Estimating Wine Quality from a 12-Dimensional Data Set

Report of CS212 Intelligent Data Analysis

MADE BY

Ziyuan Ye

11610203@mail.sustc.edu.cn

UNDER THE GUIDANCE OF

Peter Tiňo

AND

Guoji Fu



Department of Computer Science and Engineering

Southern University of Science and Technology

Shenzhen, China, July 2018

Abstract

This assignment focus on analysing the data set with the technique such as **K-means** Clustering, Self Organizing Map (SOM) and Principal Component Analysis (PCA). I try to combine the above methods work out some problems of a data set which is called **wine-quality**.)) These attributes of the data set seem not correlate with each other. However, there must exist some hidden relationship in the data set. **eg: wine quality must be influence by some attribute**. With the reduce of dimension, gradually the relationship between the attributes were disclosed. There are two data set in my folder, in these assignment I only focus on the red wine-quality data set.

Detail of Red Wine Quality Data Set

12 Dimensions

- Fixed acidity (g(tartaric acid)/dm3), range from 4.6 to 15.9 mean value is 8.3
- Volatile acidity (g(acetic acid)/dm3), range from 0.1 to 1.6 mean value is 0.5
- Citric acid (g/dm3), range from 0.0 to 1.0 mean value is 0.3
- Residual sugar (g/dm3), range from 0.9 to 15.5 mean value is 2.5
- Chlorides (g(sodium chloride)/dm3), range from 0.01 to 0.61 mean value is 0.08
- Free sulfur dioxide (mg/dm3), range from 1 to 72 mean value is 14
- Total sulfur dioxide (mg/dm3), range from 6 to 289 mean value is 46
- Density (g/cm3), range from 0.990 to 1.004 mean value is 0.996
- pH, range from 2.7 to 4.0 mean value is 3.3
- Sulphates (g(potassium sulphate)/dm3), range from 0.3 to 2.0 mean value is 0.7
- Alcohol (vol.%), range from 8.4 to 14.9 mean value is 10.4
- Quality, range from 3 to 8 mean value is 5.6

Data Set Source:

https://archive.ics.uci.edu/ml/machine-learning-databases/wine-quality/

Two Main Topic

In my assignment, there are two question I want to analysis:

- How does the quality of wine influenced by all the attribute in the data set?
- If there are any attribute that influence total sulfur dioxide?

Preprocessing Data

Normalization

The **unit** and **scale** of the data is different, this method can make different characters have the same scale. Only in this way, the comparision can be valid.

$$\widehat{E[X_i]} \approx \frac{1}{N} \sum_{j=1}^{N} X_i^j \widehat{Var[X_i]} \approx \frac{1}{N} \sum_{j=1}^{N} (X_i^j - \widehat{E[X_i]})^2 = \sigma^2 \widehat{X_{i,nor}^j} \approx \frac{X_i^j - E[X_i]}{\sigma}$$

Chapter 1

Topic 1: How does the quality of wine influenced by all the attribute in the data set?

1.1 Labelling Strategy

When analysing the first problem, the category of quality is the main attribute we want to focus. By analysis the data set of red wine, I discover that quality is range from 3 to 8. So there come up with two strategy that I would like to use.

1.1.1 Labelling Strategy 1

Obviously, the quality of wine can be separated into two class: Acceptable, Recommended. According to the number of different quality of wine. I intuitively partition them as follow.

- Class 1: Quality in range [3, 5].
- Class 2: Quality in range (5, 8].

The final labelling result is:

- Class 1: 744 cases.
- Class 2: 855 cases.

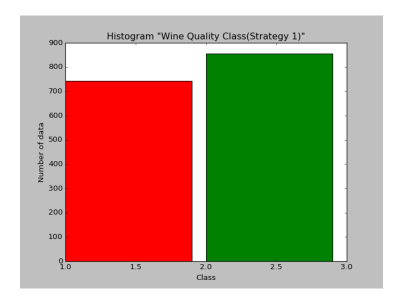


Figure 1.1: Labeling Strategy 1

Histogram of Quality

1.1.2 Labelling Strategy 2

According to James Halliday from James Halliday Annual Wine Companion, he separate wine quality into 5 class which are:Outstanding wines, Highly recommended, Recommended, Acceptable and Others. Thus I separate the data into 5 class as he did.

