

Healthcare Cost and Utilization Project Code Preparation

Drexel University LeBow College of Business
Fall Term 2023

Type 1 Diabetes Table of Contents

Merging Data	2
Diabetes K-Modes Clustering	2
Diabetes Chi-Square Hypothesis Testing.....	3
Diabetes Codes: Jupyter Notebook.....	4

Asthma Table of Contents

Asthma Initial Setup.....	11
Asthma Statistical Testing Using Chi-Squared Test	11
Asthma K-Means Clustering.....	11
Asthma K-Modes Clustering.....	11
Asthma Codes: SQL and Jupyter Notebook.....	11

Merging Data

Dataset Importation:

- Import 'dask.dataframe' library for parallel computing with larger datasets.
- Using Dask, three datasets — **Hospital**, **Core**, and **Severity** — are read into memory.

Merging Hospital with Severity Dataset:

- The Hospital and Severity datasets are merged on the 'HOSP_KID' column. It is a 'left join', which means all entries from the Hospital dataset and corresponding entries from the Severity dataset will be included.
- The resultant merged dataset is saved as a **Hospital_Severity.csv** file in the specified output directory (*Dataset/Data Cleaning/Merged*).

Merging Hospital with Core Dataset:

- Similarly, the Hospital and Core datasets are merged on the 'HOSP_KID' column using a 'left join'.
- This merged dataset is also saved as a **Hospital_Core.csv** file in the designated output location (*Dataset/Data Cleaning/Merged*).

Diabetes K-Modes Clustering

Dataset Importation:

- Import libraries such as Pandas, Dask, and Numpy and read the hospital datasets using Dask for efficient memory.
- Apply age filters to isolate patients between 0 to 18 years and filter diagnosis codes for type 1 diabetes.

Data Transformation:

- Categorize patients into age groups ('Infants', 'Children', and 'Adolescents'). Extract relevant diagnosis codes and convert Dask dataframes to Pandas dataframes for further processing.

Elbow Method:

- Use the Elbow method to determine the optimal number of clusters by plotting the cost function against the number of clusters.

KModes Algorithm:

- Apply the KModes clustering algorithm using the optimal number of clusters. Ensure that the algorithm is initialized with a consistent random seed for reproducibility.

Export Predicted Cluster Dataset:

- Export the predicted clustering dataframe with assigned clusters to a **HCUP_Clustering.csv** file for further cluster data analysis.

Diabetes Chi-Square Hypothesis Testing

Dataset Importation:

- Import libraries such as Pandas, and read the cluster datasets **HCUP_Clustering.csv** to dataframe. It includes relevant columns for the analysis, including 'GENDER', 'ALL_PATIENT_SEVERITY', 'TYPE1_DIABETES_CODES', 'HOSPITAL_REGION', 'RACE_DESC', 'LOS', and 'AGE_GROUP'.
- To conducting hypothesis tests, first to import scipy.stats library from stats.

Gender and Severity of Diagnosis:

- Create a contingency table for 'GENDER' and 'ALL_PATIENT_SEVERITY' whether the test is between two different categorical variables are associate or not.
- Perform a chi-square test and interpret the results.

Gender and Type 1 Diabetes Codes:

- Create a contingency table for 'GENDER' and 'TYPE1_DIABETES_CODES'.
- Perform a chi-square test and interpret the results.

Hospital Region vs. Ethnicity:

- Create a contingency table for 'HOSPITAL_REGION' and 'RACE_DESC'.
- Perform a chi-square test and interpret the results.

Ethnicity vs. Type 1 Diabetes Codes:

- Create a contingency table for 'RACE_DESC' and 'TYPE1_DIABETES_CODES'.
- Perform a chi-square test and interpret the results.

Length of Stay vs. Age Group:

- Create a contingency table for 'LOS' and 'AGE_GROUP'.
- Perform a chi-square test and interpret the results.

Length of Stay vs. Severity:

- Create a contingency table for 'LOS' and 'ALL_PATIENT_SEVERITY'.
- Perform a chi-square test and interpret the results.

Age Group vs. Severity:

- Create a contingency table for 'AGE_GROUP' and 'ALL_PATIENT_SEVERITY'.
- Perform a chi-square test and interpret the results.

Interpreting the Results:

- These hypothesis tests will output the Chi-square statistic, p-value, degrees of freedom, and a conclusion to accept or reject the null hypothesis based on 5% significance level

Diabetes Codes: Jupyter Notebook

■ Merging data source

```
In [1]: 1 import pandas as pd
2 import dask.dataframe as dd
3 import os
4
5 # File paths
6 hospital_file = "C:/Users/Lenovo/OneDrive - Drexel University/Fall 2023/BSAN-710/Capstone Project/Production (Codes and EDA)
7 core_file = "C:/Users/Lenovo/OneDrive - Drexel University/Fall 2023/BSAN-710/Capstone Project/Production (Codes and EDA)/Dat
8 severity_file = "C:/Users/Lenovo/OneDrive - Drexel University/Fall 2023/BSAN-710/Capstone Project/Production (Codes and EDA)
9 merged_severity_output_file = "C:/Users/Lenovo/OneDrive - Drexel University/Fall 2023/BSAN-710/Capstone Project/Production (
10 merged_core_output_file = "C:/Users/Lenovo/OneDrive - Drexel University/Fall 2023/BSAN-710/Capstone Project/Production (Code
11
```

Define dictionary

```
In [2]: 1 # Define the dtype dictionary
2 dtype_dict = {
3     'AGE_NEONATE': 'str', 'AMONTH': 'str', 'ELECTIVE': 'str', 'LOS': 'str', 'PAY1': 'str', 'PRDAY10': 'str',
4     'PRDAY11': 'str', 'PRDAY8': 'str', 'PRDAY9': 'str', 'RACE': 'str', 'TOTCHG': 'str', 'ZIPINC_QRTL': 'str',
5     'DQTR': 'str', 'PRDAY12': 'str', 'PRDAY13': 'str', 'PRDAY14': 'str', 'PRDAY15': 'str', 'PRDAY16': 'str', 'PRDAY17': 'str',
6     'PRDAY18': 'str', 'PRDAY19': 'str', 'PRDAY20': 'str', 'PRDAY21': 'str', 'PRDAY22': 'str', 'PRDAY23': 'str',
7     'PRDAY24': 'str', 'PRDAY25': 'str', 'PRDAY2': 'str', 'PRDAY3': 'str', 'PRDAY4': 'str', 'PRDAY5': 'str',
8     'PRDAY6': 'str', 'TRAN_IN': 'str', 'TRAN_OUT': 'str', 'FEMALE': 'str', 'DISPUNIFORM': 'str', 'DIED': 'str'
9 }
```

