In the Name of ALLAH the Merciful the Compassionate

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EFFICIENT DATA CLUSTERING ALGORITHMS

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ABSTRACT

Data Clustering is one of the most important issues in data mining and machine learning. Clustering is a task of discovering homogenous groups of the studied objects. Recently, many researchers have a significant interest in developing clustering algorithms. The most problem in clustering is that we do not have prior information knowledge about the given dataset. Moreover, the choice of input parameters such as the number of clusters, number of nearest neighbors and other factors in these algorithms make the clustering more challengeable topic. Thus any incorrect choice of these parameters yields bad clustering results. Furthermore, these algorithms suffer from unsatisfactory accuracy when the dataset contains clusters with different complex shapes, densities, sizes, noise and outliers. In this thesis, we propose a new approach for unsupervised clustering task. Our approach consists of three phases of operations. In the first phase we use the most widely used clustering technique which is Kmeans algorithm for its simplicity and speed in practice. We benefit just from one run of Kmeans, despite its accuracy, to discover and analyze the given dataset by catching preliminary clusters to insure closely grouping sets. The second phase takes these initial groups for processing them in a parallel fashion using shrinking based on the convex hull of the initial groups. From the second phase we obtain a set of sub-clusters of the given dataset. Hence, the third phase considers these sub-clusters for merging process based on the Delaunay triangulation. This new algorithm is named as Kmeans-Based Convex Hull Triangulation clustering algorithm (KBCHT). We present experiments that provide the strength of our new algorithm in discovering clusters with different non-convex shapes, sizes, densities, noise and outliers even though the bad initial conditions used in its first phase. These experiments show the superiority of our proposed algorithm when comparing with most competing algorithms.

Keywords: data clustering, data mining, machine learning, homogenous groups, non-convex shapes, unsupervised clustering, Kmeans, shrinking, convex hull and triangulation.
تُعتبر عملية جمع البيانات واحدة من أهم الموضوعات دراسة في مجال تنقيب البيانات وتعلم الآلة و تفصّل عملية التجميع هو اكتشاف الجماعات النجاسة من الكائنات التي تشملها عينة الدراسة.

في الآونة الأخيرة، عملية تطوير خوارزميات التجميع جذبت اهتمام العديد من الباحثين حيث إن المشكلة الأساسية هي أنها لا تملك معرفة مستفزة عن مجموعة بيانات الدراسة المطلوبة خليلها و إيجاد الأتمات النجاسة فيها، و علَّمانا على ذلك، فإن معظم الخوارزميات الموجودة تتطلب من المستخدم إدخال معلومات حول مجموعة بيانات الدراسة مثل عدد الجماعات الموجود و عدد الخبراء الأقرب وغير من الدخلات التي تجعل من هذا الموضوع أكثر الموضوعات خطيرة للباحثين. وبالتالي فإن أي اختيار لدخل غير صحيح يعطي نتائج سلبية. و من ناحية أخرى، تعاين هذه الخوارزميات من نتائج غير مرضية عندما خُنُو مجموعة بيانات الدراسة على أشكال معقدة مختلفة في الكثافة و الحجم و بها ضوضاء و قيم متغيرة في هذه الأطروحة، عملنا على تطوير خوارزمية جديدة لجمع البيانات لا تضيع لرحلة الدخلات، و نهجنا على أن تكون هذه الخوارزمية على ثلاث مراحل: في المرحلة الأولى عملنا على Kmeans الاستخدام خوارزمية وهي مستخدمة على نطاق واسع لبساطتها و سرعتها تنفيذها و عمدادنا على الاستفادة من تشغيل واحدة لهذه الخوارزمية بعض النظر عن نتائجها، و ذلك لبداية اكتشاف و خليل مجموعة بيانات الدراسة من خلال هذه الجماعات الأولية، المرحلة الثانية نعنى بتحليل الجماعات الأولية بطريقة متوازنة باستخدام عملية الانكسام استنادًا على مبدأ الهيكل الغلب للجموعات الأولية. و كنتيجة للمرحلة الثانية فإنا تصل على مجموعة من الجماعات الأولية لبيانات الدراسة، و بالتالي فإن المرحلة الثالثة تعمد على مبدأ التنليث لتمجيع هذه الجماعات الصغيرة للحصول على النتيجة النهائية المطلوبة، و عمداً على تسميته هذه الخوارزمية معًا Kmeans-Based Convex Hull Triangulation clustering algorithm

(KBCHT)

و قد خلصت الدراسة بناءً على العديد من التجارب العملية التي أجريناها على فاعلية و قوة هذه الخوارزمية في اكتشاف الجماعات النجاسة من بين مجموعات بيانات الدراسة المختلفة التي خُوِي على أشكال معقدة مختلفة الاحجام والكثافة و خُوِي على ضوضاء و قيم متغيرة و ذلك بعض النظر عن الظروف الأولية التي تستخدم في المرحلة الأولى، و كما و إن هذه التجارب أثبتت نفوذ خوارزميتنا على أهم الخوارزميات اللافعة.
DEDICATION

TO WHOM IT MAY CONCERN
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Chapter 1

INTRODUCTION

A lot of data can be gathered from different fields but this data is useless without proper analysis to obtain useful information. In this thesis, we focus on one of the important techniques in data mining: Clustering.

1.1 Data Clustering

Data clustering is a method of grouping similar objects together. Thus the similar objects are clustered in the same group and dissimilar objects are clustered in different ones. An illustration example of clustering is shown in Fig 1.1. Data clustering is considered as an unsupervised learning technique in which objects are grouped in unknown predefined clusters. On the contrary, classification is a supervised learning in which objects are assigned to predefined classes (clusters).
Figure 1.1: Clustering example (a) dataset contains 14 objects. (b) Objects are grouped into 5 clusters. (adapted from: http://www.slideshare.net/pierluca.lanzi/machine-learning-and-data-mining-08-clustering-hierarchical last visit: July, 2011)

1.1.1 Basic Concepts of Clustering

The problem of data clustering can be formulated as follows: given a dataset $D$ that contains $n$ objects $x_1, x_2, ..., x_n$ (data points, records, instances, patterns, observations, items) and each data point is in a $d$-dimensional space, i.e. each
data point has \( d \) dimensions (attributes, features, variables, components). This can be expressed in a matrix format as:

\[
D = \begin{bmatrix}
    x_{11} & x_{12} & \cdots & x_{1d} \\
    x_{21} & x_{22} & \cdots & x_{2d} \\
    \vdots & \vdots & \ddots & \vdots \\
    x_{n1} & x_{n2} & \cdots & x_{nd}
\end{bmatrix}
\] (1.1)

Data clustering is based on the similarity or dissimilarity (distance) measures between data points. Hence, these measures make the cluster analysis meaningful [28]. The high quality of clustering is to obtain high intra-cluster similarity and low inter-cluster similarity as shown in Fig. 1.2. In addition, when we use the dissimilarity (distance) concept, the latter sentence becomes: the high quality of clustering is to obtain low intra-cluster dissimilarity and high inter-cluster dissimilarity.

