Diabetes risk factors

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June 2, 2023

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1 Setup

I am interested in medical research, so I have chosen the Diabetes prediction dataset from Kaggle as the topic of my homework project. The goal of this dataset is to predict whether someone will develop diabetes, based on key indicators of the disease.

1.1 Input data

The dataset is anonymised and contains the following data about 100,000 individuals:

- Age: Ranges from 0-80, diabetes is more common amongst older adults.
- Gender: Can also have an impact on a person's susceptibility.
- Body Mass Index (BMI): Higher BMI values are linked to higher diabetes risk.
- Hypertension: Presistently elevated blood pressure in the arteries, linked to heart disease.
- **Heart disease**: Associated with a risk of developing diabetes.
- Smoking history: Considered as a risk factor, can worsen the complications of diabetes.
- **HbA1c level**: Hemoglobin A1c, measures blood sugar level over the past 2-3 months. Over 6.5% indicates diabetes.
- Blood glucose level: Key indicator of diabetes.
- Diabetes: Target value.

These are some of the key indicators of diabetes, along with demographic data, which could be used to determine risk factors for developing diabetes.

While it is not explicitly stated, the data is definitely on Type 2 diabetes, since Type 1 is a genetic condition.

1.2 Tools

I used Python, the numpy and pandas libraries for manipulation of the dataset, matplotlib, plotly and seaborn for plotting and visualising and scipy and scikit-learn for the various statistical analysis and evaluation tools they offer.

```
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import matplotlib.cm as cm
import matplotlib.colors as colors
import seaborn as sns
import plotly.express as px
from itertools import chain, combinations
from sklearn.linear_model import LinearRegression
from sklearn.model_selection import train_test_split
from sklearn.metrics import mean_squared_error
from scipy.stats import chi2_contingency
from scipy.stats import ttest_ind
```

1.3 Source code availability

The code and the dataset is available in my Github repository here:

https://github.com/nemkin/matstat-diabetes

1.4 Data cleaning and sanity checks

The first step in working with data is making sure it is not flawed. There is no single method that we can follow and guarantee no issues persist, but essentially we need to be careful and thourough and make sure nothing weird is happening in the dataset.

First, I read in the dataset from the csv file.

```
[221]: df = pd.read_csv('../data/diabetes_prediction_dataset.csv')
      df.head(10)
[221]:
         gender
                  age hypertension heart_disease smoking_history
                                                                      bmi
      O Female 80.0
                                  0
                                                 1
                                                             never 25.19
        Female 54.0
                                  0
                                                 0
                                                           No Info 27.32
      1
           Male 28.0
      2
                                  0
                                                 0
                                                             never 27.32
        Female 36.0
                                  0
                                                 0
                                                           current
                                                                   23.45
           Male 76.0
                                  1
                                                 1
                                                           current 20.14
        Female 20.0
                                  0
                                                             never 27.32
```

6 7 8 9	Female Female Male Female	42.0	0 0 0 0	0 0 0	never No Info never never	19.31 23.86 33.64 27.32
	HbA1c_1	evel	blood_glucose_level	diabetes		
0		6.6	140	0		
1		6.6	80	0		
2		5.7	158	0		
3		5.0	155	0		
4		4.8	155	0		
5		6.6	85	0		
6		6.5	200	1		
7		5.7	85	0		
8		4.8	145	0		
9		5.0	100	0		

I like check the value ranges for the data and make sure, that the columns are correctly typed:

[222]: df.dtypes

[222]: gender object float64 age int64 hypertension heart_disease int64 smoking_history object bmi float64 HbA1c_level float64 int64 blood_glucose_level diabetes int64 dtype: object

We can use the describe method to check the value ranges of the columns:

[223]: df.describe(include='all')

[223]:		gender	age	hypertension	heart_disease	smoking_history	
	count	100000	100000.000000	100000.00000	100000.000000	100000	\
	unique	3	NaN	NaN	NaN	6	
	top	Female	NaN	NaN	NaN	No Info	
	freq	58552	NaN	NaN	NaN	35816	
	mean	NaN	41.885856	0.07485	0.039420	NaN	
	std	NaN	22.516840	0.26315	0.194593	NaN	
	min	NaN	0.080000	0.00000	0.000000	NaN	
	25%	NaN	24.000000	0.00000	0.000000	NaN	
	50%	NaN	43.000000	0.00000	0.000000	NaN	
	75%	NaN	60.000000	0.00000	0.000000	NaN	
	max	NaN	80.000000	1.00000	1.000000	NaN	

	bmi	${ t HbA1c_level}$	blood_glucose_level	diabetes
count	100000.000000	100000.000000	100000.000000	100000.000000
unique	NaN	NaN	NaN	NaN
top	NaN	NaN	NaN	NaN
freq	NaN	NaN	NaN	NaN
mean	27.320767	5.527507	138.058060	0.085000
std	6.636783	1.070672	40.708136	0.278883
min	10.010000	3.500000	80.000000	0.000000
25%	23.630000	4.800000	100.000000	0.000000
50%	27.320000	5.800000	140.000000	0.000000
75%	29.580000	6.200000	159.000000	0.000000
max	95.690000	9.000000	300.000000	1.000000