Read data

```
In [3]: 1 # Read datasets
2 hospital_df = dd.read_csv(hospital_file)
3 core_df = dd.read_csv(core_file, dtype=dtype_dict)
4 severity_df = dd.read_csv(severity_file)
```

Merging Hospital and Core

```
In [4]: 1 merged_core_df = hospital_df.merge(core_df, on='HOSP_KID', how='left')
2
3 # Check if the file exists
4 if not os.path.exists(merged_core_output_file):
5     # If the file does not exist, export the final merged dataframe to a CSV file
6     merged_core_df.compute().to_csv(merged_core_output_file, index=False)
7
```

Merging Hospital and Severity

```
In [5]: 1 # Merge the dataframes Hospital and Severity
2 merged_severity_df = hospital_df.merge(severity_df, on='HOSP_KID', how='left')
3
4 # Check if the file exists
5 if not os.path.exists(merged_severity_output_file):
6     # If the file does not exist, export the final merged dataframe to a CSV file
7     merged_severity_df.compute().to_csv(merged_severity_output_file, index=False)
8
```

■ Clustering with KModes

```
In [ ]: 1 import pandas as pd
2 import dask.dataframe as dd
3 import numpy as np
4 import matplotlib.pyplot as plt
5 import seaborn as sns
```

Load csv file

```
In [ ]: 1 hospital_core_file = "C:/Users/Lenovo/OneDrive - Drexel University/Fall 2023/BSAN-710/Capstone Project/Production (Codes and
2 hospital_severity_file = "C:/Users/Lenovo/OneDrive - Drexel University/Fall 2023/BSAN-710/Capstone Project/Production (Codes
```

Define dictionary

```
In [ ]: 1 # Define the dtype dictionary
2 dtype_dict = {
3     'AGE_NEONATE': 'str', 'AMONTH': 'str', 'ELECTIVE': 'str', 'LOS': 'str', 'PAY1': 'str', 'PRDAY10': 'str',
4     'PRDAY11': 'str', 'PRDAY8': 'str', 'PRDAY9': 'str', 'RACE': 'str', 'TOTCHG': 'str', 'ZIPINC_QRTL': 'str',
5     'DQTR': 'str', 'PRDAY12': 'str', 'PRDAY13': 'str', 'PRDAY14': 'str', 'PRDAY15': 'str', 'PRDAY16': 'str', 'PRDAY17': 'str',
6     'PRDAY18': 'str', 'PRDAY19': 'str', 'PRDAY20': 'str', 'PRDAY21': 'str', 'PRDAY22': 'str', 'PRDAY23': 'str',
7     'PRDAY24': 'str', 'PRDAY25': 'str', 'PRDAY2': 'str', 'PRDAY3': 'str', 'PRDAY4': 'str', 'PRDAY5': 'str',
8     'PRDAY6': 'str', 'TRAN_IN': 'str', 'TRAN_OUT': 'str', 'FEMALE': 'object', 'DISPUNIFORM': 'str', 'DIED': 'str'
9 }
10
```

Read dataset and filter only type 1 diabetes codes tranpose into one columns

```
In [ ]: 1 # Read hospital_core dataset
2 hospital_core_df = dd.read_csv(hospital_core_file, dtype=dtype_dict)
3 # Read severity dataset
4 hospital_severity_df = dd.read_csv(hospital_severity_file, dtype=dtype_dict)
5
6 # Filter for age
7 age_condition = (hospital_core_df['AGE'] >= 0) & (hospital_core_df['AGE'] <= 18)
8
9 # Pre-existing type 1 diabetes diagnosis codes
10 type1_preexist = ['024011', '024012', '023013', '024019', '02402', '02403']
11
12 # Vectorized condition for both type1_preexist and 'E10' for columns I10_DX1 to I10_DX5
13 dx_conditions = False
14 for column in ['I10_DX1', 'I10_DX2', 'I10_DX3', 'I10_DX4', 'I10_DX5']:
15     dx_conditions |= hospital_core_df[column].isin(type1_preexist).fillna(False)
16     dx_conditions |= hospital_core_df[column].str.startswith('E10').fillna(False)
17
18 # Combine age and diabetes diagnosis conditions
19 combined_condition = age_condition & dx_conditions
20
21 hospital_core_df = hospital_core_df[combined_condition]
22
23 # Extract all diagnosis codes related to type 1 diabetes
24 def extract_diagnosis_codes(row):
25     return [code for code in row if pd.notnull(code) and (code.startswith('E10') or code in type1_preexist)]
26
27 hospital_core_df['TYPE1_DIABETES_CODES'] = hospital_core_df[['I10_DX1', 'I10_DX2', 'I10_DX3', 'I10_DX4', 'I10_DX5']].apply(extract_diagnosis_codes, axis=1)
28
29 # Transform dask dataframe to pandas dataframe
30 hospital_core_df = hospital_core_df.compute()
31
32 # Reset the index for a unique index
33 hospital_core_df.reset_index(drop=True, inplace=True)
34
35 #hospital_core_df.head()
36
```

```
In [ ]: 1 hospital_core_df.shape[0]
```

```
In [ ]: 1 # Join Severity to Hospital_Core by 'RECNUM'
2 hospital_severity_df = hospital_severity_df.compute()
3
4 hospital_core_severity_df = hospital_core_df.merge(hospital_severity_df, on='RECNUM', how='inner')
5 hospital_core_severity_df.head()
6
```

```
In [ ]: 1 hospital_core_severity_df.info()
```

```
In [ ]: 1 # Join Severity to Hospital_Core by 'RECNUM'
2 hospital_severity_df = hospital_severity_df.compute()
3
4 hospital_core_severity_df = hospital_core_df.merge(hospital_severity_df, on='RECNUM', how='inner')
5 hospital_core_severity_df.head()
6
```

```
In [ ]: 1 hospital_core_severity_df.info()
```

```
In [ ]: 1 # Checking Null values
2 hospital_core_severity_df.isnull().sum()*100/hospital_core_severity_df.shape[0]
```

```
In [ ]: 1 def categorize_age(age):
2     if age < 1:
3         return "Infant"
4     elif 1 <= age <= 13:
5         return "Children"
6     else:
7         return "Adolescents"
8
9 # Categorization function 'AGE_GROUP'
10 hospital_core_severity_df['AGE_GROUP'] = hospital_core_severity_df['AGE'].apply(categorize_age)
11 hospital_core_severity_df['TYPE1_DIABETES_CODES'] = hospital_core_severity_df['TYPE1_DIABETES_CODES'].apply(lambda x: x[0] if x else None)
12
13
14 hospital_core_severity_df = hospital_core_severity_df.drop('AGE', axis = 1)
```

```
In [ ]: 1 select_diagnosis_columns = ['AGE_GROUP', 'GENDER', 'HOSPITAL_REGION', 'LOS', 'ALL_PATIENT_SEVERITY', 'RACE_DESC', 'TYPE1_DIAB']
2
3 hospital_core_severity_df = hospital_core_severity_df[select_diagnosis_columns]
4 hospital_core_severity_df.head()
```

```
In [ ]: 1 print("Total number of records for clustering: " + str(hospital_core_severity_df.shape))
```