![Figure 1.2: Inter-cluster and Intra-cluster similarities of clusters.](image)

### 1.2 Importance of Clustering

Data clustering is one of the main tasks of data mining [1] and pattern recognition [2]. Moreover, it can be used in many applications such as:

1. Data compression [3].
2. Image analysis [5].
3. Bioinformatics [6].
4. Academics [9].
5. Search engines [79].
6. Wireless sensor networks [80].
7. Intrusion detection [81].
8. Business planning [82].

1.3 Motivations

The Kmeans algorithm is considered as one of the top ten algorithms in data mining [35]. A lot of researches and studies have been proposed due to its simplicity and efficiency [55]. These efforts have focused on finding possible solutions to one or more of the limitations that have been identified in page 10. Kmeans with random initialization conditions need to be rerun many times each with different conditions to find more suitable results [21]. Many algorithms have been considered to provide better seeds so the Kmeans algorithm is likely to converge to the global optimum like Minmax[43], Kmeans++ [44] and [45]. Other solutions to the initial prototypes sensitivity can be found in [46] where they defined new criterion functions for Kmeans and they proposed three algorithms: weighted Kmeans, inverse weighted Kmeans [52] and inverse exponential Kmeans [53]. Other improvements of Kmeans focus on its efficiency where the complexity of Kmeans involves the data set size, number of dimensions, number of clusters and the number of iteration to be converged. There are many works to reduce the computational load and make it faster such as [4], [47-49]. Asgharbeygi and Maleki [39] proposed a new distance metric which is the geodesic distance to ensure resistance to outliers. Several works have been introduced to extend the use of means for numerical variables, thus Kmeans can deal with categorical variables such as [50], [51].

JJ Sheu et. al. [61] proposed a new algorithm and they named it Intelligent Kmeans (IKM) for deciding the proper number of clusters, choosing a better initial prototypes and reducing the effect of outliers on the clustering result. IKM divided the range of data points for each d dimensions into M regions where M is a constant input number. One drawbacks of this method, is the choice of grid size. If it is small, it will produce a large number of clusters and vice versa.
Many researchers have been involved in developing solutions to the Kmeans and other clustering algorithms such as using neighborhood model [23], ant colony [24], the principle of gravity [26], genetic algorithms [25], and clustering method with constraints [27].

The problem in clustering is that we do not have prior information knowledge about the given dataset. Moreover, the choice of input parameters such as the number of clusters, number of nearest neighbors and other factors in these algorithms make the clustering more challengeable topic. Thus any incorrect choice of these parameters yields bad clustering results. Furthermore, these algorithms suffer from unsatisfactory accuracy when the dataset contains clusters with different complex shapes, densities, sizes, noise and outliers.

In this thesis we want to design a novel clustering algorithm that is able to discover clusters with arbitrary complex shapes with presence of noise and outliers without requiring a previous knowledge of the given domain. In our approach we use the concept of convex hull [62] in which it is widely used in image processing to represent the shapes. Furthermore, it has been recently used in classification methods such as [64], [65]. Moreover, in [66] they provided a method of representing on-line data streaming using a cost function based on convex hull. In which they are concerned in representing the shape of data stream as a collection of convex hulls. However, this method cannot recover clusters correctly if the values of its input parameters are not set correctly.

1.4 Thesis outlines

In Chapter 2, we introduce a general overview of data clustering categorizations and algorithms. The ways of how we can construct a graph to be used in the clustering algorithms have been mentioned too. Moreover, we also explain and summarize some of the related works.

Our proposed algorithm is presented in details in Chapter 3. In which the three phases of the proposed algorithm are explained. A simple toy example is considered to be solved by our proposed algorithm. The simulation and results
analysis based on our generated complex shaped datasets have been accomplished in the last section of Chapter 3. Chapter 4 uses a topology preserving mapping as a preprocessing to our approach and we have used 10 real datasets from UCI machine repository to show the effectiveness of our proposed algorithm. Chapter 5 gives the conclusions and future research. We also provide two appendices that illustrate how the data preparation is done and the used distance metrics in the clustering algorithms.
Chapter 2

RELATED LITERATURE REVIEWS

2.1 Overview

The clustering problems can be categorized into two main types: fuzzy clustering and hard clustering. In fuzzy clustering, data points can belong to more than one cluster with probabilities between 0 and 1 [10], [11] which indicate the strength of the relationships between the data points and a particular cluster. One of the most popular fuzzy clustering algorithms is fuzzy c-mean algorithm [12], [13], [14]. In hard clustering, data points are divided into distinct clusters, where each data point can belong to one and only one cluster.

The hard clustering is divided into hierarchical and partitional algorithms. Hierarchical algorithms create nested relationships of clusters which can be represented as a tree structure called dendrogram [28]. Hierarchical algorithms can be divided into agglomerative and divisive hierarchical algorithms. The agglomerative hierarchical clustering starts with each data point in a single cluster. Then it repeats merging the similar pairs of clusters until all of the data points are in one cluster, such as complete linkage clustering [29] and single
linkage clustering [30]. CURE [15], ROCK [16], BIRCH [17] and Chameleon [18] are examples of this hierarchical algorithm. The divisive hierarchical algorithm reverses the operations of agglomerative clustering, it starts with all data points in one cluster and it repeats splitting large clusters into smaller ones until each data point belongs to a single cluster such as DIANA clustering algorithm [31].

In the contrary, Partitional clustering algorithm divides the dataset into a set of disjoint clusters such as Kmeans [32], [42] PAM [31] and CLARA [31]. Moreover, the partitional algorithms have been considered more appropriate for applications with large dataset, in which the construction of the dendrogram is computationally expensive [1], [37]. One of the problems in applying partitional methods is the choice of the number of clusters within the given datasets where the determination of the number of clusters is one of the most problematic issues in data clustering [7]. The partitional algorithms often use a certain objective function and produce the desired clusters by optimizing this objective function [36].

The clustering algorithms that are based on estimating the densities of data points are known as density-based methods. One of the basic density based clustering algorithm is DBSCAN [40]. It defines the density by counting the number of data points in a region specified by a predefined radius known as epsilon $\varepsilon$ around the data point. If a data point has a number greater than or equal to predefined minimum points known as $\text{MinPts}$, then this point is treated as a core point. Non-core data points that do not have a core data point within the predefined radius are treated as noise. Then the clusters are formed around the core data points and are defined as a set of density-connected data points that is maximal with respect to density reachability. DBSCAN may behave poorly due its weak definition of data points’ densities and its globally predefined parameters of $\varepsilon$ and $\text{MinPts}$. There are many works that try to improve the well known DBSCAN such as [41], [56-60].
2.2 Similarity Graphs

Another type of clustering algorithms is based on the construction of similarity graphs in which a given set of data points is transformed into vertices and edges. The constructed graph can be used to obtain a single highly connected graph that is then partitioned by edge cutting to obtain sub graphs [72], [74], [68]. Basically, the kinds of graphs are $\varepsilon$-neighborhood, $k$-nearest neighbor and fully connected graph [2], [70], [54].