- Class 1: Quality in range [3, 4].
- Class 2: Quality equals 5
- Class 3: Quality equals 6
- Class 4: Quality equals 7
- Class 5: Quality equals 8

The final labelling result is:

- Class 1: 63 cases.
- Class 2: 681 cases.
- Class 3: 638 cases.
- Class 4: 199 cases.
- Class 5: 199 cases.

Histogram of Alcohol

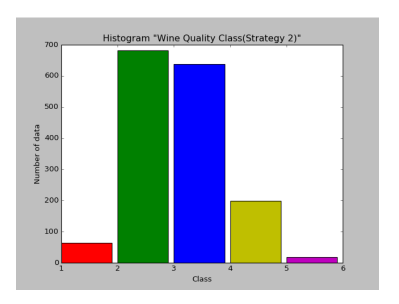


Figure 1.2: Labeling Strategy 2

1.2 Visualization

1.2.1 Principal Component Analysis

After normalization, the co-variance of matrix X can be estimated as:

$$\widehat{C_{j,k}} \approx \frac{1}{N} \sum_{i=1}^{N} (x_j^i - \mu_j) \cdot (x_k^i - \mu_k) = \frac{1}{N} \sum_{i=1}^{N} x_j^i \cdot x_k^i \implies \widehat{C} \approx \frac{1}{N} X X^T$$

If the covariance result is positive, that means that X,Y are positively correlated. The large the covariance result is, the dependency between X and Y will be.

After covariance result calculated, linalg.eig() function innumpy was called to generate the eigenvalues, eigenvectors of the co-variance matrix.

Once finish SVD decomposition, Cov[X] can be calculated as a matrix whose diagonal elements are exactly the eigenvalues of Cov[X].

Many of the features here are related to class tags, but there is noise or redundancy. In this case, a feature dimensionality reduction method is needed to reduce the number of features, reduce noise and redundancy, and reduce the possibility of over-fitting.

The idea of PCA is to map n-dimensional features to the k dimension (k < n), which is a new orthogonal feature. This k-dimensional feature is called the pivot element, and it is a reconstructed k-dimensional feature, rather than simply removing the k dimension features from the n-dimensional feature

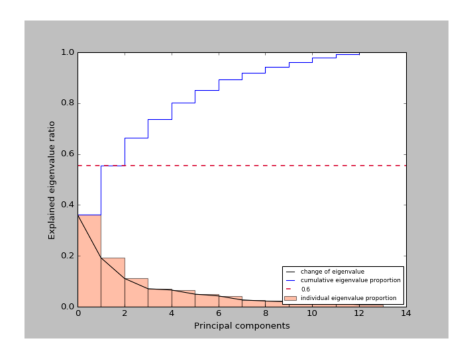
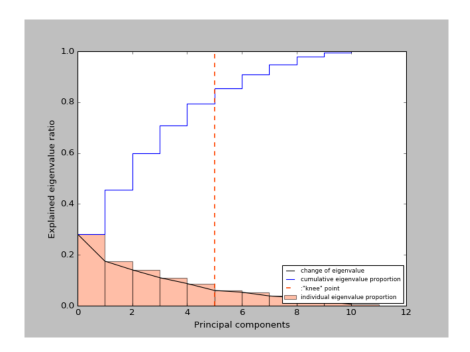


Figure 1.3.1: Cumulative Eigenvalues of Co-variance Matrix

Figure 1.3.2: Knee Points



The above **Figure 1.3.1** shows the cumulative eigenvalues of the co-variance matrix. It illustrates that if we just take two principal components base on this data set, it would not represent the original data set roughly. Hence, in the **Figure 1.3.1**, the knee point can be found according to the decrease ratio as the figure present.

The most important and useful information from PCA is the ranked imporance for the attributes. It means how many separable or variance information is allocated in a certain attribute from the dataset. In the following table, I list the value of the first 2 eigenvectors of the covariance matrix. The first column here shows coefficients of linear combination that defines principal component 1, and the second column shows coefficients for principal component 2.