Copy Data

```
In [ ]: 1 # Keep a copy of data
        2 hospital_core_severity_df_copy = hospital_core_severity_df.copy()

In [ ]: 1 hospital_core_severity_df.head()
```

Find Optimal k by Elbow Method

```
In [ ]: 1 from kmodes.kmodes import KModes
        2 import matplotlib.pyplot as plt
        3
        4 np.random.seed(42)
        5 cost = []
        6 for num_clusters in list(range(1,8)):
        7     kmode = KModes(n_clusters=num_clusters, init = "Cao", n_init = 5, verbose=0)
        8     kmode.fit_predict(hospital_core_severity_df)
        9     cost.append(kmode.cost_)
       10
       11 y = np.array([i for i in range(1,8,1)])
       12
       13 plt.figure(figsize=(10,6))
       14 #plt.plot(y, cost)
       15 plt.plot(y, cost, marker='o')
       16 plt.xlabel('Number of Clusters')
       17 plt.ylabel('Cost')
       18 plt.title('Elbow Method For Optimal k')
       19 plt.grid(False)
       20 plt.show()
       21
       22
```

Model fitting

```
In [ ]: 1 optimal_k = 3 # as per elbow method

In [ ]: 1 # fit the KModes Clustering algorithm with 5 optimal clusters.
        2 np.random.seed(42)
        3 kmd = KModes(n_clusters=optimal_k, init = "Huang", n_init = 5, verbose=0)
        4 kmd_clusters = kmd.fit_predict(hospital_core_severity_df)

In [ ]: 1 # Combining predicted clusters with the original dataframe
        2 hospital_core_severity_df = hospital_core_severity_df_copy.reset_index()

In [ ]: 1 kmd_clusters_df = pd.DataFrame(kmd_clusters)
        2 kmd_clusters_df.columns = ['CLUSTERS']
        3 kmd_clusters_df = pd.concat([hospital_core_severity_df, kmd_clusters_df], axis = 1).reset_index()
        4 kmd_clusters_df = kmd_clusters_df.drop(['index', 'level_0'], axis = 1)
        5 # added row numbers
        6 kmd_clusters_df['ROWNUM'] = range(1, len(kmd_clusters_df) + 1)
        7
        8 # Create a mapping from old to new cluster labels
        9 label_mapping = {0: 1, 1: 2, 2: 3}
       10
       11 # start with cluster 1 instead of 0. map the old cluster labels to the new ones
       12 kmd_clusters_df['CLUSTERS'] = kmd_clusters_df['CLUSTERS'].map(label_mapping)
```

Extract the clusters dataset

```
In [ ]: 1 import os
        2 output_file = 'C:/Users/Lenovo/OneDrive - Drexel University/Fall 2023/BSAN-710/Capstone Project/Production (Codes and EDA)/D
        3
        4 # Check if the file exists
        5 if not os.path.exists(output_file):
        6     # If the file does not exist, create HCUP_Clustering.csv file
        7     kmd_clusters_df.to_csv(output_file, index=False)
        8
        9 kmd_clusters_df.head()
```