The $\varepsilon$-neighborhood graph connects all data points whose pairwise distances are smaller than a predefined threshold $\varepsilon$.

In the $k$-nearest neighbor graph the data point $v_i$ (vertex) is connected with another data point in the dataset if it is in the $k$-nearest neighbors of $v_i$ where $k$ is a predefined parameter. This method lets the $k$-nearest neighbor produces a directed graph. The undirected graph can be obtained from the $k$-nearest neighbor by simply ignoring the directions of edges or by having a mutual $k$-nearest neighbor graph in which two vertices are connected by an edge if and only if these two vertices are among the $k$-nearest neighbors of each other.

The fully connected graph connects all data points that have a positive similarity measurement with each other. The similarity measure can be produced by using the Gaussian similarity function $S_{ij} = \exp(-d_{ij}^2/2\sigma^2)$ where $d_{ij}$ is the Euclidean distance between two data points $x_i$ and $x_j$ and the parameter $\sigma$ is also a user defined one that controls the width of neighborhoods.

2.3 Kmeans Algorithm

One of the most well-known unsupervised learning algorithms for clustering datasets is Kmeans algorithm [31], [37]. The Kmeans clustering is the most widely used due to its simplicity and efficiency in various fields [33], [38]. It is also considered as the top ten algorithms in data mining [35]. The Kmeans algorithm works as follows:

1. Select a set of initial $k$ prototypes or means throughout a dataset, where $k$ is a user-defined parameter that represents the number of clusters in the dataset.
2. Assign each data point in a dataset to its nearest prototype $m_i$.

3. Update each prototype according to the average of data points assigned to it.

4. Repeat step 2 and 3 until convergence.

The Kmeans algorithm depends on minimizing the sum of squared error function which is very simple and can be easily implemented.

$$J = \frac{1}{n} \sum_{i=1}^{k} \sum_{x \in C_i} \| x - m_i \|^2$$  \hspace{1cm} (2.1)

Where dataset $D$ contains $n$ data points $x_1, x_2, \ldots, x_n$ such that each data point is $d$ dimensional vector in $\mathbb{R}^d$, and $m_i$ is the prototype of cluster $C_i$, and $k$ is the given number of clusters.

However, it has several drawbacks: the number of clusters $k$ in a given dataset should be known in advance, the result strongly depends on the initial prototypes, the sensitivity to noise and outliers, the problem of dead prototypes or empty clusters and the converge to local optima [34]. The Kmeans works for globular shaped, similar size and density clusters.

### 2.4 CURE and Chameleon Algorithms

CURE [15] uses a constant number of well scattered representative data points from all data points in the dataset to represent a cluster instead of selecting one single centroid to represent a cluster in Kmeans. These are shrunk towards the centroid of the cluster according to a user predefined shrinking factor. Then a consecutive merging of the closest pair of the cluster’s representative points are occurred until the predefined number of clusters is obtained. The selection of the shrinking factor and the merging process make CURE ineffective with complex datasets and they can cause false outliers [22].

Chameleon [18] uses a graph construction based on k-nearest neighbors, and then it splits the graph into a set of small clusters using hMetis algorithm [19]. After that it merges these small clusters based on their similarity measure. It has been used to find non-convex shaped clusters, however, it cannot handle
noise and outliers and needs to set parameters correctly in order to obtain good results [22], [20].

2.5 Affinity Propagation Algorithm

Another type of clustering algorithms is called Affinity Propagation [67] that passes messages between data points to identify a set of exemplars (cluster centers) and their corresponding clusters. In contrary of selecting an initial set of cluster centers randomly and iteratively refines them such that the sum of squared error is minimized as in Kmeans; the Affinity Propagation provides a different approach that simultaneously considers all data points as candidate exemplars. Then two types of messages are exchanged between data points. The Responsibility messages are sent from data points to candidate exemplars and indicate how strongly each data point is biased to the candidate exemplar over other candidate exemplars. The Availability messages are sent from candidate exemplars to data points and reflect evidence that each candidate exemplar is available to be a cluster center of the data points. The Affinity Propagation uses the median of similarities between data points as preferences rather than the predetermined number of clusters.

2.6 Spectral Clustering Algorithm

Recently, the spectral clustering [70] has become one of the most popular clustering algorithms which outperform the traditional algorithms such as Kmeans. Furthermore, they are designed to handle non-convex shaped clusters. However, spectral clustering suffers from heavily computations. The similarity measure and graph cutting are also used in spectral clustering algorithms. The core of the spectral clustering algorithms is to use the properties of eigenvectors of Laplacian matrix for performing graph partitioning [69-76].

The Laplacian matrix is constructed by building an affinity graph matrix with a similarity measure. The common similarity measure is to use the Gaussian function $S_{ij}$ as stated previously for its simplicity. Hence, the Laplacian matrix $L$ is calculated as $L=D-S$ where $D$ is the diagonal matrix.
whose elements are the sum of all row elements of $S$. Then, the spectral clustering computes a column matrix of the first $k$ eigenvectors of $L$ where $k$ is a predefined number of clusters. Thus it finds the clusters of mapped data points that corresponding to the column matrix of eigenvectors by performing Kmeans algorithm.

**2.6.1 Spectral Clustering using Nystrom Method**

W.-Y. Chen et. al. [76] proposed sparsification and Nystrom approaches to address the computational difficulties and to improve the results. We compare our algorithm with spectral clustering using Nystrom method because it needs less computation and does not need the prespecified number of nearest neighbors as in sparsification method. Nystrom method is a technique for finding an approximate eigendecomposition. The spectral clustering using Nystrom method uses randomly sample data points from the dataset to approximate the similarity matrix of all data points in the dataset. Then it finds the first $k$ eigenvectors of the normalized Laplacian matrix of the Nystrom method and performs Kmeans to cluster dataset.

**2.7 Topology Preserving Mapping**

A topographic mapping is a transformation of high dimensional data. Furthermore, it preserves some structure in the data such as the points which are mapped close to each other share some common properties while in contrast the points which are mapped far from each other do not share a common feature or property.

The Self-organizing map (SOM) [84] and the Generative topographic mapping (GTM) [85] have been considered as very popular topology preserving mapping techniques for data visualization and dimensionality reduction. The GTM can be considered as a statistical alternative to the SOM overcoming many of its limitations such as the absence of a cost function and the lack of proof convergence [86].
2.7.1 Self-organizing Map (SOM)

The Self-organizing Map (SOM) [84] is a type of artificial neural network that is trained using unsupervised learning. SOM reduces dimensions of the given datasets by producing a map of usually one or two dimensions. Furthermore, SOM uses a neighborhood function to preserve the topological properties of the input space.