This shows, that gender and smoking_history are actually categorical columns (they have a few unique values), while hypertension, heart_disease, and diabetes are meant to be boolean, and finally age should be an integer.

```
[224]: df['age'] = df['age'].astype(int)
    df['gender'] = df['gender'].astype('category')
    df['smoking_history'] = df['smoking_history'].astype('category')
    df['hypertension'] = df['hypertension'].astype(bool)
    df['heart_disease'] = df['heart_disease'].astype(bool)
    df['diabetes'] = df['diabetes'].astype(bool)
```

```
[224]: gender
                               category
                                  int32
       age
       hypertension
                                   bool
       heart_disease
                                   bool
       smoking_history
                               category
       bmi
                                float64
       HbA1c_level
                                float64
       blood_glucose_level
                                  int64
       diabetes
                                   bool
       dtype: object
```

Let's check our categorical values:

```
[225]: columns = df.select_dtypes(include='category').columns.tolist()

for column in columns:
    values = sorted(list(df[column].unique()))
    print(column)
    print(values)
    print()
```

gender

```
['Female', 'Male', 'Other']
smoking_history
['No Info', 'current', 'ever', 'former', 'never', 'not current']
```

I would like to fix the inconsistent values in smoking_history:

```
gender
['Female', 'Male', 'Other']
smoking_history
```

['current', 'ever', 'former', 'never', 'no_info', 'not_current']

1.4.1 Analysing the column values and their distribution

When our data comes from a third party, and we have not collected it ourselves, it is a good idea to check for any inconsistencies in the values.

First, I checked that no values were missing.

```
[227]: df.isna().sum()
[227]: gender
                               0
                               0
       age
       hypertension
                               0
       heart_disease
                               0
       smoking_history
                               0
                               0
       bmi
       HbA1c_level
       blood_glucose_level
                               0
       diabetes
                               0
       dtype: int64
```

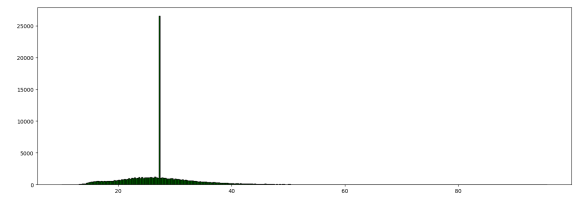
If there were missing values, we could drop them with df.dropna(inplace=True), but it is not needed.

BMI The first weirdness I noticed was with BMI.

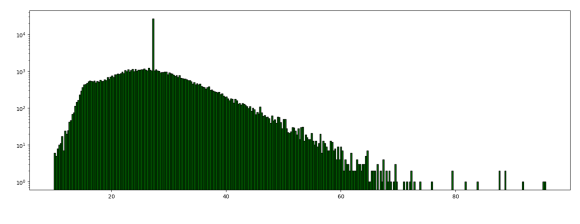
Let us plot the values of BMI in a histogram:

```
[228]: values = df['bmi']
counts, bins, _ = plt.hist(values, bins='auto', color='darkgreen', ___
→edgecolor='black')

plt.gcf().set_size_inches(18, 6)
plt.show()
```



That doesn't look good. I kept increasing the bins, before leaving them auto, but that singular peak will not disappear. Let's try logarithmic scale, so we can at least see something:



Is that a single value of BMI? Let's count the most common values:

```
[230]: max_count_bin = np.argmax(counts)
      max_count = counts[max_count_bin]
      max_bin_range = (bins[max_count_bin], bins[max_count_bin + 1])
      max_bin_range
      unique_values, counts = np.unique(values, return_counts=True)
      sorted_indices = np.argsort(counts)[::-1]
      sorted_values = unique_values[sorted_indices]
      sorted_counts = counts[sorted_indices]
      top_10_values = sorted_values[:10]
      top_10_counts = sorted_counts[:10]
      print(top_10_values)
      print(top_10_counts)
                                                              24.5]
                   27.12 27.8 24.96 22.4
                                            25.
                                                  25.6 26.7
                                               99
      [25495
               103
                     101
                           100
                                  100
                                         99
                                                     98
                                                           94
                                                                  94]
```