Descriptive Analysis

Cluster Proportion

```
In [ ]: 1 # Calculate the proportions
2 np.random.seed(42)
3 cluster_size = kmd_clusters_df['CLUSTERS'].value_counts().sort_index()
4 cluster_size_proportion = (cluster_size / cluster_size.sum()) * 100
5
6
7 # Plotting
8 plt.figure(figsize=(10, 6))
9 cluster_size_proportion.plot(kind='bar')
10
11 # Adding titles and labels
12 plt.title('Cluster Size Proportions (%)')
13 plt.xlabel('Cluster')
14 plt.ylabel('Proportion (%)')
15 plt.xticks(rotation=0)
16
17 # show percentages above the bars
18 for i, value in enumerate(cluster_size_proportion):
19     plt.text(i, value + 0.5, f'{value:.2f}%', ha='center', va='bottom')
20
21 plt.tight_layout()
22 plt.show()
23
In [ ]: 1 kmd_clusters_df.head()
In [ ]: 1 # Heatmap of AGE_GROUP
2 crosstab = pd.crosstab(kmd_clusters_df['AGE_GROUP'], kmd_clusters_df['CLUSTERS'], normalize='index') * 100
3
4 # matrix with percentages
5 annot_array = crosstab.values
6 annotations = [[f'{val:.2f}%' for val in row] for row in annot_array]
7
8 plt.figure(figsize=(10, 6))
9 sns.heatmap(crosstab, annot=annotations, cmap="YlGnBu", fmt="")
10 plt.title("Distribution of Age Group across Clusters")
11 plt.show()
12
13 # Heatmap of GENDER
14 crosstab = pd.crosstab(kmd_clusters_df['GENDER'], kmd_clusters_df['CLUSTERS'], normalize='index') * 100
15
16 # matrix with percentages
17 annot_array = crosstab.values
18 annotations = [[f'{val:.2f}%' for val in row] for row in annot_array]
19
20 plt.figure(figsize=(10, 6))
21 sns.heatmap(crosstab, annot=annotations, cmap="YlGnBu", fmt="")
22 plt.title("Distribution of Gender across Clusters")
23 plt.show()
24
25 # Heatmap of RACE_DESC
26 crosstab = pd.crosstab(kmd_clusters_df['RACE_DESC'], kmd_clusters_df['CLUSTERS'], normalize='index') * 100
27
28 # matrix with percentages
29 annot_array = crosstab.values
30 annotations = [[f'{val:.2f}%' for val in row] for row in annot_array]
31
32 plt.figure(figsize=(10, 6))
33 sns.heatmap(crosstab, annot=annotations, cmap="YlGnBu", fmt="")
34 plt.title("Distribution of Race across Clusters")
35 plt.show()
36
37 # Heatmap of RACE_DESC
38 crosstab = pd.crosstab(kmd_clusters_df['HOSPITAL_REGION'], kmd_clusters_df['CLUSTERS'], normalize='index') * 100
39
40 # matrix with percentages
41 annot_array = crosstab.values
42 annotations = [[f'{val:.2f}%' for val in row] for row in annot_array]
43
44 plt.figure(figsize=(10, 6))
45 sns.heatmap(crosstab, annot=annotations, cmap="YlGnBu", fmt="")
46 plt.title("Distribution of Hospital Region across Clusters")
47 plt.show()
48
49 # Heatmap of Severity
50 crosstab = pd.crosstab(kmd_clusters_df['ALL_PATIENT_SEVERITY'], kmd_clusters_df['CLUSTERS'], normalize='index') * 100
51
52 # matrix with percentages
53 annot_array = crosstab.values
54 annotations = [[f'{val:.2f}%' for val in row] for row in annot_array]
55
56 plt.figure(figsize=(10, 6))
57 sns.heatmap(crosstab, annot=annotations, cmap="YlGnBu", fmt="")
58 plt.title("Distribution of Health Severity across Clusters")
59 plt.show()
60
61 # Heatmap of LOS
62 kmd_clusters_df['LOS'] = kmd_clusters_df['LOS'].astype(int)
63
```

```

64 def categorize_los(los):
65     if los <= 5:
66         return "Short-term"
67     elif 5 < los <= 15:
68         return "Mid-term"
69     else:
70         return "Long-term"
71
72 kmd_clusters_df['LOS_Category'] = kmd_clusters_df['LOS'].apply(categorize_los)
73
74 # crosstab with the LOS_category and Clusters
75 crosstab = pd.crosstab(kmd_clusters_df['LOS_Category'], kmd_clusters_df['CLUSTERS'])
76 # Convert the crosstab values to proportions
77 crosstab_percentage = (crosstab / crosstab.sum().sum()) * 100
78
79 # Generate annotations with percentage symbol
80 annotations = crosstab_percentage.applymap(lambda x: f"{x:.2f}%")
81
82 plt.figure(figsize=(10, 6))
83 sns.heatmap(crosstab_percentage, annot=annotations, cmap="YlGnBu", fmt="s")
84 plt.title("Distribution of LOS across Clusters in Percentages")
85 plt.xlabel("Cluster")
86 plt.ylabel("LOS Category")
87 plt.show()
88
89
90 # Heatmap of TYPE1 DIABETES
91 crosstab = pd.crosstab(kmd_clusters_df['TYPE1_DIABETES_CODES'], kmd_clusters_df['CLUSTERS'], normalize='index') * 100
92
93 # matrix with percentages
94 annot_array = crosstab.values
95 annotations = [[f'{val:.2f}%' for val in row] for row in annot_array]
96
97 plt.figure(figsize=(10, 6))
98 sns.heatmap(crosstab, annot=annotations, cmap="YlGnBu", fmt="")
99 plt.title("Distribution of Type1 Diabetes Diagnosis across Clusters")
100 plt.show()
101
102
103
104

```

■ Chi-Square Testing

```

In [ ]: 1 import pandas as pd

In [ ]: 1 hospital_cluster_data = pd.read_csv("C:/Users/Lenovo/OneDrive - Drexel University/Fall 2023/BSAN-710/Capstone Project/Produ
2 hospital_cluster_data.head()

```

Hypothesis 1: A disparity in gender concerning the severity of the diagnosis of type 1 diabetes.

create the contingency tables for each relationship, performed the chi-square test to determine the significance of associations, and provided the interpretations based on the p-values

```

In [ ]: 1 import scipy.stats as stats
2
3 # Create a contingency table for GENDER vs ALL_PATIENT_SEVERITY
4 contingency_table_gender_sev = pd.crosstab(hospital_cluster_data['GENDER'], hospital_cluster_data['ALL_PATIENT_SEVERITY'])
5 print("Contingency Table for Gender and Severity:")
6 print(contingency_table_gender_sev)
7
8 # Conduct the chi-square test
9 chi2, p, df_1, _ = stats.chi2_contingency(contingency_table_gender_sev)
10 print(f"\nHypothesis 1 chi-square: {chi2}, p-value: {format(p, '.10f')}, degrees of freedom: {df_1}\n")
11 print("-" * 50)
12
13 # Interpret the result gender vs severity
14 alpha = 0.05
15 if p < alpha:
16     print("Reject the null hypothesis: There is a significant relationship between gender and severity.")
17 else:
18     print("Fail to reject the null hypothesis: There is no significant relationship between gender and severity.")
19
20 #-----
21
22 # Create a contingency table for GENDER vs TYPE1_DIABETES_CODES
23 contingency_table_gender_dia = pd.crosstab(hospital_cluster_data['GENDER'], hospital_cluster_data['TYPE1_DIABETES_CODES'])
24 print("\nContingency Table for Gender and Type 1 Diabetes Codes:")
25 print(contingency_table_gender_dia)
26
27 # Conduct the chi-square test
28 chi2_gender, p_gender, df_2, _ = stats.chi2_contingency(contingency_table_gender_dia)
29 print(f"\nchi-square for Gender vs. Type 1 Diabetes Codes: {chi2_gender}, p-value: {format(p_gender, '.10f')}, degrees of freedom: {df_2}\n")
30 print("-" * 50)
31
32 # Interpret the result for gender vs type 1 diabetes
33 alpha = 0.05
34 if p_gender < alpha:
35     print("Reject the null hypothesis: There is a significant relationship between gender and Type 1 diabetes codes.")
36 else:
37     print("Fail to reject the null hypothesis: There is no significant relationship between gender and Type 1 diabetes codes.")
38 print("-" * 50)
39
40 #-----
41

```



```
In [ ]: 1 print("\nConclusion:")
2
3 print("These findings support Hypothesis 1, gender and severity have correlations with Type 1 diabetes.")
```

