# DAT565/DIT407 Assignment 5

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This paper is addressing the assignment 3 study queries within the *Introduction to Data Science & AI* course, DIT407 at the University of Gothenburg and DAT565 at Chalmers. The main source of information for this project is derived from the lectures and Skiena [1]. Assignment 5 is about distance and network methods.

#### Problem 1: Preprocessing the dataset

To preprocess the dataset, we first load the data from the file seeds.csv using the pandas library. We then normalize the data using the StandardScaler from sklearn.

# Problem 2: Determining the appropriate number of clusters

The normalized data is then used to calculate the inertia for different numbers of clusters. The inertia is plotted in Figure 1. Using the elbow method, we can see that the optimal number of clusters is 3.

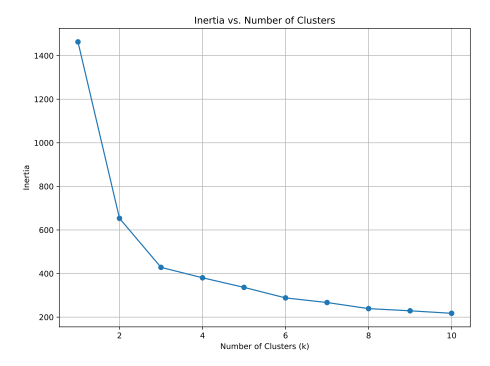


Figure 1: Invertia vs. Number of clusters

#### Problem 3: Visualizing the classes

To visually represent the classes, we initially project all the features onto a 2D plane. It becomes evident that the seeds form distinct clusters corresponding to the three species groups. However, certain features exhibit less distinct separation than others, such as asymmetry and compactness. Notably, the scatter plot depicting the relationship between seed length and width clearly illustrates that seed sizes are readily discernible. This is illustrated in Figure 2.

Subsequently, we employ Gaussian random projection to condense the data into a 2D space. The outcome is depicted in Figure 3, where the different seed species remain distinguishable. Finally, the UMAP projection is presented in Figure 4, highlighting the distinctiveness of the seed species.

The seeds are indeed distinguishable in the 2D space, particularly through size-related features (area, perimeter, length, width), which exhibit a linear relationship. Conversely, other features demonstrate less separability

The ability to distinguish seeds based on size-related features in a 2D space implies that clustering algorithms, particularly those sensitive to geometric relationships, can effectively separate seed samples into distinct groups. Algorithms like K-means or hierarchical clustering can leverage these separable features to identify natural clusters within the data. However, features that are less separable might pose challenges for clustering algorithms, potentially leading to overlapping clusters.

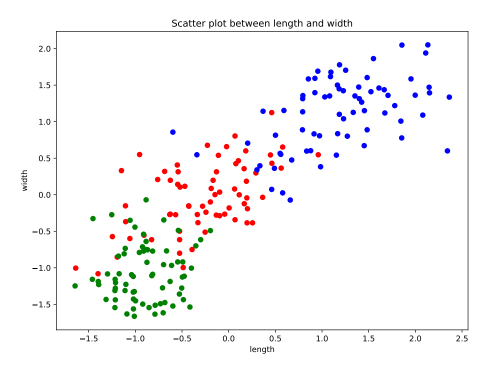


Figure 2: Scatter plot between lenght and width

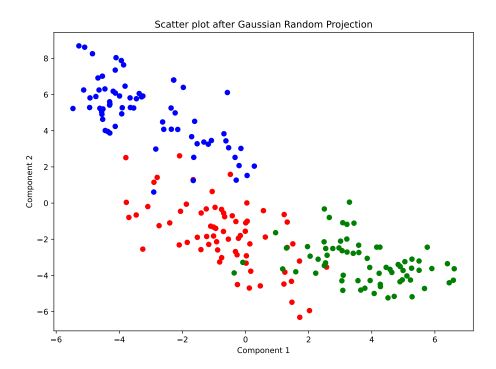


Figure 3: Gaussian random projection

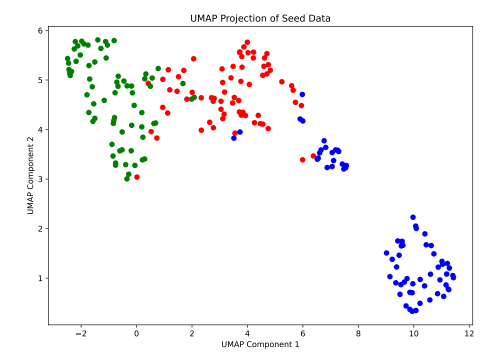


Figure 4: UMAP projection of Seeds

### Problem 4: Evaluating clustering

To apply k-means clustering to the data, we use the KMeans function from sklearn with 3 as the number of clusters, and then build the model on the normalized data.