If the value is positive, there is a positive correlation between the value and the principal component it project to. While, if it is negative, things will opposite the positive one. However, whatever the sign of a value is, I just care about how large the absolute value it will be. The large the absolute value is, the large effect it will do to principal component.

| Top 2 Largest Eigenvalues and Eigenvectors | | | | |
|--|-----------------------|-------------------------|----------------------|--|
| 1st largest: 3.09913244 | | 2nd largest: 1.92590969 | | |
| Dimensions | Values | Dimensions | Values | |
| fixed acidity | 0.48931421519678553 | density | 0.5694869591070468 | |
| citric acid | 0.4636316563339583 | total sulfur dioxide | -0.5236044991201001 | |
| sulphates | -0.43851962406530764 | volatile acidity | 0.3986719794517932 | |
| рН | 0.3953530087692868 | density | -0.37857017927744485 | |
| alcohol | 0.242921330946993 | 'chlorides' | 0.20275695801841312 | |
| volatile acidity | -0.23858436259606988 | citric acid | -0.1691026594058847 | |
| chlorides | 0.21224658194729165 | sulphates | -0.1637023809766136 | |
| residual sugar | 0.14610715358517834 | residual sugar | 0.10175002720609787 | |
| total sulfur dioxide | -0.11323206500149913 | рН | -0.07645796788428783 | |
| free sulfur dioxide | -0.036157524410518845 | free sulfur dioxide | 0.055788721251921254 | |
| density | 0.023574853564211597 | fixed acidity | 0.04560728999309162 | |

Table 1.3.1: Top two eigenvectors

Analysing the **Table 1.3.1: Top two eigenvectors** In new axis 1, **fixed acidity** index is the most important part because the coeffcient is **0.48931421519678553** which do largest contribution to the **1st principal component**. Similarly, **density** is the most important part of **2 nd principal component** which is **0.5694869591070468**. Combine with two column as the table display. We can only find that just **residual sugar** appear in the **last four value** both in 1st and 2 nd column. Thus, probably **residual sugar** is the most useless feature in all data set.

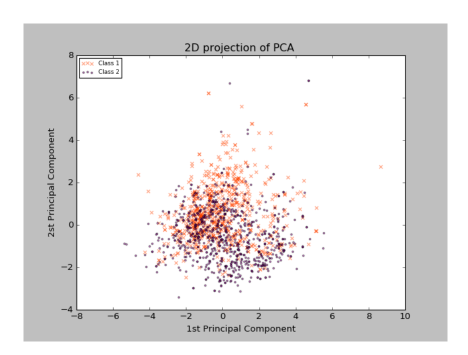


Figure 1.3.3: 2D PCA projection

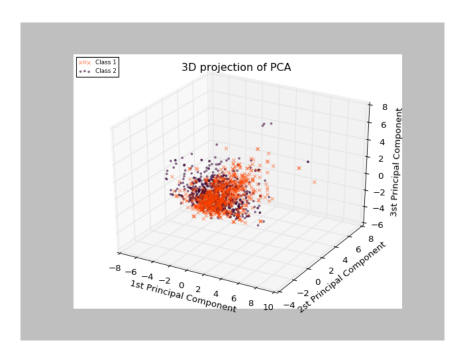


Figure 1.3.4: 3D PCA projection

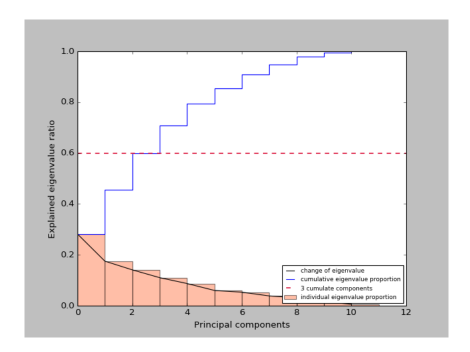


Figure 1.3.5: Ratio of 3 cumulative components

1.2.2 Self-organizing Map

To compare directly put original data set to som method and som after pca, I do the directly som firstly.

It is quite clear that the data be separated into two region, but actually we want to see the data can be analysis or observe in the lower dimension. Directly som seems to reduce the dimension, however it is a **fake**. With the check of normal vectors of the grids most of the cos value of the normal vectors are negative values or very small values, which means that the angle between each grid is pretty large, which confirm my judgement.

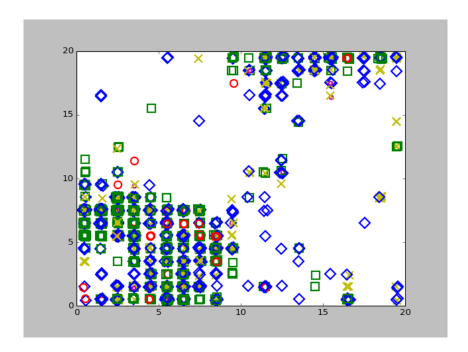


Figure 1.3.6: Direct SOM of 11 Dimensional data

Next, I used PCA to make the data reduce to **5 dimensions** with the use of label strategy 2. Here, the figure cover iteration 10times, 30 times, 50times.