Hypothesis 2: Factors such as the regions of hospitals and ethnicity are found to have a correlation with Type 1 diabetes within the group.

```
In [ ]: 1 # Create a contingency table for HOSPITAL_REGION vs Ethnicity
2 contingency_table_hosp_race = pd.crosstab(hospital_cluster_data['HOSPITAL_REGION'], hospital_cluster_data['RACE_DESC'])
3 print("\nContingency Table for Hospital Region vs. Ethnicity:")
4 print(contingency_table_hosp_race)
5
6 # Conduct the chi-square test
7 chi2, p, df_3, _ = stats.chi2_contingency(contingency_table_hosp_race)
8 print(f"\nChi-square for Hospital Region vs. Ethnicity: {chi2}, p-value: {format(p, '.10f')}, degrees of freedom: {df_3}\n")
9 print("-" * 50)
10
11 # Interpret the result for hospital region
12 alpha = 0.05
13 if p < alpha:
14     print("Reject the null hypothesis: There is a significant relationship between hospital region and ethnicity.")
15 else:
16     print("Fail to reject the null hypothesis: There is no significant relationship between hospital region and Ethnicity.")
17
18 print("-" * 50)
19
20 #-----
21 # Create a contingency table for RACE_DESC vs TYPE1_DIABETES_CODES
22 contingency_table_race_dia = pd.crosstab(hospital_cluster_data['RACE_DESC'], hospital_cluster_data['TYPE1_DIABETES_CODES'])
23 print("\nContingency Table for Ethnicity vs. Type 1 Diabetes Codes:")
24 print(contingency_table_race_dia)
25
26 # Conduct the chi-square test
27 chi2_race, p_race, df_4, _ = stats.chi2_contingency(contingency_table_race_dia)
28 print(f"\nChi-square for Ethnicity vs. Type 1 Diabetes Codes: {chi2_race}, p-value: {format(p_race, '.10f')}, degrees of free
29 print("-" * 50)
30
31 # Interpret the result for ethnicity
32 if p_race < alpha:
33     print("Reject the null hypothesis: There is a significant relationship between ethnicity and Type 1 diabetes codes.")
34 else:
35     print("Fail to reject the null hypothesis: There is no significant relationship between ethnicity and Type 1 diabetes co
36
```

```
In [ ]: 1 print("\nConclusion:")
2
3 print("These findings support Hypothesis 2, hospital regions and ethnicity have correlations with Type 1 diabetes within the
```

Hypothesis 3: Distribution of the length of a patient's stay in the hospital and severity of diabetes diagnoses among different age groups.

```
In [ ]: 1 # Define the bins and their Labels
2 bins = [-1, 0, 7, 14, 30, float('inf')] # the -1 ensures that 0 is included in the first bin
3 labels = ['0 days', '1-7 days', '8-14 days', '15-30 days', '31+ days']
4
5
6 # Create a new column for the binned LOS
7 hospital_cluster_data['LOS_Binned'] = pd.cut(hospital_cluster_data['LOS'], bins=bins, labels=labels)
8
9 # Create a contingency table for the binned LOS vs AGE_GROUP
10 contingency_table_los_age = pd.crosstab(hospital_cluster_data['LOS_Binned'], hospital_cluster_data['AGE_GROUP'])
11 print("\nContingency Table for Length of Stay vs. Age Group:")
12 print(contingency_table_los_age)
13
14 # Conduct the chi-square test
15 chi2, p, df_5, _ = stats.chi2_contingency(contingency_table_los_age)
16 print(f"\nChi-square for Length of stay vs. Age Group: {chi2}, p-value: {format(p, '.10f')}, degrees of freedom: {df_5}\n")
17 print("-" * 50)
18
19 # Interpret the result for LOS
20 alpha = 0.05
21 if p < alpha:
22     print("Reject the null hypothesis: There is a significant relationship between length of stay and patient age group.")
23 else:
24     print("Fail to reject the null hypothesis: There is no significant relationship between length of stay and patient age group.")
25
26 print("-" * 50)
27
28 #-----
29
30 # Create a contingency table for Los vs severity
31 contingency_table_los_sev = pd.crosstab(hospital_cluster_data['LOS_Binned'], hospital_cluster_data['ALL_PATIENT_SEVERITY'])
32 print("\nContingency Table for Length of stay vs. Severity:")
33 print(contingency_table_los_sev)
34
35 # Conduct the chi-square test
36 chi2_los, p_los, df_6, _ = stats.chi2_contingency(contingency_table_los_sev)
37 print(f"\nChi-square for Length of stay vs. Severity: {chi2_los}, p-value: {format(p_los, '.10f')}, degrees of freedom: {df_6}\n")
38 print("-" * 50)
39
40 # Interpret the result for Los and severity
41 if p_los < alpha:
42     print("Reject the null hypothesis: There is a significant relationship between length of stay and severity.")
43 else:
44     print("Fail to reject the null hypothesis: There is no significant relationship between length of stay and severity.")
45
46 #-----
47
48 # Create a contingency table for Age group vs severity
49 contingency_table_age_sev = pd.crosstab(hospital_cluster_data['AGE_GROUP'], hospital_cluster_data['ALL_PATIENT_SEVERITY'])
50 print("\nContingency Table for Age Group vs. Severity:")
51 print(contingency_table_age_sev)
52
53 # Conduct the chi-square test
54 chi2_ag, p_ag, df_7, _ = stats.chi2_contingency(contingency_table_age_sev)
55 print(f"\nChi-square for Length of stay vs. Severity: {chi2_ag}, p-value: {format(p_ag, '.10f')}, degrees of freedom: {df_7}\n")
56 print("-" * 50)
57
58 # Interpret the result for age group and severity
59 if p_ag < alpha:
60     print("Reject the null hypothesis: There is a significant relationship between age group and severity.")
61 else:
62     print("Fail to reject the null hypothesis: There is no significant relationship between age group and severity.")
63
```

Asthma Initial Setup

Similar to the merging done for Diabetes, the datasets were combined together. Afterward, a basic summary of the dataset was created to determine the spread of the data. In addition, data cleaning and preprocessing were done prior to statistical testing and clustering analysis.

Asthma Statistical Testing Using Chi-Squared Test

Similar to the statistical testing done for Diabetes, chi-squared test for statistical analysis, examining the relationships between categorical variables. The test generated a p-value and a Chi-Square value. A low p-value (<0.05) suggested significant associations, enabling us to make informed decisions about the validity of patterns within our dataset and test our initial hypothesis.

Asthma K-Means Clustering

Applying K-Means clustering, we segmented the data into distinct groups based on similarities. This unsupervised machine learning technique allowed us to identify inherent patterns and groupings within the dataset.

For better results we created bins for variables like Age and Total Charges. The clustering algorithm divided the entire data for asthmatic patients into 5 clusters.