Yes, for some reason, we have over $\frac{1}{4}$ entries of the dataset with the exact BMI of 27.32. Let's look at these:

```
[231]: weird_entries = df[df['bmi'] == 27.32]
weird_entries.head(10)
```

```
[231]:
           gender
                    age hypertension heart_disease smoking_history
                                                                           bmi
       1
           Female
                                 False
                                                 False
                                                               no_info
                                                                         27.32
       2
             Male
                     28
                                 False
                                                 False
                                                                  never
                                                                         27.32
       5
           Female
                     20
                                 False
                                                 False
                                                                         27.32
                                                                  never
           Female
       9
                     32
                                False
                                                False
                                                                  never
                                                                         27.32
       10 Female
                     53
                                False
                                                False
                                                                  never
                                                                         27.32
       14 Female
                                False
                                                                         27.32
                     76
                                                False
                                                                no_info
             Male
                                                                no_info 27.32
       15
                                False
                                                False
       18
           Female
                     42
                                False
                                                 False
                                                                no_info 27.32
             Male
                                False
                                                                        27.32
       26
                     67
                                                  True
                                                           not_current
             Male
       38
                     50
                                  True
                                                False
                                                                current 27.32
                         blood_glucose_level
           HbA1c_level
                                               diabetes
                    6.6
                                           80
                                                   False
       1
                    5.7
       2
                                          158
                                                   False
       5
                    6.6
                                           85
                                                   False
       9
                    5.0
                                          100
                                                   False
       10
                    6.1
                                           85
                                                   False
       14
                    5.0
                                          160
                                                   False
       15
                    6.6
                                          126
                                                   False
                    5.7
       18
                                           80
                                                   False
                    6.5
                                                    True
       26
                                          200
```

38 5.7 260 True

These all seem like real entries, not duplicates of the same entry. My best guess for what happened, is that when the BMI was missing, they put down the mean of the rest of the dataset, which is exactly 27.32.

```
[232]: df[df['bmi'] != 27.32]['bmi'].mean()
```

[232]: 27.321029595329176

In a real life scenario, I would clarify this with the person who gave me the data, however for this homework project, I will assume that when a person looks average weight, they may not bother measuring their BMI, so the average is probably a good approximation, so I will accept this.

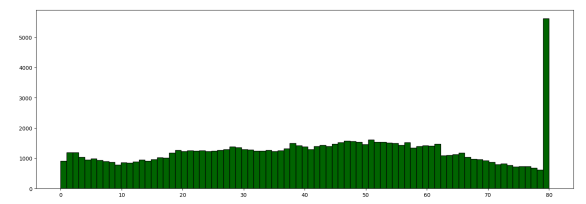
Another concern I had was with extreme cases of BMI, such as the values 80 and above. I choose to keep these, because they are not overrepresented and diabetes is linked to extreme obesity, so these are important entry points for prediction.

Age Next issue is with the age.

Let's plot the values:

```
[233]: values = df['age']
num_bins = values.nunique()
counts, bins, _ = plt.hist(values, bins=num_bins, color='darkgreen', □
→edgecolor='black')

plt.gcf().set_size_inches(18, 6)
plt.show()
```



We have a lot of people aged 80. It looks like older people are overrepresented in this data and probably not all of them are exactly 80, they just cut off the age to fit into a [0,80] interval.

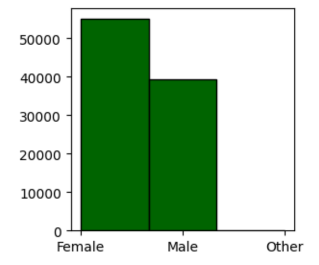
```
[234]: df[df['age']==80]['age'].count()
```

[234]: 5621

In this case, I choose to remove these values, because I feel like this could cause some skewing.

```
[235]: df = df[df['age'] < 80]
```

Gender Let's plot the values:



The category Other can mean many things. For medical research, we are concerned about the biological sex of the participants, so I will remove these entries too.

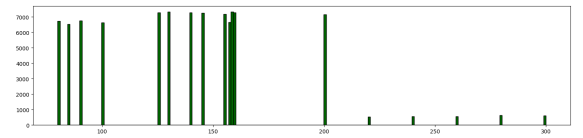
```
[237]: df = df[df['gender'] != 'Other']
df['gender'] = df['gender'].cat.remove_categories('Other')
df['gender'].unique()
```

Blood glucose level

```
[238]: values = df['blood_glucose_level']
counts, bins, _ = plt.hist(values, bins=200, color='darkgreen', 

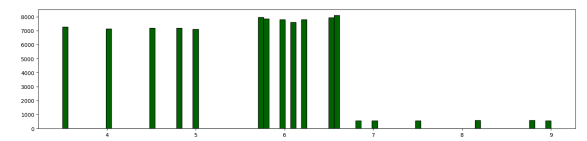
→edgecolor='black')
```

```
plt.gcf().set_size_inches(18, 4)
plt.show()
```



It seems like this is not a continuous spectrum. There is also a concentration of values around 155-160. I'm not sure how to interpret this. The values are probably in mg/dl. For fasting levels, 80-100 is normal, 101-125 is elevated and above 126 is high. Other than that, I would need to ask someone with medical knowledge on why this is happening. For now, I have to accept these, as there is no clear way on how to fix them or if they need to be fixed at all.