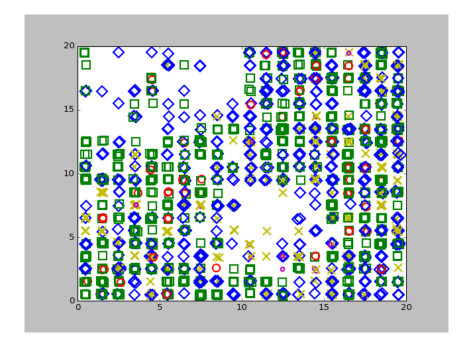


Figure 1.3.7: SOM (Iteration: 10 times)

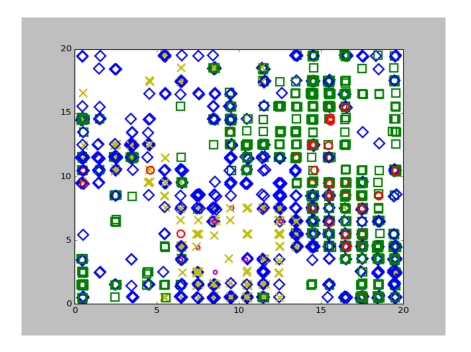
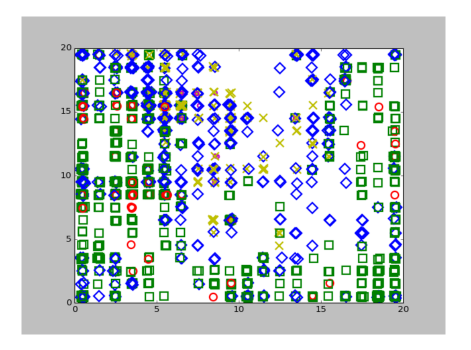


Figure 1.3.8: SOM(Iteration: 30 times)

Figure 1.3.9: SOM(Iteration: 50 times)



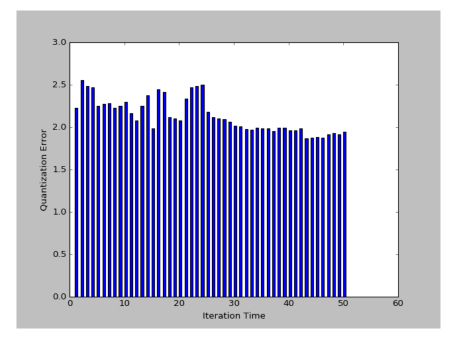


Figure 1.3.10: SOM Quantization Error)

In conclusion, SOM can reduce the high level dimension data set to lower level dimension which will make us easier to see the data feature in the current lower coordinate. Although in some coordinate there exist some small cluster be separated, we can find the link of nodes with same label dominate to whole graph. And we can simply find the the data be separated into mainly 3 clusters: yellow, green, blue.

1.2.3 The Comparision PCA and SOM

PCA and SOM are different technique to reduce the dimension of data. There exist some common and difference between them. My conclusion is as follow.

- When we focus a data set with **high dimension**. If we want to reduce the dimension, we **should not use SOM alone**. We should use PCA first to reduce the dimension as your wish, then do the SOM to make it easier to see the data feature in the current lower coordinate.
- Both of them can reduce the dimension of data. However SOM is more likely to use for compressing and stretching the data to suit the new coordinate. PCA can just combine the current related dimension of coordinate to form a new coordinate which is rather than a projection but a combination.

1.3 Clustering

According to the conclusion of SOM after PCA, we simply find that there exist 3 clusters in the data set.

In order to find the difference of clustering with and without PCA, here I made a contrast group as follow.