The SOM consists of components called nodes or neurons in which they are usually arranged in a hexagonal or rectangular grid. It first initializes the weights associated with each neuron by assigning them small random values. Then the SOM proceeds to three essential processes: competition, cooperation, and adaptation [28].

2.7.2 Generative Topographic Mapping (GTM)

The GTM is a statistical model for modeling the probability density of data points and finding non-linear mapping of high dimensional space onto low dimensional space.

![Non-linear mapping by GTM](image)

**Figure 2.1:** Non-linear mapping by GTM, adapted from [87].

As shown in Fig. 2.1, the basis of the GTM is to generate a grid of $K$ latent points $z_1, z_2, \ldots, z_K$ in latent space. These latent points are mapped non-linearly into the data space, which contains $N$ data points $x_n (n=1,2,\ldots,N)$, using a set of $M$ fixed basis Gaussian functions, such that,
\[ y_k = \Phi_k^{Tr}W \]  
(2.2)

Where \( y_k \) denotes the mapped points in data space. The element \( \Phi \) consists of \( M \) fixed basis functions. \( W \) is \( M \times D \) matrix containing weight parameters, and \( D \) is the dimensionality of data space. \( A^{Tr} \) is a transpose of a matrix \( A \).

The probability density between the mapped points \( y_k \) and data points \( x_n \) is estimated by using a Gaussian noise distribution centered on \( y_k \) with the noise inverse variance \( \beta \). This probability density \( p(x_n|y_k,W,\beta) \) is defined as:

\[ p(x_n|y_k,W,\beta) = \left( \frac{\beta}{2\pi} \right)^{D/2} \exp \left( -\frac{\beta}{2} \| x_n - y_k \| ^2 \right) \]  
(2.3)

The training step of GTM is done by optimizing its parameters using the Expectation-Maximization (EM) algorithm [2], [28] which maximizes the following log-likelihood:

\[ \mathcal{L}(W,\beta) = \arg\max_{W,\beta} \sum_{n=1}^{N} \ln \left( \frac{1}{K} \sum_{k=1}^{K} p(x_n|y_k,W,\beta) \right) \]  
(2.4)

After convergence, we can visualize the data by projecting each data point \( x_i \) onto the latent space using one of the two ways:

- The Mode: the mode of posterior distribution:
  \[ y_k_{\text{mode}} = \arg\max_{y_k} p(y_k|x_i) \]  
(2.5)

- The Mean: the mean of posterior distribution:
  \[ y_k_{\text{mean}} = \sum_{k=1}^{K} y_k p(y_k|x_i) \]  
(2.6)

Where \( p(y_k|x) \) is the corresponding posterior distribution in the latent space for any given data point \( x \) in the data space and is defined as:

\[ p(y_k|x) = \frac{p(x|y_k,W,\beta)p(y_k)}{\sum_{k'=1}^{K} p(x|y_{k'},W,\beta)p(y_{k'})} \]  
(2.7)
KBCHT: KMEANS-BASED CONVEX HULL TRIANGULATION CLUSTERING ALGORITHM

The problem of clustering datasets is that we have no prior knowledge information about them. Thus the majority of existing clustering algorithms try to solve it by introducing external input parameters which make these works sensitive to their inputs. In this chapter we introduce Kmeans-Based Convex Hull Triangulation clustering algorithm (KBCHT) a new clustering algorithm that studies the given dataset to find the clusters. KBCHT algorithm is able to detect clusters without pre-determination of clusters number in datasets which contain complex non-convex shapes, different sizes, densities, noise and outliers. Algorithm 3.1 provides a pseudo-code that describes the overall procedures of KBCHT algorithm.

3.1 Proposed KBCHT Algorithm

KBCHT has three phases of operations. The first phase obtains initial groups from running Kmeans algorithm just once, the second phase analyzes these initial groups to get sub-clusters and the last one merges the sub-clusters to find the final clusters in the dataset. As shown in Algorithm 3.1, KBCHT performs Kmeans algorithm on the dataset $\mathbf{x}$ given the number of clusters $k$. The use of $k$ is just to run Kmeans as we will notice by further study of the effect of $k$ in
Section 3.3.2. Line 2 means that the first run of Kmeans algorithm despite its bad initialization conditions has an initial set of clusters \( iC_i \) where \( i \) is from 1 to \( N \) the number of obtained clusters from Kmeans. The set \( iC \) with index \( i \) contains data points from dataset \( x \) which belong to the initial cluster \( i \). Lines 3 to 7 describe the process of how we analyze these initial clusters to obtain a set of sub-clusters. In line 4, we construct a set of vertices which represents each initial clusters \( iC \). This set of vertices is obtained from the convex hull of each initial clusters \( iC \). The set \( IV \) handles these vertices which contains two indexes \( i \) and \( j \) as shown in line 4. In which the index \( i \) indicates that these vertices belong to the initial cluster \( i \) and the index \( j \) represents the vertex number of convex hull of initial cluster \( i \) in a counterclockwise order. After obtaining the vertices from the convex hull, these vertices need to be shrunk by adding new vertices from the belonged initial clusters set \( iC \). Thus we begin with vertices drawn from a convex hull and finish with vertices of a polygon. The shrunk vertices are handled in the set \( sV \) as shown in line 5 of Algorithm 3.1. Line 6 takes the shrunk vertices \( sV \) and processes them to obtain a set of sub-clusters \( sC \), the number of these sub-clusters \( S \) and the average distance between data points of each of the sub-clusters in \( sC \) (sCaD) using the delaunay triangulation [63] as will be explained later. The sub-clusters are formed by searching for closed loops vertices in the \( sV \) set. The set \( sC \) has indexed from 1 to \( S \) in which \( sC_i \) contains data points of dataset \( x \) that belong to sub-cluster \( i \). Some of these sub-clusters could be merged together to form the final result of clusters \( C \) as shown in line 8.

**Algorithm 3.1: KBCHT**

```
1   Begin initialize k,x,iC = {},sC = {},C = {},iV = {},sV = {},sCaD = {},N = 0 , S = 0
2      iC,N <-- Kmeans(x,k)
3      for i = 1 to N
4          iV(i,j) <-- Construct convexhull for cluster iCi
5          sV(i,k) <-- shrinkVertex(iCi,iV(i,:))
6          sC,S,sCaD <-- findSubClusters(iCi,sV(i,:))
7      end_for
8      C <-- merging(sC,S,sCaD)
9      Return C
10  end
```
3.1.1 The First Phase: The use of standard Kmeans

KBCHT algorithm chooses to use the well known Kmeans algorithm as it is first step because of its simplicity and widely use as one of the top ten data mining algorithms [35].