Blood sugar level



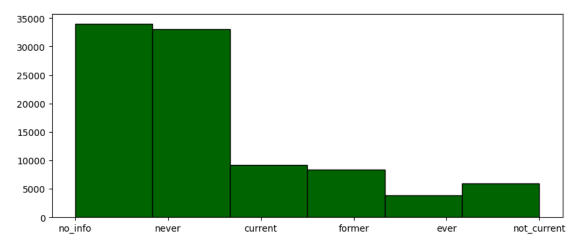
For HbA1c, less than 5.6% is normal, between 5.7% - 6.4% is elevated (prediabetes) and above 6.5% is high (diabetes).

Although increasing the granuality does reveal that not every value is represented, but the precision is only one digit after the integer, so I think this looks okay.

Smoking history

```
[240]: values = df['smoking_history']
num_bins = values.nunique()
counts, bins, _ = plt.hist(values, bins=num_bins, color='darkgreen', □
→edgecolor='black')

plt.gcf().set_size_inches(10, 4)
plt.show()
```



We have a large amount of entries with no_info as a value. I was debating whether to remove these, or not, but I ended up keeping them, because they represent a large chunk of our data.

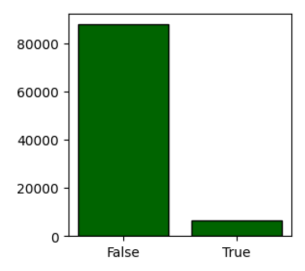
I was also questioning what ever means, given the other possibilities present, I believe it means they have tried smoking in the past, but weren't addicted.

Hypertension

```
[241]: df['hypertension'].value_counts()

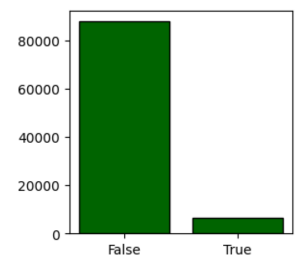
value_counts = df['hypertension'].value_counts()

plt.figure(figsize=(3,3))
plt.bar(value_counts.index, value_counts.values, color='darkgreen',u edgecolor='black')
plt.xticks(ticks=[0,1], labels=['False', 'True'])
plt.show()
```



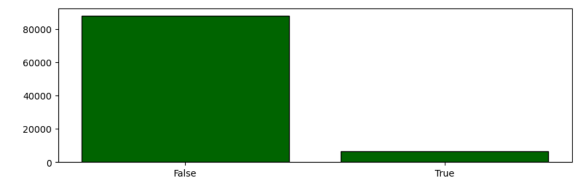
It seems like a reasonable distribution.

Heart disease



Similarly, this looks okay.

Diabetes Diabetes is our target value.



This seems like a good ratio, similar to the generic population.

2 Analysis

2.1 Complete multivariate linear regression

The first thing I will try is create a multivariate linear regression model, using all of the available variables.

In order to do this, we must convert the categorical variables into 0/1 variables. This can be done with the get_dummies function, as seen below.

```
[244]: df_encoded = pd.get_dummies(df) df_encoded.head()
```

```
[244]:
                                                                      blood_glucose_level
                hypertension
                               heart_disease
                                                  bmi
                                                        HbA1c_level
       1
           54
                        False
                                        False
                                                27.32
                                                                6.6
                                                                                         80
                                                                                             \
                                                27.32
                                                                5.7
       2
            28
                        False
                                        False
                                                                                        158
       3
                                        False
                                                23.45
                                                                5.0
            36
                        False
                                                                                        155
       4
           76
                         True
                                         True
                                                20.14
                                                                4.8
                                                                                        155
       5
            20
                        False
                                        False
                                                27.32
                                                                6.6
                                                                                         85
```

```
gender_Female gender_Male smoking_history_no_info
1
      False
                       True
                                    False
                                                                True
      False
                                     True
                                                               False
2
                      False
3
      False
                       True
                                    False
                                                               False
4
      False
                      False
                                     True
                                                               False
5
      False
                       True
                                    False
                                                               False
   smoking_history_current
                             smoking_history_ever smoking_history_former
                                             False
                                                                       False
1
                      False
2
                      False
                                             False
                                                                       False
3
                       True
                                             False
                                                                       False
4
                       True
                                             False
                                                                       False
5
                      False
                                             False
                                                                       False
   smoking_history_never
                           smoking_history_not_current
                    False
                                                   False
1
2
                     True
                                                   False
3
                    False
                                                   False
4
                    False
                                                   False
5
                                                   False
                     True
```

Then, we separate our target variable.

```
[245]: X = df_encoded.drop('diabetes', axis=1)
y = df_encoded['diabetes']
```

And split the data into training and testing datasets, with the ratio of 80% and 20%. It is good practice to seed the pseudorandom generator, so it will always result in the same split, across multiple runs of our notebook.

```
[246]: X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, 

→random_state=42)
```

Then we create our multivariate linear regression model, using scikit-learn.