Asthma K-Modes Clustering

We also employed K-Modes clustering, specifically designed for categorical data. This approach further enriched our understanding by revealing patterns in non-numeric variables, providing a holistic view of the dataset.

The clustering algorithm divided the entire data for asthmatic patients into 3 clusters.

Asthma Codes: SQL and Jupyter Notebook

SQL Queries:

--Query to create separate table for respiratory diseases based on diagnosis code:
CREATE TABLE capstone2.RESPIRATORY_DISEASES AS

```
SELECT HOSP_KID  
      ,RECNUM  
      ,DXCCSR_RSP001  
      ,DXCCSR_RSP002  
      ,DXCCSR_RSP003
```

```

,DXCCSR_RSP004
,DXCCSR_RSP005
,DXCCSR_RSP006
,DXCCSR_RSP007
,DXCCSR_RSP008
,DXCCSR_RSP009
,DXCCSR_RSP010
,DXCCSR_RSP011
,DXCCSR_RSP012
,DXCCSR_RSP013
,DXCCSR_RSP014
,DXCCSR_RSP015
,DXCCSR_RSP016
,DXCCSR_RSP017
FROM `capstone - 400517. capstone2.kid_GPRS`

```

--QUERY FOR asthma cases by Gender

```

SELECT FEMALE
      ,count(*)
FROM `capstone - 400517. capstone2.table_core`
WHERE Asthma <> '0'
      AND Asthma IS NOT NULL
GROUP BY 1

```

--QUERY FOR asthma cases by INCOME

```

SELECT ZIPINC_QRTL
      ,count(*)
FROM `capstone - 400517. capstone2.table_core`
WHERE Asthma <> '0'
GROUP BY 1

```

--QUERY FOR asthma cases by race

```

SELECT RACE
      ,count(*)
FROM `capstone - 400517. capstone2.table_core`
--WHERE Asthma <> '0'
GROUP BY 1

```

--query for asthma and infectious diseases

```

SELECT Asthma
      ,INF_All
      ,COUNT(*) AS INFANDASTHMA
FROM `capstone - 400517. capstone2.table_core`
WHERE Asthma <> '0'
      AND INF_All <> 0
GROUP BY 1
      ,2

```

--COMORBIDITY ANALYSIS FOR ASTHMA AND OTHER DISEASES

```

SELECT RECNUM
FROM `capstone - 400517. capstone2.table_core`
WHERE (
      Blood_All <> 0
      OR MAL_All <> 0
      OR MBD_ALL <> 0
      OR END_ALL <> 0
      OR INF_All <> 0

```

```

        OR DIG_All <> 0
        OR NVS_All <> 0
        OR PRG_All <> 0
        OR CIR_All <> 0
        OR SKN_All <> 0
        OR MUS_All <> 0
        OR Tumor_All <> 0
        OR EYE_All <> 0
        OR EAR_All <> 0
    )
    AND Asthma <> '0'

```

--Data cleaning in gender

```

DELETE
FROM `capstone - 400517. capstone2.table_core`
WHERE FEMALE IN (
    'A'
    , 'C'
    , 'nan'
);

```

--query to get total charges by payer

```

SELECT
    --RECNUM
    Pay1
    ,sum(TotalChargesInteger)
FROM `capstone - 400517. capstone2.table_core`
WHERE Asthma <> '0'
    AND (
        Blood_All = 0
        AND MAL_All = 0
        AND MBD_All = 0
        AND END_All = 0
        AND INF_All = 0
        AND DIG_All = 0
        AND NVS_All = 0
        AND PRG_All = 0
        AND CIR_All = 0
        AND SKN_All = 0
        AND MUS_All = 0
        AND Tumor_All = 0
        AND EYE_All = 0
        AND EAR_All = 0
    )
    AND PAY1 IN (
        '1'
        , '2'
    )
--and race <> 'nan'
--and Race not like '%nan'
GROUP BY 1

```

--Query to check count of each endocrine disease

```

SELECT SUM(CASE
    WHEN CAST(DXCCSR_END001 AS INT64) = 1

```

```

        THEN 1
    ELSE 0
END) AS Count_1
,SUM(CASE
    WHEN CAST(DXCCSR_END002 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_2
,SUM(CASE
    WHEN CAST(DXCCSR_END003 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_3
,SUM(CASE
    WHEN CAST(DXCCSR_END004 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_4
,SUM(CASE
    WHEN CAST(DXCCSR_END006 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_6
,SUM(CASE
    WHEN CAST(DXCCSR_END007 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_7
,SUM(CASE
    WHEN CAST(DXCCSR_END008 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_8
,SUM(CASE
    WHEN CAST(DXCCSR_END009 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_9
,SUM(CASE
    WHEN CAST(DXCCSR_END010 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_10
,SUM(CASE
    WHEN CAST(DXCCSR_END011 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_11
,SUM(CASE
    WHEN CAST(DXCCSR_END012 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_12
,SUM(CASE
    WHEN CAST(DXCCSR_END013 AS INT64) = 1
        THEN 1
    ELSE 0
END) AS Count_13
,SUM(CASE

```

```

        WHEN CAST(DXCCSR_END014 AS INT64) = 1
            THEN 1
        ELSE 0
        END) AS Count_14
    ,SUM(CASE
        WHEN CAST(DXCCSR_END015 AS INT64) = 1
            THEN 1
        ELSE 0
        END) AS Count_15
    ,SUM(CASE
        WHEN CAST(DXCCSR_END016 AS INT64) = 1
            THEN 1
        ELSE 0
        END) AS Count_16
    ,SUM(CASE
        WHEN CAST(DXCCSR_END017 AS INT64) = 1
            THEN 1
        ELSE 0
        END) AS Count_17
    ,
FROM `capstone - 400517. capstone2.ENDOCRINE_DISEASES`
--Query to analyze asthma and length of stay
SELECT RECNUM
    ,LOS
FROM `capstone - 400517. capstone2.table_core`
WHERE (
    Blood_All <> 0
    OR MAL_All <> 0
    OR MBD_ALL <> 0
    OR END_ALL <> 0
    OR INF_All <> 0
    OR DIG_ALL <> 0
    OR NVS_ALL <> 0
    OR PRG_All <> 0
    OR CIR_All <> 0
    OR SKN_All <> 0
    OR MUS_All <> 0
    OR Tumor_All <> 0
    OR EYE_All <> 0
    OR EAR_All <> 0
)
AND Asthma <> '0'
AND Asthma IS NOT NULL
--Query to get count of individual infectious diseases
SELECT SUM(CASE
    WHEN CAST(DXCCSR_INF001 AS INT64) <> 0
        THEN 1
    ELSE 0
    END) AS Count_INF001
    ,SUM(CASE
    WHEN CAST(DXCCSR_INF002 AS INT64) <> 0
        THEN 1
    ELSE 0
    END) AS Count_INF002
    ,SUM(CASE
    WHEN CAST(DXCCSR_INF003 AS INT64) <> 0
        THEN 1
    ELSE 0
    END) AS Count_INF003