The rand index is obtained by applying the rand\_score function on the labels of the clustering and the true labels. Its value is 0.90.

Finally we iterate over all the possible permutations in the range [0..4] to find the best accuracy score. With the permutation  $\{0,1,2,3\} \rightarrow \{2,3,1,0\}$ , the accuracy is equal to 0.92.

## Problem 5: Agglomerative clustering

We iterate over the linkage options and calculate the accuracy value after finding the right permutation for each of the linkage options. The best linkage option is the ward method, with an accuracy of 0.93. The dendrogram is shown in Figure 5

By looking at the 2-dimension projections from Problem 3, some of the points are close neighbors to points that don't belong to the same cluster, and the boundaries between clusters are not clearly defined. Therefore the "single" linkage option which merge clusters depending on the minimum distance gives a low accuracy value of 0.35. Other linkage options give roughly the same accuracy (around 0.9).

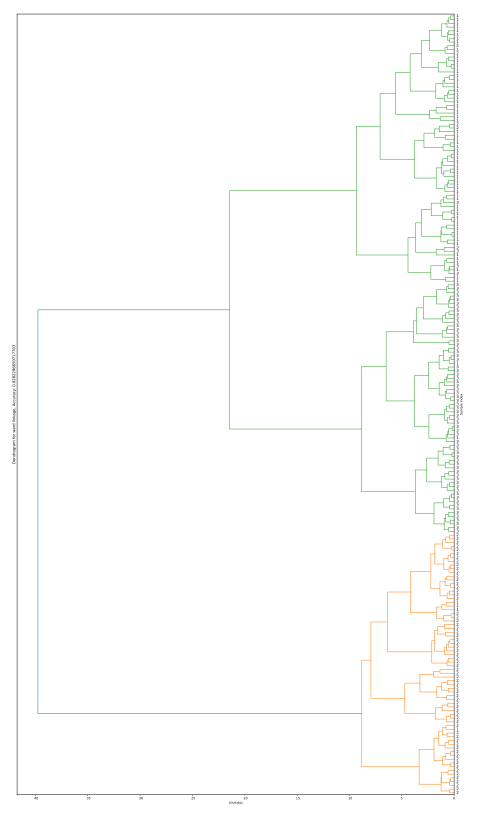


Figure 5: Dendrogram 6

#### References

[1] Steven S Skiena. The Data Science Design Manual. Retrieved 2024-01-20. 2024. URL: https://ebookcentral.proquest.com/lib/gu/detail.action?docID=6312797.

