values[value])
        converted_columns[col] = converted_values
```

```
In [45]:
```

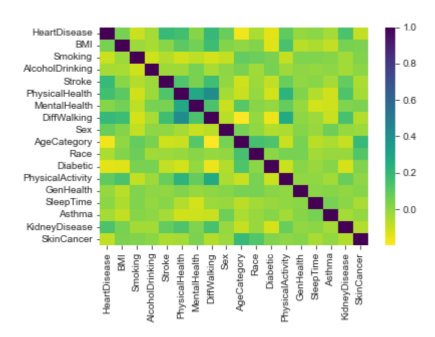
```
cleaned_df, conversion_index = data_convert(heart_df)
```

In [46]: ▶

```
sns.heatmap(cleaned_df.corr(), cmap='viridis_r')
```

Out[46]:

<AxesSubplot:>



Using the thermodynamic diagram, we can intuitively see the correlation between various variables. It can be seen that diffwalking is highly correlated with physical health. The correlation between heartdisease and agecategory is small.

Part 3. Model establishment and algorithm analysis