```

```

SUM(CASE
    WHEN CAST(DXCCSR_INF004 AS INT64) <> 0
    THEN 1
    ELSE 0
END) AS Count_INF004
SUM(CASE
    WHEN CAST(DXCCSR_INF005 AS INT64) <> 0
    THEN 1
    ELSE 0
END) AS Count_INF005
SUM(CASE
    WHEN CAST(DXCCSR_INF006 AS INT64) <> 0
    THEN 1
    ELSE 0
END) AS Count_INF006
SUM(CASE
    WHEN CAST(DXCCSR_INF007 AS INT64) <> 0
    THEN 1
    ELSE 0
END) AS Count_INF007
SUM(CASE
    WHEN CAST(DXCCSR_INF008 AS INT64) <> 0
    THEN 1
    ELSE 0
END) AS Count_INF008
SUM(CASE
    WHEN CAST(DXCCSR_INF009 AS INT64) <> 0
    THEN 1
    ELSE 0
END) AS Count_INF009
SUM(CASE
    WHEN CAST(DXCCSR_INF010 AS INT64) <> 0
    THEN 1
    ELSE 0
END) AS Count_INF010
SUM(CASE
    WHEN CAST(DXCCSR_INF011 AS INT64) <> 0
    THEN 1
    ELSE 0
END) AS Count_INF011
SUM(CASE
    WHEN CAST(DXCCSR_INF012 AS INT64) <> 0
    THEN 1
    ELSE 0
END) AS Count_INF012
FROM `capstone - 400517. capstone2.Infectious_DISEASES`
--Query to clean total charges col.
--ALTER TABLE `capstone-400517.capstone2.table_core`
--ADD COLUMN TotalChargesInteger INT64;
-- Copy data from the existing column to the new column
UPDATE `capstone - 400517. capstone2.table_core`
SET TotalChargesInteger = CASE
    WHEN TOTCHG = 'nan'
    THEN NULL -- or set to a default value
    WHEN TOTCHG = 'A'
    THEN NULL
    WHEN TOTCHG = 'C'
    THEN NULL
    WHEN TOTCHG = 'TOTCHG'

```



```

        THEN NULL
    ELSE CAST(TOTCHG AS INT64)
    END
WHERE 1 = 1

```

--Query to get LOS AND TOTAL CHARGE FOR PATIENTS WITH ASTHMA AND OTHER DISEASES

```

SELECT t1.DXCCSR_RSP009
      ,COUNT(*)
FROM (
    SELECT RECNUM
          ,DXCCSR_Default_DX1
          ,DXCCSR_RSP009
    FROM `capstone - 400517. capstone2.kid_GPRS` A11
    WHERE A11.DXCCSR_Default_DX1 = 'RSP009'
    ) t1
GROUP BY 1

```

--query to create separate table for CongenitalMalfunction

CREATE TABLE capstone2.CongenitalMalfunction_DISEASES AS

```

SELECT HOSP_KID
      ,RECNUM
      ,DXCCSR_MAL001
      ,DXCCSR_MAL002
      ,DXCCSR_MAL003
      ,DXCCSR_MAL004
      ,DXCCSR_MAL005
      ,DXCCSR_MAL006
      ,DXCCSR_MAL007
      ,DXCCSR_MAL008
      ,DXCCSR_MAL009
      ,DXCCSR_MAL010
FROM `capstone - 400517. capstone2.kid_GPRS`

```

--query to make new table for infectious diseases

CREATE TABLE capstone2.Infectious_DISEASES AS

```

SELECT HOSP_KID
      ,RECNUM
      ,DXCCSR_INF001
      ,DXCCSR_INF002
      ,DXCCSR_INF003
      ,DXCCSR_INF004
      ,DXCCSR_INF005
      ,DXCCSR_INF006
      ,DXCCSR_INF007
      ,DXCCSR_INF008
      ,DXCCSR_INF009
      ,DXCCSR_INF010
      ,DXCCSR_INF011
      ,DXCCSR_INF012
FROM `capstone - 400517. capstone2.kid_GPRS`

```

--Query to make range for LOS

```

UPDATE capstone2.table_core
SET LOS_Range = CASE
    WHEN LOS BETWEEN '0'

```

```

        AND '1'
        THEN '0-1'
    WHEN LOS BETWEEN '2'
        AND '3'
        THEN '2-3'
    WHEN LOS BETWEEN '4'
        AND '9'
        THEN '4-9'
    WHEN LOS > '10'
        THEN '10+'
    END
WHERE 1 = 1

```

```

--query to make range for total charges
UPDATE capstone2.table_core
SET TotalChargeRange = CASE
    WHEN TotalChargesInteger BETWEEN 0
        AND 12000
        THEN 1
    WHEN TotalChargesInteger BETWEEN 12001
        AND 20000
        THEN 2
    WHEN TotalChargesInteger BETWEEN 20001
        AND 40000
        THEN 3
    WHEN TotalChargesInteger BETWEEN 40001
        AND 80000
        THEN 4
    WHEN TotalChargesInteger > 80001
        THEN 5
    END
WHERE 1 = 1

```

```

# Python script for calculating Chi-Squared value and p value for asthma by gender
#import library
from scipy.stats import chi2_contingency
import matplotlib.pyplot as plt
import numpy as np

observed_data = [[101427, 1333965], [103168, 1248246]]

# Perform chi-square test
chi2, p, _, _ = chi2_contingency(observed_data)

#print p value
print(f"P-value: {p}")

print(f"Chi-squared value: {chi2}")

```

```

#Python script to check chi-square and P value for asthma by hospital region
#import library
from scipy.stats import chi2_contingency
import matplotlib.pyplot as plt
import numpy as np

observed_data = [[46993, 463696], [50325, 645687], [81598, 1116237], [25679, 356591]]

```

```

# Perform chi-square test
chi2, p, dof, expected = chi2_contingency(observed_data)
#print p value
print(f"P-value: {p}")