The model's coefficients are as follows:

print(intercept)

Coefficients:

```
Variable
                                  Coefficient
                                      0.133472
2
                   heart_disease
1
                    hypertension
                                      0.103687
                     HbA1c_level
4
                                      0.076937
         smoking_history_former
11
                                      0.015804
7
                     gender_Male
                                      0.006450
3
                                      0.004050
                             bmi
5
            blood_glucose_level
                                      0.002163
13
    smoking_history_not_current
                                      0.001371
0
                                      0.001358
           smoking_history_ever
10
                                     -0.001578
9
        smoking_history_current
                                     -0.001865
12
          smoking_history_never
                                     -0.003399
                   gender_Female
6
                                     -0.006450
8
        smoking_history_no_info
                                     -0.010332
```

Intercept:

```
Variable Coefficient
Unitercept -0.814089
```

It is important to note here immediately, that the scale of these coefficients depends on their value sets. For example, gender_Male is between 0 and 1, while BMI is between 10 and 100. For this reason, we cannot compare the relative values of the coefficients, without taking into account the possible values of the variables behind them.

However, the signs (positive or negative) can be looked at and compared, which we will do shortly.

But first, the precision of this model on the test dataset is as follows:

```
[249]: y_pred = model.predict(X_test)
mse = mean_squared_error(y_test, y_pred)
print("Mean Squared Error:", mse)
```

Mean Squared Error: 0.04693052558568325

And on the train dataset:

```
[250]: y_pred = model.predict(X_train)
mse = mean_squared_error(y_train, y_pred)
print("Mean Squared Error:", mse)
```

Mean Squared Error: 0.048474640595590854

We can conclude, that the model was not overfit and the error is relatively low.

2.2 Independence testing

2.2.1 **Gender**

Examining the coefficients, the first thing I noticed is the gender variables. It seems that, with all other variables present, being male slightly increases, while being female slightly reduces the risk of diabetes.

I wonder if without the other variables, what could we say about the influence of gender on diabetes? I will test the independence of the **gender** and **diabetes** variables in the original dataset using Chisquare test of independence.

At first, I will create the frequency table of their values:

```
[251]: crosstab = pd.crosstab(df['gender'], df['diabetes'])
crosstab
```

```
[251]: diabetes False True gender
Female 51188 3856
Male 35697 3620
```

Then, I will use scipy, to compute a Pearson's chi-squared statistic.

The null hypothesis is that the variables are independent.

```
[252]: chi2, p, dof, expected = chi2_contingency(crosstab)
    print(f"Chi-square value: {chi2}")
    print(f"P-value: {p}")
    print(f"Degrees of freedom: {dof}")
    print("Expected:")
    print(expected)
```

```
Chi-square value: 152.1271942474673
P-value: 5.94328803769177e-35
Degrees of freedom: 1
Expected:
[[50682.9933977 4361.0066023]
[36202.0066023 3114.9933977]]
```

With a p-value of $p = 5,94 \cdot 10^{-35}$ we can say, that we have very strong evidence against our null hypothesis, therefore we can conclude that gender and diabetes are indeed dependent:

The gender of a person influences the likelihood of developing diabetes.

As we can see in the Expected matrix, that is the expected frequencies for the same population size, were these variables independent of each other. We can see that we got less than expected females with diabetes and more than expected males with diabetes.

2.2.2 Smoking history

The other interesting thing I noticed about the multivariate linear regression coefficients is that smoking_history_current has a negative coefficient, while smoking_history_former and

smoking_history_not_current have positive coefficients. At least smoking_history_never has a negative coefficient too, which makes perfect sense.

I do have tales from friends and family saying that when someone chooses to stop smoking is when the problems start. A more likely explanation is that when people stop smoking, the damage has already been done. When someone is still smoking, maybe they have less years behind them and quitting smoking **earlier** should help lower the risk of diabetes and heart disease (amongst many diseases).

I will also run the same independence test on smoking_history and diabetes:

```
[253]: crosstab = pd.crosstab(df['smoking_history'], df['diabetes'])
crosstab
```

```
[253]: diabetes
                          False
                                 True
       smoking_history
       no_info
                          32751
                                   1260
       current
                           8227
                                    924
       ever
                           3376
                                    436
       former
                           7004
                                   1372
       never
                          30181
                                   2878
       not_current
                           5346
                                    606
```

```
[254]: chi2, p, dof, expected = chi2_contingency(crosstab)
    print(f"Chi-square value: {chi2}")
    print(f"P-value: {p}")
    print(f"Degrees of freedom: {dof}")
    print("Expected:")
    print(expected)
```

Chi-square value: 1844.0298560488386

P-value: 0.0

Degrees of freedom: 5

Expected:

```
[[31316.38849737 2694.61150263]

[8425.98780216 725.01219784]

[3509.98420958 302.01579042]

[7712.3892286 663.6107714]

[30439.81321732 2619.18678268]

[5480.43704497 471.56295503]]
```

The p-value is so small, the numerical representation displays it as 0. This means smoking history has indeed an influence on whether or not someone develops diabetes.