1.3.1 Clustering Without PCA

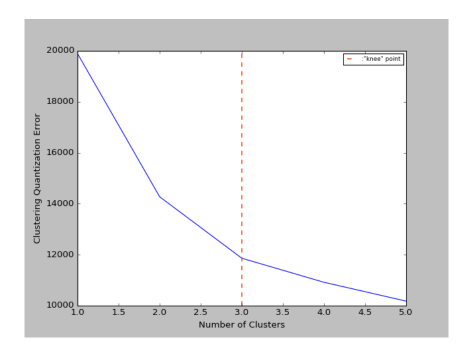


Figure 1.3.1:Clustering Quantization Error without PCA

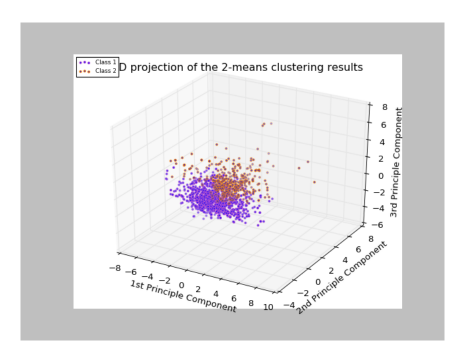


Figure 1.3.2: 2-Means Clustering

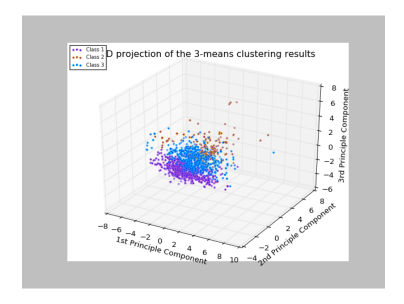


Figure 1.3.3: 3-Means Clustering

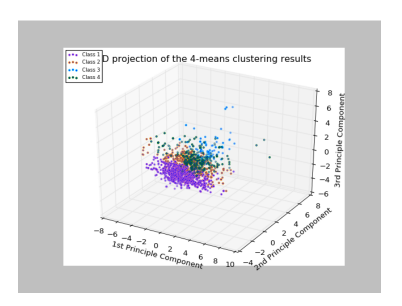


Figure 1.3.4: 4-Means Clustering

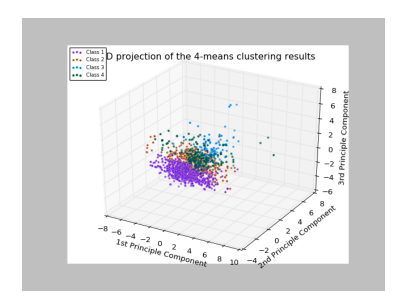


Figure 1.3.5: 5-Means Clustering

1.3.2 Conclusion

Without doing the pca, the data we observe in the 3D level always aim to cover themselves. So that we can obsever the data straight forward which is not a good method.

1.3.3 Clustering with PCA

We have see the result of clustering without pca. It seems not be a good method to obsever. Hence, I try to do PCA first to reduce the dimension of data, to see if it can improve the observation.

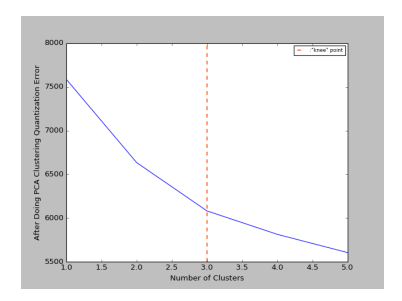


Figure 1.3.6: Clustering Quantization Error with PCA

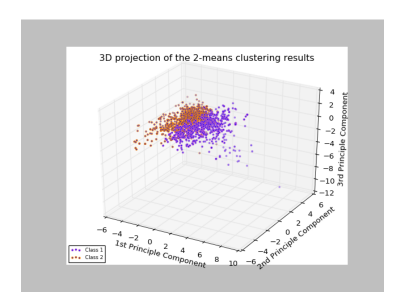


Figure 1.3.7: 2-Means Clustering

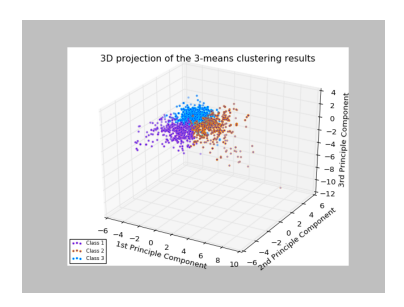


Figure 1.3.8: 3-Means Clustering

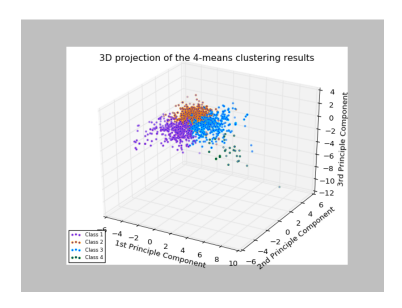


Figure 1.3.9: 4-Means Clustering

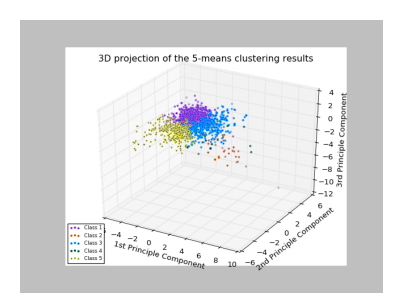


Figure 1.3.10: 5-Means Clustering

1.3.4 Conclusion

After doing the clustering with or without PCA, my conclusion as follow.