```

```

# Python script for chi-squared test for income and asthma
#import library
from scipy.stats import chi2_contingency
import matplotlib.pyplot as plt
import numpy as np
observed_data = [[74825, 804022], [48867, 634510], [45054, 617432], [33955, 499721]]
# Perform chi-square test
chi2, p, _, _ = chi2_contingency(observed_data)
#print p value
print(f"P-value: {p}")

print(f"Chi-squared value: {chi2}")

```

```

# Python script for chi-squared test: asthma with race
#import library
from scipy.stats import chi2_contingency
import matplotlib.pyplot as plt
import numpy as np
observed_data = [[78861, 1227805], [60569, 434845], [38607, 458916], [4366, 94119], [1550, 18515], [10290, 160175]]
# Perform chi-square test
chi2, p, _, _ = chi2_contingency(observed_data)
#print p value
print(f"P-value: {p}")

```

Code for clustering:

```

# Filtering only for RSP009
results = job.to_dataframe()
results = results[results['DXCSR_Default_DX1'] == 'RSP009']

```

```

# Import numpy to handle NaN values
import numpy as np

# Replace all instances of 'nan' with NaN in the results DataFrame
results.replace('nan', np.nan, inplace=True)

```

```

# Drop NA values from table
results.dropna(inplace=True)

```

```
results.replace('A', np.nan, inplace=True)
```

```
# Drop NA values form table  
results.dropna(inplace=True)
```

```
# check if we have NA values among unique values  
  
cols = results.columns  
for i in cols:  
  
    print(i, results[i].unique())
```

```
#Converting all values into numeric so that we can run ML algorithms  
  
import pandas as pd  
  
# Convert all columns to numeric, coercing errors to NaN  
results = results.apply(pd.to_numeric, errors='coerce')  
  
# Convert to Int64  
results = results.astype('Int64')
```

```
#Checking if the values converted into numeric  
  
cols = results.columns  
for i in cols:  
  
    print(i, results[i].unique())
```

```
#Converting all values into numeric so that we can run ML algorithms  
  
import pandas as pd  
  
# Convert all columns to numeric, coercing errors to NaN  
results = results.apply(pd.to_numeric, errors='coerce')  
  
# Convert to Int64  
results = results.astype('Int64')
```

```
#Checking if the values converted into numeric  
  
cols = results.columns  
for i in cols:
```

```
print(i, results[i].unique())
```

```
# Separating the features that we need for K-Mode algorithm
```

```
df_kmode = results[['AGE', 'FEMALE', 'RACE', 'HOSP_REGION', 'ZIPINC_QRTL', 'LOS', 'TOTCHG',  
'PAY1']]
```

```
# Creating bins for Length of stay column
```

```
## labels = ['0-1', '2-3', '4-9', '10 and over']
```

```
bins = [-1, 1, 3, 9, float('inf')] # Using -1 as the lower bound to include 0  
labels = [1, 2, 3, 4]
```

```
# Create a new column with the binned categories
```

```
df_kmode.loc[:, 'LOS'] = pd.cut(df_kmode['LOS'], bins=bins, labels=labels)
```

```
# Creating bins for "Total charge" feature
```

```
## labels = ['0-12000', '12,000-20,000', '20,000-40,000', '40,000-80,000', '80,000+']
```

```
bins = [-1, 12000, 20000, 40000, 80000, float('inf')] # Using -1 as the lower bound to include 0  
labels = [1, 2, 3, 4, 5]
```

```
# Create a new column with the binned categories
```

```
df_kmode.loc[:, 'TOTCHG'] = pd.cut(df_kmode['TOTCHG'], bins=bins, labels=labels)
```

```
# Creating bins for "AGE" feature of the model
```

```
## labels = ['0-1', '1-10', '11-20']
```

```
bins = [-1, 1, 10, 21] # Using -1 as the lower bound to include 0  
labels = [1, 2, 3]
```

```
# Create a new column with the binned categories
```

```
df_kmode.loc[:, 'AGE'] = pd.cut(df_kmode['AGE'], bins=bins, labels=labels)
```

```
# Install K-Modes ML library
```

```
!pip install kmodes
```

```
# Import necessary libraries from KModes and create clusters
```

```
from kmodes.kmodes import KModes
```

```
import pandas as pd
from sklearn.preprocessing import OneHotEncoder

encoder = OneHotEncoder(sparse=False)
df_encoded = encoder.fit_transform(df_kmode)

km = KModes(n_clusters=3, init='Huang', n_init=5, verbose=1)
```

```
#Check of the clusters created in vector format
```

```
df_encoded
```

```
# Fit the cluster
```

```
clusters = km.fit_predict(df_encoded)
```

```
# Checking the optimal number of clusters using Elbow method
```

```
import matplotlib.pyplot as plt
from kmodes.kmodes import KModes

costs = []
for num_clusters in range(1, 10):
    km = KModes(n_clusters=num_clusters, init='Huang', n_init=5, verbose=0)
    km.fit(df_encoded)
    costs.append(km.cost_)

plt.plot(range(1, 10), costs, marker='o')
plt.xlabel('Number of clusters')
plt.ylabel('Cost')
plt.title('Elbow Method For Optimal Number of Clusters')
plt.show()
```

```
# Join the cluster results to the original table
```

```
df_kmode['Cluster'] = clusters
```

```
# Using dataframe df_kmode: save to csv file for future analysis and visualization
```

```
df_kmode.to_csv('df_kmode_Asthma_primary_3.csv', index=False)
```

```
#Importing Library for kMeans
```

```
import pandas as pd
from sklearn.preprocessing import StandardScaler
from sklearn.cluster import KMeans
import matplotlib.pyplot as plt
```

```
#Scaling the dataset
scaler = StandardScaler()
scaled_features = scaler.fit_transform(df_asthma)
```

```
# Elbow Method
inertia = []
for i in range(1, 11):
    kmeans = KMeans(n_clusters=i, random_state=0)
    kmeans.fit(scaled_features)
    inertia.append(kmeans.inertia_)

plt.plot(range(1, 11), inertia)
plt.title('Elbow Method')
plt.xlabel('Number of clusters')
plt.ylabel('Inertia')
plt.show()
```

```
# Initialize KMeans with k clusters
kmeans = KMeans(n_clusters=5, random_state=0)

# Fit the model
kmeans.fit(scaled_features)

# Get the cluster labels for each data point
labels = kmeans.labels_
```

```
# Add the cluster labels as a new column in the original DataFrame
df_asthma['Cluster'] = labels

#Export result to CSV
df_asthma.to_csv("/df_cluster_new.csv")
```

-----End-----