### Appendix: Source Code

```
from umap import UMAP
1
   import pandas as pd
3 import matplotlib.pyplot as plt
   {\bf from} \ \ {\bf sklearn.preprocessing} \ \ {\bf import} \ \ {\bf StandardScaler}
   from sklearn.cluster import KMeans
6 from sklearn.random_projection import GaussianRandomProjection
   from sklearn.metrics import rand_score
   import itertools
   from sklearn.metrics import accuracy_score
10
   from scipy.cluster.hierarchy import dendrogram, linkage
   from sklearn.cluster import AgglomerativeClustering
12
13 # Load the seeds dataset
14
   random_state = 79
   15
17
18 X = seeds.drop(columns=['species']) # Features
19
   y = seeds['species']
20
21 # Normalize the data
22
    scaler = StandardScaler()
23
    X_normalized = scaler.fit_transform(X)
24
25
    seeds_normalized = pd.DataFrame(X_normalized, columns=X.columns)
26
    seeds_normalized['species'] = y
27
28
   X = seeds_normalized.drop(columns=['species'])
29
30
    def plot_inertia(X):
        inertia_values = []
31
32
        for k in range (1, 11):
            kmeans = KMeans(n_clusters=k, random_state=random_state).
33
                \hookrightarrow fit (X)
            inertia_values.append(kmeans.inertia_)
34
35
        plt.plot(range(1, 11), inertia_values, marker='o')
        plt.xlabel('Number of Clusters (k)')
plt.ylabel('Inertia')
plt.title('Inertia's Number of Clusters')
37
38
39
        plt.grid(True)
40
41
        plt.show()
42
    def plot_features(features, y, colors):
43
        num_features = len(features)
44
45
        num\_rows = num\_features - 1
46
        num\_cols = num\_features - 1
47
48
        fig, axes = plt.subplots(num_rows, num_cols, figsize=(15, 15))
49
        for i in range(num_rows):
```

```
51
                for j in range(num_cols):
                     if i != j:
 52
                          ax = axes[i, j]

ax.scatter(X[features[i]], X[features[j]], c=y.map(
 53
54
                               ⇔ colors))
                          ax.set_xlabel(features[i])
55
 56
                          ax.set_ylabel(features[j])
57
                          ax.set_title(f'Scatter-plot-between-{features[i]}-

    and { features [ j ] } ')

 58
59
           plt.tight_layout()
60
           plt.show()
 61
     \begin{array}{lll} \textbf{def} & \texttt{plot\_gaussian\_random\_projection} \, (X, \ y, \ \texttt{colors} \,) \colon \\ \end{array}
62
63
           grp = GaussianRandomProjection(n_components=2, random_state=
                → random_state)
64
           projected = grp.fit_transform(X)
 65
          \begin{array}{ll} plt.\,fig\,ure\,(\,fig\,siz\,e\,=\,(8,\ 6)\,)\\ plt.\,scatter\,(\,projected\,[:\,,\ 0]\,,\ projected\,[:\,,\ 1]\,,\ c=\!y\,.\\ map(\,colors\,)\,)\\ plt.\,xlabel\,(\,\,'Component\,\cdot\,1\,\,') \end{array}
66
67
 68
           plt.ylabel('Component-2')
plt.title('Scatter-plot-after-Gaussian-Random-Projection')
 69
 70
 71
           plt.show()
 72
 73
     def plot_umap(X, y, colors):
           umap_model = UMAP(n_components=2)
 74
 75
           umap = umap_model.fit_transform(X)
 76
 77
           plt.figure(figsize=(8, 6))
 78
           plt.scatter(umap[:, 0], umap[:, 1], c=y.map(colors))
 79
           plt.xlabel('UMAP-Component-1
           plt.ylabel('UMAP-Component-2')
plt.title('UMAP-Projection-of-Seed-Data')
80
 81
           plt.show()
 82
83
 84
 85
 86
      def find-permutation(n_clusters, true_labels, cluster_labels):
87
           permutations = itertools.permutations(range(n_clusters))
88
           best_permutation = None
 89
           best_accuracy = 0
 90
           for permutation in permutations:
                {\tt permuted\_labels} = [\, {\tt permutation} \, [\, {\tt label} \, ] \  \, \begin{array}{c} \textbf{for} \\ \textbf{label} \end{array} \, \textbf{in}
91
                    92
                accuracy = accuracy_score(permuted_labels, true_labels)
93
                if accuracy > best_accuracy:
 94
                     best_accuracy = accuracy
95
                     best_permutation = permutation
96
           return best_permutation, best_accuracy
97
98
      def plot_dendrogram(n_clusters, X, y):
99
100
           linkage_options = ['ward', 'complete', 'average', 'single']
101
           best\_accuracy = 0
102
           best_linkage = None
103
104
           for linkage_option in linkage_options:
105
                clustering = AgglomerativeClustering(n_clusters=len(y.

    unique()), linkage=linkage_option)

106
                cluster = clustering.fit(X)
107
                permutation, accuracy = find_permutation(n_clusters, y,
```

```
⇔ cluster.labels_)
108
109
             if accuracy > best_accuracy:
110
                 best_accuracy = accuracy
111
                 best_linkage = linkage_option
112
         Z = linkage(X, method=best_linkage)
113
         plt.figure(figsize=(12, 6))
114
         dendrogram (Z, labels=y.values, leaf_rotation=90, leaf_font_size
115
             \hookrightarrow =8)
         plt.title(f"Dendrogram for {best_linkage}-linkage, Accuracy: {
116
        117
118
119
         plt.show()
120
121
    plot_inertia(X)
    colors = {1: 'red', 2: 'blue', 3: 'green'}
122
123
    features = seeds_normalized.columns
    \verb|plot_features| ( features , y, colors )
124
125
    plot_gaussian_random_projection(X, y, colors)
126
    plot_umap(X, y, colors)
127
128
129
    kmeans = KMeans(n_clusters=len(y.unique()), random_state=
        → random_state)
130
    kmeans. fit (X)
    kmeans_labels = kmeans.labels_
131
132
133
    rand_index = rand_score(y, kmeans_labels)
134
    print("Rand-score:", rand_index)
135
    all_labels = pd. Series (kmeans_labels)._append(y)
136
137
    all_unique_labels = all_labels.unique()
138
    best\_permutation, best\_accuracy = find\_permutation(len(
139

→ all_unique_labels), y, kmeans_labels)
140
    print("Best-Accuracy:", best_accuracy)
141
142
    print("Best - Permutation:", best_permutation)
143
    plot_dendrogram(len(all_unique_labels), X, y)
144
```