Comparing the knee point in Figure 1.4.1 and Figure 1.4.6. It is obvious that clustering with PCA reduce the quantization error. It might because of the reduction of dimension the PCA have done simultaneously reduce the noise. The knee point in graph of quantization error can not be a judgement of result of clustering. So we see the Figure 1.4.8 to estimate if this clustering method is suitable to this data set. We can find that the data set be separated unambiguous. Thus, k-means clustering is suitable for this data set.

Chapter 2

If there are any attribute that influence total sulfur dioxide?

2.1 Labelling

Strategy of labelling the total sulfur dioxide (0,160], (160 440]

2.2 Visualization

2.2.1 Principal Component Analysis

Just like what I did in Section 1.3, I used the same technique to complete this part. The result as follow. The first two feature affect to the data are fixed acidity and alcohol, which are 0.48809550387543194 and -0.5528903530300658

| Top 2 Largest Eigenvalues and Eigenvectors | | | | |
|--|-----------------------|-------------------------|----------------------|--|
| 1st largest: 3.12113839 | | 2nd largest: 2.11175471 | | |
| Dimensions | Values | Dimensions | Values | |
| fixed acidity | 0.48809550387543194 | alcohol | -0.5528903530300658 | |
| citric acid | 0.4734325262220812 | quality | -0.5236044991201001 | |
| pН | -0.4326046485510123 | volatile acidity | 0.3986719794517932 | |
| density | 0.3698915265676169 | density | -0.37857017927744485 | |
| volatile acidity | -0.2655231833551456 | 'chlorides' | 0.20275695801841312 | |
| sulphates | -0.2545312162478427 | citric acid | -0.1691026594058847 | |
| chlorides | 0.19715263821767132 | sulphates | -0.1637023809766136 | |
| residual sugar | 0.13860211721461613 | residual sugar | 0.10175002720609787 | |
| quality | 0.11318644768411748 | pН | -0.07645796788428783 | |
| alcohol | -0.07242073902021279 | free sulfur dioxide | 0.055788721251921254 | |
| free sulfur dioxide | -0.047260478851762794 | fixed acidity | 0.04560728999309162 | |

Table 2.3.1: Top two eigenvectors

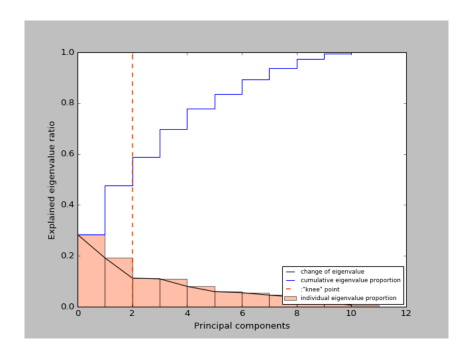


Figure 2.3.2: Cumulative eigenvalues of co-variance matrix

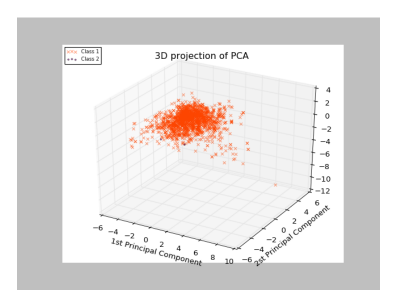


Figure 2.3.3: 3D PCA projection of total sulfur dioxide

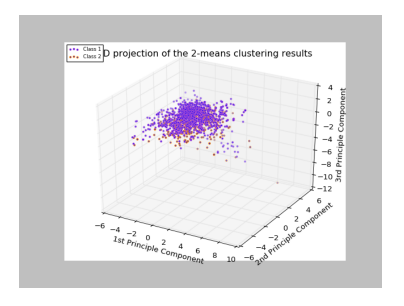


Figure 2.3.4: Clustering of total sulfur dioxide

2.3. Conclusion 32

2.3 Conclusion

As what I mention, The first two feature affect to the total sulfur dioxide are fixed acidity and alcohol, which are 0.48809550387543194 and -0.5528903530300658

Because of the limitation of working time and similarity of works, SOM and clustering were only present one figure.

However, I cover all the requirements.

If it is possible, I am looking forward to be your students in the future.

Finally, I sincerely express my thanks here to GuoJi and Peter.

Code & Related Files:

https://github.com/Voldet/data_analysis