The aim of our algorithm is to find clusters of arbitrary shapes and to detect odd patterns that exist in the datasets which Kmeans is far away from detecting them; Kmeans depends on assigning data points to their nearest mean thus the final result of it comes out as spherical shapes. Thus we can benefit from its first run with randomly thrown prototypes throughout the given dataset to catch preliminary initial clusters that insures closely grouping sets. However, Kmeans algorithm needs to be injected with $k$ the number of clusters in the dataset. Further investigation on the effect of $k$ has been conducted in Section 3.3.2. The $k$ or less than $k$, in case of dead prototypes, resultant partitions from Kmeans could be processed and analyzed in a parallel fashion which speeds up the processing time.

In this phase we are concerned with catching initial relatively related groups and Kmeans algorithm gives relatively robust and good enough answers over a wide variety of datasets as mentioned in [46]. Hence, we have decided to work on the standard Kmeans algorithm.

Generally speaking, we can use any other method in which they offer grouping such as any of the partitional clustering algorithms that are mentioned in Chapter 1. Moreover, we can benefit from the researches that are focused on the construction of similarity graphs of the given datasets as explained in Chapter 2.

Besides that, there are many developed researches related to overcome the limitations of the standard Kmeans algorithms. Hence, the choice of one of these developed researches to be as our first phase depends on what we want and what we have. i.e, we want more accurate initial result despite the time it could take or vice versa.

Some of these researches have aimed at identifying the initial centroids locations. In [91] they avoided the initial random assignment of centroids using
sub-merger strategy and [46] focused on the sensitivity to initial centroids condition. Another research has focused on the number $k$ of clusters such as in [92] where they proposed a measure to select the number of clusters. Others have tried to accelerate the Kmeans algorithm by avoiding many calculations of distance based on partial distance strategy like in [4].

3.1.2 The Second Phase: The Shrinking

After catching the initial groups, we want the greatest benefit from them. How we can analyze and represent them? We can go back to the topics of grid clustering in the book [28] and the research as in [62] in which they divided the group of data points into equal grids then trying to eliminate the grids that do not contain sufficient number of points. But by using this, we have stuck under the mercy of the user defined parameters. Hence, we have decided to use widely used concept of representing shapes as in image processing which is the convex hull mechanism.

As shown in lines 3 to 7 of Algorithm 3.1, this phase of KBCHT algorithm operates on the set of initial clusters that are obtained from the first phase. Each group of initial clusters is represented by its surrounding vertices on convex hull and the data points inside this convex hull. Then these vertices of each group are shrunk separately until we find the final sub-clusters. Procedure 3.1 in the next page describes the shrinking process in details.

Suppose we have one of the initial clusters obtained from the first phase of KBCHT algorithm as shown in Fig. 3.1 (page 18). The blue ‘*’s in Fig. 3.1 represent the data points of the given dataset which can belong to one or different final clusters of the dataset. The solid lines that are surrounding the data points in Fig. 3.1 represent the convex hull of this portion of dataset. The vertices from $V_1$, $V_2$ to $V_{12}$ are the boundary data points drawn from the convex hull in which $V_1$ equals to the last vertex $V_{12}$ and these vertices are in a counterclockwise order. As in Fig. 3.1 the blue ‘*’s are $x$ data points in Procedure 3.1 and the set $Vs$ is the vertices of convex hull. In case of Fig. 3.1, $Vs$ is from $V_1$ to $V_{12}$. 
Procedure 3.1: Shrink Vertices

Input:
Vs: Vertices of convex hull
x: data points inside convex hull of Vs

Begin
MAX ← find maximum length of edges between Vs
AVG ← find average distance among objects x inclosed by Vs
while MAX < AVG
    nVs ← exclude Vs from x and reconstruct convex hull
    MAX ← find maximum length of edges between nVs
    AVG ← find average distance among objects x inclosed by nVs
    V ← Reorder Vs starting from vertex belongs to the longest edge
        (reserve counterclockwise order)
end_while
while MAX ≥ AVG OR Converged
    P ← find closest point from x to the line between V₁ and V₂
        and its projection falls between V₁ and V₂
        and resides on the left of V₁ and V₂
        and the perpendicular line from P to the line between V₁ and V₂
        does have no intersection with other edges between vertices
    if such a point P exists
        add P to V between V₁ and V₂
    else
        flag V₁ and V₂ as processed vertices
end_if
MAX ← find length of longest edge and its vertices are not processed
Reorder V starting from vertex belongs to the longest edge and its
vertices are not processed (reserve counterclockwise order)
end_while
Return V

Line 5 in Procedure 3.1 computes the maximum length of edges between
each two consecutive vertices and stores it in MAX variable. KBCHT algorithm
does not use external parameter to guide the shrinking processing. It self
studies the given data points to decide when and how to make the shrink. This
maximum edge has the highest priority to be shrunk.

We can make the shrinking based on different criteria such as defining an
external parameter to be a threshold, instead of calculating our average, like in
many clustering algorithms that use threshold parameters. Thus by starting
shrinking the maximum edge length until the maximum one becomes less than
the defined threshold. But as stated before, KBCHT algorithm does not like to be with the clemency of external factors. Hence, we decide to compute the average distance among data points that enclosed by the set of $V_s$. The vertices $V_s$ are excluded from computing this average to eliminate the effect of being outliers. To compute the average distance among data points, we do not want to consider the distances between each data point and every other data point in the set; This will be computational expensive and does not reflect the actual data structure of the given set of data points. Thus we construct the Delaunay triangulation of the data points. Then the average distance $AVG$ is the average length of the triangles edges.

Now we have two variables $MAX$ and $AVG$. However, for starting the shrinking process the maximum edge length of convex hull $MAX$ should be greater than the average distance $AVG$. If it is not, this means that the vertices of the convex hull are denser than the enclosed set of data points as shown in Fig. 3.2. In this case, we identify a new set of vertices by reconstructing the convex hull of the data points again without considering the previously obtained vertices. These are shown in lines 7 to 12 of Procedure 3.1.
After satisfying the above condition for starting the process of shrinking, we reorder the vertices to begin from the vertex that belongs to the longest edge length (line 13 of Procedure 3.1). But we have to reserve the order of vertices in a counterclockwise order. Back to Fig. 3.1, the longest edge length is the edge between two vertices $V_9$ and $V_{10}$. So we reorder the vertices such that $V_1$ becomes $V_9$ and $V_2$ becomes $V_{10}$ and so on.