I would like to further investigate this relationship. Since diabetes is a binary variable, it is hard to see exactly the effects of smoking on it. However, we have a few numerical variables in the dataset, which are strong indicators of diabetes, such as the various measurements of blood levels and BMI.

So I will continue exploring the effects of smoking on these variables further.

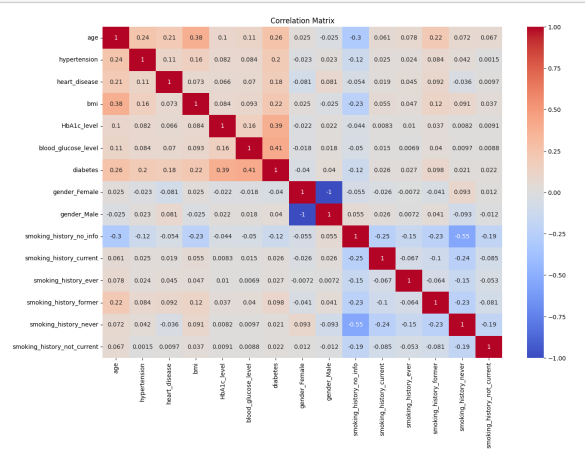
2.3 Correlation matrix

In order to establish which numerical variables have a strong relationship with diabetes, I will be using a correlation matrix heatmap to visualise these 1-1 relationships.

```
[255]: correlation_matrix = df_encoded.corr()

plt.figure(figsize=(15, 10))
    sns.heatmap(correlation_matrix, annot=True, cmap='coolwarm', center=0)

plt.title('Correlation Matrix')
    plt.show()
```



We can see on the heatmap, that blood_glucose_level and HbA1c_level have the strongest correlation to diabetes, so I will be exploring the effects of smoking on these variables.

2.4 Testing the mean of categories

Smoking history is a categorical variable, which means it defines different groups of people. For these groups, I can calculate the mean of blood_glucose_level and HbA1c_level, then, I can investigate whether or not the differences in the results are statistically significant.

```
[256]: df.groupby('smoking_history')[['blood_glucose_level']].mean().

→sort_values('blood_glucose_level')
```

Here, we can see that the average blood glucose level slightly increases, as someone has more history with smoking. Interestingly again, current smokers rank below former smokers. This can again be due to the fact that former smokers may have a longer smoking history than current smokers.

```
[257]: df.groupby('smoking_history')[['HbA1c_level']].mean().sort_values('HbA1c_level')
```

[257]:		HbA1c_level
	smoking_history	
	no_info	5.457690
	never	5.531640
	current	5.546694
	not_current	5.557023
	ever	5.571800
	former	5.646036

However, for HbA1c_level, the level of ever, which I assume means "smoked a bit a long time ago" seems really high on the ranking.

I would like to know if there is a statistically significant difference between the average $HbAlc_levels$ of the $smoking_history$ categories. ANOVA would not tell me which specific groups are different, so instead I will perform pairwise hypothesis testing for the equivalence of the mean between all $5 \cdot 5$ category pairs and display the resulting p-values in a matrix heatmap format, similar to the correlation matrix.

The diagonal of this matrix will be comparing the same variables, but this is also the case for the correlation matrix, so I will keep those values in as well.

I could use paired t-tests, but that would require knowing that the variances are the same. I will be using Welch-test instead, which is more general and has a similar built-in function in scipy.

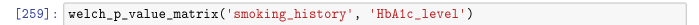
I will be calling this the Welch p-value matrix. I have tried finding such a thing in the literature, but I wasn't able to. For example MANOVA tests multiple dependent variables, while I have a single dependent variable and I want all pairwise relations between the various categories.

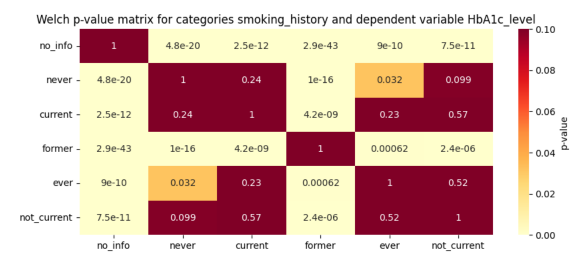
Let us first define the function:

```
[258]: def welch_p_value_matrix(category, target):
    categories = df[category].unique()
```

```
p_val_matrix = np.zeros((len(categories), len(categories)))
for i, category_i in enumerate(categories):
  for j, category_j in enumerate(categories):
     category_data_i = df[df[category] == category_i][target]
     category_data_j = df[df[category] == category_j][target]
     # Performs Welch's t-test, when equal_var = False is set.
     t_stat, p_val = ttest_ind(category_data_i, category_data_j,_
→equal_var=False)
     p_val_matrix[i, j] = p_val
plt.figure(figsize=(10, 4))
sns.heatmap(p_val_matrix, cmap='YlOrRd', annot=True, fmt=".2g",
           xticklabels=categories, yticklabels=categories, vmax=0.1,
           cbar_kws={'label': 'p-value'})
plt.title(f"Welch p-value matrix for categories {category} and dependent,
→variable {target}")
plt.show()
```