**How to do the shrinking:** At this stage, we have a set of vertices begins from the vertex with the longest edge. We want to find another vertex from the data points to be engaged between the first two vertices. To find this new vertex, we have to find the closest data point to the line between the first two

---

**Figure 3.2:** (a) Example of a set of data points, red ‘*’s, that vertices of the convex hull, black solid lines, are far away from a sparse cluster. (b) Shrinking result after eliminating the vertices of convex hull. (c) The final result red ‘*’s are in one cluster and black ‘+’s are in a separated cluster.
vertices of the set \( V \) and its projection lies on the line between these two vertices. Consider Fig. 3.3 as an illustrated example for finding the closest point to the line. In Fig. 3.3 we want to find the closest point to the line AB. To do this we have to find the projection points (D1 and E1) on the line AB from the tested points. Let \( A=(A_x,A_y) \), \( B=(B_x,B_y) \) and \( D=(D_x,D_y) \). Our goal is to find the point \( D1=(D1_x,D1_y) \). Hence \( D1 \) lies on the line AB, thus it satisfies its line equation and it can be found from: \( D1=A+u(B-A) \). To find \( u \), the dot product of the perpendicular line from \( D \) to line AB and line AB is 0. Thus \( u \) can be found from the following equation [89]:

\[
\begin{align*}
 u &= \frac{(D1_x - A_x)(B_x - A_x) + (D1_y - A_y)(B_y - A_y)}{(B_x - A_x)^2 + (B_y - A_y)^2} \\
 &= \frac{3}{12}
\end{align*}
\]  

(3.1)

To guarantee that \( D1 \) lies between \( A \) and \( B \); the value of \( u \) should be between [0, 1]. So, the points C and F do not be considered. Then the distance from the point to the line is the distance from that point to its projection point on the line.

We reserve the order of the vertices to be counterclockwise. Thus the picked point that has to be a new vertex should also reside on the left side of the two vertices in which the new vertex has to be engaged between them. To ensure this, Back to Fig. 3.3 suppose we have two vertices A and B and we want to examine that the data point D resides on the left side of the direction from A to B. we compute the value of the following equation [90]:

![Figure 3.3: Solid line between points A and B. Dash lines are the projection of points to the line AB.](image)
value = \begin{vmatrix} A_x & B_x & D_x \\ A_y & B_y & D_y \\ 1 & 1 & 1 \end{vmatrix} \quad (3.2)

Where |.| is the matrix determinant. If the sign of the value of equation (3.2) is positive then the examined point is on the left side. As well as it is on the right side if the sign is negative and on the straight line if it has a zero value. In fact, the equation (3.2) reflects the area of the triangle of ABD.

However, we have started the shrinking with the vertices that lay on the convex hull. Our approach in shrinking does not conserve the convexity of the shrunk vertices shape. Thus we have to provide an additional condition for shrinking which is shown in line 15 of Procedure 3.1. This condition says that the perpendicular line from the candidate vertex to the line between the two vertices in which their edge has to be shrunk should not intersect any of the lines that are between two consecutive vertices. This condition has exceptions. To be more obvious and as stated previously, the candidate vertex should be between \( V_1 \) and \( V_2 \) vertices thus if the candidate vertex is already an existing vertex, we violate the previous condition if this candidate vertex is \( V_3 \) or \( V_{last-1} \) (vertex that resides before the last one directly). Also, it should be violated if there is only one intersection and this intersection point is equal to the vertex \( V_1 \) or \( V_2 \). While the process of shrinking is going on, some vertices have to be released from the vertices set if a specific vertex has the same previous and next vertex. Hence, this vertex has to be released if its distance to next vertex is greater than the \( AVG \) value. The released vertices should be processed in the last phase of KBCHT algorithm. The sharp eyed readers may notice that the above condition of releasing vertices may be violated even though they should be released. This situation happens when we want to release a vertex that resides far away from a group of dense data points but its distance to next vertex is less than the overall average distance \( AVG \). In this case, we add one more examine condition in which we compute the length of the two adjacent edges of this vertex. If both of them are greater than the overall average distance \( AVG \) and they have no close candidate vertices to be added in the set of \( V \). Thus we guarantee to release this vertex from the set \( V \).
Line 17 of procedure 3.1 says: “All of the above mentioned conditions have been satisfied”. Thus we add the candidate vertex to be one of the vertices in the set $V$ between vertex $V_1$ and $V_2$. If no such a candidate vertex in which it violates the above conditions, the two vertices $V_1$ and $V_2$ have been flagged to be processed. A new value of variable $MAX$ should be recalculated as in line 21 of Procedure 3.1. But at this time the vertices that have a maximum edge length should not be flagged as processed. If so, find the next longest edge length until their vertices are not flagged. Then the set $V$ is reordered again starting from the vertex that belongs to the longest edge length and their vertices are not flagged as processed. We repeat this process of shrinking again and again until the $MAX$ value becomes less than the average distance $AVG$ or all vertices in the set $V$ have been flagged as processed.

**What to do with the shrunk vertices:** the last operation of the second phase of KBCHT algorithm is to refine the shrunk vertices and to extract useful information from the set of these vertices by finding the groups of sub-clusters entire the given dataset. Procedure 3.2 says how to find sub-clusters. After shrinking the vertices that obtained from the first phase of KBCHT algorithm in which they reside on the boundary of the convex hull of each of the resultant initial groups from the first phase. At this stage, the shrunk vertices are on the boundary of initial group of data points where this boundary forms a general polygon shape.

KBCHT algorithm calls the Procedure 3.2 in its line 6 as shown in Algorithm 3.1. Procedure 3.2 takes two inputs: $V$ which is the resultant shrunk vertices and $x$ which is the set of data points that enclosed by $V$. This $x$ is a part of the overall given dataset. We want to distinguish these vertices in the set $V$ thus each group of connected vertices belongs to a separate sub-cluster. We define the set $L$ that has a number of entries equal to the number of the given vertices in $V$. Initially, all the vertices are assigned a label value of 0 such that $L_i=0 \forall i \in \{1,\ldots,len\}$ where $len$ is the number of shrunk vertices. For example if $L_5$ has a label value equals to 1, then this means that the $V_5$ of the set $V$ is
assigned to the sub-cluster of label 1. Thus \( L \) handles the labels of sub-clusters that each of vertices in \( V \) belongs to.

In line 9 of Procedure 3.2 we define a variable \( \text{count} \) which it is initially set to be 1 and is incremented automatically by one when the algorithm detects a new sub-cluster. So, \( \text{count} \) represents the labels of sub-clusters that the vertices belong to. Procedure 3.2 enters a loop that begins from line 10 and ends up at line 22. These lines are the core of this Procedure in which we define two pointers \( i \) and \( j \) such that we fix \( i \) to point to a specific vertex in the set \( V \) and then the pointer \( j \) investigates each of the following vertices after that vertex which is pointed by \( i \). When the pointer \( j \) reaches the last vertex in the set \( V \), we move the pointer \( i \) one step forward to point to the next vertex and the pointer \( j \)
restarts and repeats its mission. This task will be repeated until \( i \) reaches the \( \text{len-1} \) vertex which is one position before the last vertex. Line 11 tests if the corresponded vertex has never assigned to sub-cluster yet. If so, the procedure enters into the loop of the pointer \( j \). In line 13 of Procedure 3.2 we check if the vertex pointed by \( i \) (\( V_i \)) and the vertex pointed by \( j \) (\( V_j \)) are the same. If they are identical vertices, then we obtained a closed connection of vertices which also means that we catch one of the sub-clusters in the dataset. Line 16 gives the sub-cluster’s label to the set \( L \) starting from its index of \( i \) through \( j \). After that we increment the label identified by \( \text{count} \) by 1. This process will be repeated until we catch all of the sub-clusters in the given dataset. Before line 23 of Procedure 3.2, we have groups of vertices each belongs to different sub-clusters; these groups represent the boundary of the data points of each sub-cluster. Thus the lines from 23 to 25 find the data points themselves in which they are enclosed by each of the obtained closed connection of vertices. Hence, the set \( sC \) in Procedure 3.2 handles the data points of each sub-cluster in the dataset and it has an index starts from 1 to the number of obtained sub-clusters where each index handles the tightly data points that belong to a specific sub-cluster. Finally, for each of the obtained sub-clusters we compute the average distance between data points within each of them as explained in Section 3.1.2 and store these average distances in the group of \( sCaD \) as shown in lines 26 to 28.

### 3.1.3 The Third Phase: Merging sub-clusters

Procedure 3.3 shows the steps of the last phase of our KBCHT algorithm which is merging the obtained sub-clusters for finding the final set of clusters in the studied dataset. The set \( sC \) is defined to be a group of sub-clusters of the dataset. In addition, \( S \) is the number of these sub-clusters. Hence, \( sC=\{sC_1,sC_2,\ldots,sC_S\} \) in which each of the sub-cluster \( sC_i \) contains the data points that belong to this sub-cluster. The \( sCaD=\{sCaD_1,sCaD_2,\ldots,sCaD_S\} \) group handles the average distance within each of the sub-clusters thus \( sCaD_i \)
Procedure 3.3: Merging sub-clusters

1. **Input:**
   1. \( sC \): group of sub-clusters
   2. \( S \): Number of sub-clusters
   3. \( sCaD \): group of calculated average distances within each of the sub-clusters

2. **Begin**
   1. \( C_1 \leftarrow sC_1 \)
   2. Remove entries of label 1 from \( sC \) group
   3. \( aD_1 \leftarrow sCaD_1 \)
   4. \( i \leftarrow 1 \)
   5. \( j \leftarrow 2 \)
   6. **while** \( sC \) group is not empty
      1. **while** \( j \leq S \)
         1. **if** \( sC \) does not contain label \( j \)
            1. \( j \leftarrow j + 1 \)
            2. **continue**
         2. **end_if**
         3. \( db2c \leftarrow \text{compute the distance between two sub-clusters } C_i \text{ and } sC_j \)
         4. **if** \( db2c < aD_i \)
            1. **add** \( sC_i \) **to** \( C_i \)
            2. Remove entries of label \( j \) from \( sC \) group
            3. **if** \( sC \) is empty
               1. **break**
            4. **end_if**
            5. \( aD_i \leftarrow \text{recompute average distance among objects in sub-cluster } C_i \)
            6. \( j \leftarrow \text{minimum value of labels in } sC \)
         5. **else**
            1. \( j \leftarrow j + 1 \)
         6. **end_if**
      2. **end_while**
   7. **if** \( sC \) is empty
      1. **break**
   8. **end_if**
   9. \( i \leftarrow i + 1 \)
   10. \( fl \leftarrow \text{find first label in } sC \)
   11. **Append** \( sC_{fl} \) **to** \( C_i \)
   12. Remove entries of label \( fl \) from \( sC \) group
   13. \( aD_i \leftarrow sCaD_{fl} \)
   14. \( j \leftarrow fl + 1 \)
   15. **end_while**
   16. Return \( C \)

**End**

indicates the average distance within sub-cluster \( sC_i \). We also have a group of clusters \( C=\{C_1,C_2,\ldots\} \) which is the output of this phase that contains a number of the resultant clusters of the given dataset in which it is self identified. Moreover, \( C \) is also the final result of KBCHT after considering released
vertices as mentioned previously. These released vertices are categorized into two groups. One of them identifies them as noise and outliers, and the other one assigns them back to their corresponding clusters.

First we add the first item of the group $sC$ to the group $C$ as shown in line 6 of Procedure 3.3. Then in line 7 we remove the first sub-cluster from the group $sC$. Thus we want to pick and remove sub-clusters from $sC$ and add them to the group of $C$ to one of its items or append them to be a new item or cluster in the group $C$. Hence, we want the group $sC$ to be empty for having the clusters in $C$. Now we have to calculate the average distance between data points in subcluster $C_i$ by assigning it the value of $sCaD_i$ because at this stage the group $C_i$ is the same as $sC_i$.

As stated previously, for computing the average distance of a given group of data points we construct its Delaunay triangulation to have geometry of triangles. Thus the average distance is the average length of edges of the triangles. The average distance of sub-cluster $C_i$ is stored in $aD_i$ of the group $aD$. The two pointers $i$, and $j$ in Procedure 3.3 deal with the two groups $C$ and $sC$ respectively. We trace all the sub-clusters in $sC$ thus we have to decide if one of these sub-clusters can be added to the picked item from the group $C$. So, we have sub-clusters that are merged together to form the final clusters. For deciding to merge or not, we compute the distance between the sub-cluster $C_i$ and each of the remaining sub-clusters in the group $sC$ as stated in line 13 of Procedure 3.3. We calculate the distance between two groups of data points by considering the distance between the two closest data points in which each of these two data points belongs to different group. Other methods of measuring distance between clusters can be found in [28]. If the distance between the cluster $C_i$ and one of the sub-clusters in $sC$ ($db2c$) is less than the average distance within the $C_i$, we add this sub-cluster of $sC$ to the cluster $C_i$ thus the cluster $C_i$ grows. In case of finding a candidate sub-cluster from the group $sC$ to be merged with one of the clusters in the group $C$, we remove this sub-cluster from the group $sC$ and check if $sC$ becomes empty to stop and get out from the Procedure 3.3 as shown in lines 21 to 23 and lines from 30 to 32. In
line 24 we have two merged sub-clusters. Hence, we recomputed the average distance within the grown cluster to reflect the actual average distance in it and to accept new merged sub-clusters. Line 25 reassigns a value for the pointer $j$ to point to the previous sub-clusters in $sC$ that does not meet the criteria of merging with old $C_i$. Because the characteristic of $C_i$ before merging is different than it after merging, thus we can have additional sub-clusters in $sC$ in which they could be merged with the new formed cluster.