Then let's run this function for the HbA1c_level!





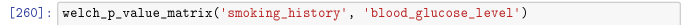
If the cell is red, it means there is no statistically significant difference between the two categories for the HbA1c_level. If the cell is orange, we can say that there is a statistically significant difference (p < 0.05). Finally, when the cell is light yellow, the p value is even smaller, p < 0.01.

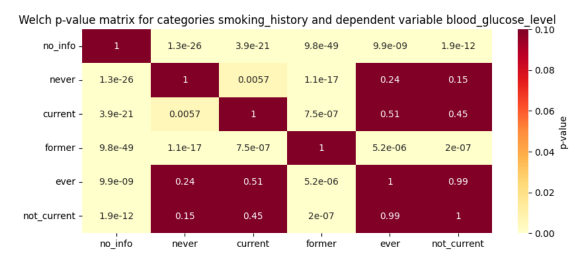
Interestingly, if someone is a former smoker, which I assume means "has been addicted to smoking and quit", the increase in the average HbA1c_levels is statistically significant, relative to all other groups, including current smokers!

Other than this, no_info had a similar result. Since no_info has the smallest average among all of the categories, so I believed no_info should be similar to never, however they statistially differ.

Finally, ever and never also have a statistically significant difference, ever having the higher average between the two. So if someone has smoked in the past, that can still put them in a higher risk category for increased HbA1c_levels.

Now, let us look at the target variable blood_glucose_level!





We have similar results for the categories no_info and former. However, this time current and never also have a statistically significant difference, current having an increased average blood_glucose_level.

After looking at these results, my conclusion is that categorical representation of smoking_history seems unfit for statistical purposes. A better way to represent smoking would be giving the number of months smoked during someone's lifetime and the number of months after their last smoke. I believe these variables would better represent the differences than these categories.

2.5 Selected multivariate linear regression

Finally, I would like to form the best multivariate regression model for predicting diabetes.

How do I define best?

I want to use only a subset of the variables, minimizing the number of them needed, which still do a relatively good job at minimizing the MSE.

In order to find the best subset, I will first calculate the MSE for all subset of variables (the power set of the columns), then I will plot these against the number of variables needed, on a scatter plot.

First, I have created a function, that can do the same linear regression I did previously, but on the variables, specified on the input parameter:

```
[264]: def linear_regression(variables, should_print=False):
        X = df_encoded[variables]
        y = df_encoded['diabetes']
        X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2,_
       \rightarrowrandom_state=42)
        model = LinearRegression()
        model.fit(X_train, y_train)
        coefficients = pd.DataFrame({'Variable': X.columns, 'Coefficient': model.
       intercept = pd.DataFrame({'Variable': ['Intercept'], 'Coefficient': model.
       →intercept_})
        y_pred = model.predict(X_test)
        mse = mean_squared_error(y_test, y_pred)
        if should_print:
          print("Coefficients:")
          print(coefficients)
          print()
          print("Intercept:")
          print(intercept)
          print()
          print("Mean Squared Error:")
          print(mse)
        return mse
```

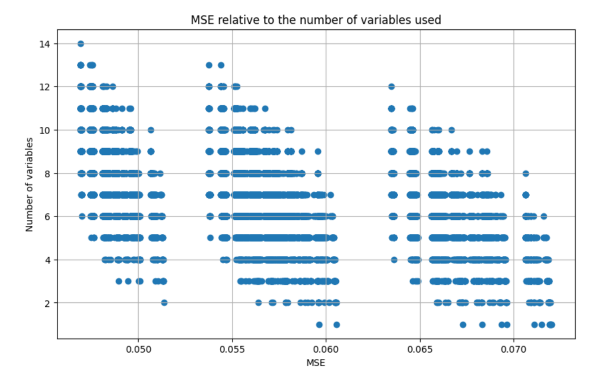
Then I ran this for all column subsets (excluding the empty one) and collected the results:

```
})
```

Using the following code, I was able to display the actual variables needed as hover text for every data point. However, this plot is only visible in the Jupyter notebook it is not displayed when I convert it to a PDF. For the PDF, a similar plot, without the hover follows next.

For the PDF, here is the same plot, without the hover text:

```
[276]: results_df = pd.DataFrame(results)
    plt.figure(figsize=(10, 6))
    plt.scatter(results_df['mse'], results_df['num_vars'])
    plt.xlabel('MSE')
    plt.ylabel('Number of variables')
    plt.title('MSE relative to the number of variables used')
    plt.grid(True)
    plt.show()
```



After examining the labels for the three distinct diamond-shaped blobs on the image above, I see

that the first blob, with the least MSE contains both HbA1c_level and blood_glucose_level, the second blob contains one of the two, finally the third blob with the most MSE contains neither.

It is clear, that these two variables are necessary, to improve prediction. Moreover, they result in the lowest MSE when using two variables only, an MSE of 0.051. This is the bottom dot of the first diamond.

Adding age to them results in an MSE of 0.049 for three variables, adding bmi results in an MSE of 0.048 for four variables, adding heart_disease results in an MSE of 0.0475 for 5 variables, finally adding hypertension results in 0.0469 for six variables.

Further variables don't decrease the MSE significantly.

I will print these best linear regression models for all number of variables:

```
[277]: min_mse_columns = results_df.groupby('num_vars').apply(lambda x: x.loc[x['mse'].
        →idxmin(), 'columns'])
[278]: n vars = 1
      cols = min_mse_columns[n_vars]
      print(f"Number of variables: {n_vars}")
      print()
      _ = linear_regression(cols, True)
      Number of variables: 1
      Coefficients:
                    Variable Coefficient
         blood_glucose_level
                                 0.002731
      Intercept:
          Variable Coefficient
      0 Intercept
                      -0.296492
      Mean Squared Error:
      0.05965061175137981
[279]: n_{vars} = 2
      cols = min_mse_columns[n_vars]
      print(f"Number of variables: {n_vars}")
       _ = linear_regression(cols, True)
      Number of variables: 2
      Coefficients:
                    Variable Coefficient
                 HbA1c level
                                 0.084451
        blood_glucose_level
                                 0.002379
      Intercept:
```

```
0 Intercept
                   -0.714191
      Mean Squared Error:
      0.051362749971308484
[280]: n_{vars} = 3
       cols = min_mse_columns[n_vars]
       print(f"Number of variables: {n_vars}")
       print()
       _ = linear_regression(cols, True)
      Number of variables: 3
      Coefficients:
                    Variable Coefficient
      1
                 HbA1c_level
                                0.080259
      0
                                 0.002462
                         age
      2 blood_glucose_level
                                 0.002255
      Intercept:
          Variable Coefficient
      0 Intercept
                      -0.771513
      Mean Squared Error:
      0.048955894608910165
[281]: n_{vars} = 4
       cols = min_mse_columns[n_vars]
       print(f"Number of variables: {n_vars}")
       print()
       _ = linear_regression(cols, True)
      Number of variables: 4
      Coefficients:
                    Variable Coefficient
                 HbA1c_level
                                 0.079186
      1
                         bmi
                                 0.004354
      3
        blood_glucose_level
                                 0.002223
                         age
                                 0.001940
      Intercept:
          Variable Coefficient
      0 Intercept
                      -0.859603
      Mean Squared Error:
      0.04820938328094974
```

Variable Coefficient

```
[282]: n_{vars} = 5
       cols = min_mse_columns[n_vars]
       print(f"Number of variables: {n_vars}")
       print()
       _ = linear_regression(cols, True)
      Number of variables: 5
      Coefficients:
                     Variable Coefficient
      1
               heart_disease
                                  0.147063
      3
                  HbA1c level
                                  0.078263
      2
                          bmi
                                  0.004424
      4
         blood_glucose_level
                                  0.002196
                                  0.001681
                          age
      Intercept:
          Variable
                    Coefficient
         Intercept
                       -0.847087
      Mean Squared Error:
      0.04750629195281252
[283]: n_{vars} = 6
       cols = min_mse_columns[n_vars]
       print(f"Number of variables: {n_vars}")
       print()
       _ = linear_regression(cols, True)
      Number of variables: 6
      Coefficients:
                     Variable Coefficient
      2
               heart_disease
                                  0.137909
      1
                 hypertension
                                  0.105400
      4
                  HbA1c_level
                                  0.077108
                                  0.004134
      3
                          bmi
      5
         blood_glucose_level
                                  0.002167
                                  0.001437
                          age
      Intercept:
          Variable
                     Coefficient
         Intercept
                       -0.826001
      Mean Squared Error:
      0.04699444731206997
```

In conclusion, the blood level variables seem the most important and also most difficult to measure. Adding trivially measurable variables, such as the age and bmi improves the MSE by 0.003, so by

default I would use the 4 variable model.

If history of heart-related issues is known, then I would also measure hypertension, which is directly related to heart disease and use the 6 variable model for diabetes prediction.