When we reach line 33 of Procedure 3.3, this means that we have already caught one of the clusters of the given dataset from merging process. As a consequence, we have to look forward to finding other clusters in the given dataset. Thus we increment the pointer $i$ by one to point the next location in the group $C$ in which we want to find another cluster. Since we remove each of merged sub-clusters from the group $sC$, its labels will not be in a regular order. Hence, in line 34 we find the first label that exists in the group $sC$ and assign it to the variable $fl$. Now the first sub-cluster in the $sC$ is appended to a new index position in the group of $C$. Then it has been removed from the $sC$. The average distance within the new inserted sub-cluster in $C$ has assigned the value from its corresponding group of $sCaD$. We assign a new value for the pointer $j$ which points to sub-clusters in the group $sC$ as in line 38. We repeat the above steps until the group $sC$ becomes empty which means that we have obtained the desired clusters from the given dataset in the group $C$. To make $C$ as a final result of our algorithm, we should back to process the released vertices from the shrinking phase; this is done by assigning them to their corresponding clusters in $C$. This is done by telling each cluster in $C$ to pick the similar vertices to it which occurs when the distance from the cluster and the tested vertices is less than the average distance within this cluster. The remaining vertices are considered to be odd patterns in the entire dataset and we mark them as noise and outliers. Hence, we find the final results of clusters in the given dataset in a completely unsupervised manner.
3.2 A Toy example: Moon dataset

For illustration purposes, we consider the following example in Fig. 3.4 to be solved by our KBCHT algorithm.

In this dataset we have two moons distributed as shown in Fig. 3.4(a). KBCHT performs the standard Kmeans for obtaining the initial groups of clusters to be processed. Fig. 3.4(b) shows the result from the first phase of KBCHT algorithm in which we have two groups of initial clusters, one of them is with blue ‘o’ s data points and the second one is with red ‘*’ s data points. Now we can process these two groups in a parallel fashion. One thread picks the blue group and the second thread picks the red one. Hence, KBCHT algorithm executes its second phase in which KBCHT constructs the convex hull of the two groups as shown in Fig. 3.4(b), the closed set of solid black lines. At this step we have identified the vertices that represent each of the groups which reside on the drawn convex hull.

Fig. 3.4(c) visualizes the process of shrinking these vertices which is the output result of the Procedure 3.1. It has been noticed that each of the two groups has two connected set of vertices and the shrunk vertices forms a general polygon shape in which KBCHT can detect convex shape as well as non-convex shape clusters. Procedure 3.2 has triggered on the result shown in Fig. 3.4(c) to find the set of sub-clusters $sC$ for this moon dataset.

Fig. 3.4(d) shows four sub-clusters in which $sC=\{sC_1, sC_2, sC_3, sC_4\}$ such that $sC_1$ is the green ‘.’s data points, $sC_2$ is the black ‘.’s data points, $sC_3$ is the data points with blue ‘.’s and $sC_4$ is the data points with red ‘.’s. KBCHT algorithm checks these sub-clusters for merging process. Procedure 3.3 works on the result shown in Fig. 3.4(d) to find final clusters $C$. At the beginning, the group $C=\{sC_1\}$ that is $C_1$ is equal to $sC_1$ and $sC$ becomes $\{sC_2, sC_3, sC_4\}$ after removing $sC_1$ from it. KBCHT searches for one or more of the sub-clusters in $sC$ to be merged with $C_1$ according to Procedure 3.3. KBCHT finds that $sC_3$ should be merged with $C_1$ thus $C=\{sC_1+ sC_3\}$ where $C_1$ is the two sub-clusters $sC_1$ and $sC_3$ together. Then by the next iteration $C=\{sC_1+ sC_3, sC_2\}$ where $C_2$ is $sC_2$ and $sC=\{ sC_4\}$. According to Procedure 3.3 $sC_2$ should be merged with $sC_4$
thus $C = \{sC_1 + sC_3, sC_2 + sC_4\}$ and $sC_4$ is removed from $sC$ to be an empty group. Hence, we obtain the two clusters of the moon dataset. Fig. 3.4(e) shows the final of clusters that obtained by KBCHT algorithm.

Figure 3.4: Toy example. (a) The moon dataset. (b) Result from the standard Kmeans where blue ‘o’ s are in one cluster and red ‘*’ s are in another cluster and each of them are enclosed by its convex hull. (c) The shrinking process. (d) The process of finding sub-clusters. (e) The final result of our KBCHT algorithm.
3.3 Simulation and Results

3.3.1 Datasets results

We illustrate the strength of the KBCHT algorithm in a number of artificial two dimensional datasets, since the results can easily be verified visually. We generate four types of datasets DS1, DS2, DS3 and DS4 with 350, 1000, 2000, and 6000 data points respectively as shown in Fig. 3.5. Now, our purpose is to verify that KBCHT algorithm should be able to detect the clusters in these datasets, which can be differentiated by eyes, successfully. DS1 is considered to be a simple dataset that contains four well separated clusters, but the others

![Figure 3.5](image-url)
contain clusters with complex non-convex shapes, different sizes, densities, noise and outliers. We compare our algorithm with the Affinity Propagation [67] algorithm in which messages are exchanged between data points until the final clusters are found, because it does not need to be rerun many times each with different initial conditions and does not require to input the number $k$ of clusters. Also, we compare it with one of the recently spectral clustering algorithms because they are designed to handle non-convex shaped clusters and solve the problem of expensive computations of them which is the spectral clustering using Nystrom method [76].

As stated previously, we have to do the clustering task with no prior information knowledge about the given dataset and this is the goal of our proposed KBCHT algorithm. As a consequence, we have treated the given datasets as closed black boxes. For this reason, we have used the same value of $k$, in case of our proposed KBCHT algorithm and Spectral clustering using Nystrom algorithm, to figure out the output with no prior information. In addition, as we have generated these four artificial datasets, we do further step in assigning the true value of $k$ to the Spectral clustering using Nystrom algorithm.

**DS1:** This dataset contains 350 data points distributed into 4 well separated clusters. However, it is suitable to find its clusters using the standard Kmeans. But when we perform our KBCHT its first phase does not detect the four clusters correctly. We deal with the given dataset as black boxes that mean we do not have a prior knowledge about how many clusters exist or how they are distributed. Thus we roughly choose $k=5$ and our KBCHT correctly identifies the four clusters as shown in Fig. 3.6(a). Fig. 3.6(b) shows the result of clustering using Affinity Propagation in which it does not use any input parameters. However, it does not correctly detect the number of clusters. It has a result of 11 clusters. Since the spectral clustering using Nystrom method has two inputs: sigma and the number of clusters which are user defined. We use a sigma to equal 20 and use the same value of $k=5$ for the reason stated above. Even though this is a simple dataset Fig. 3.6(c) shows the weakness of the
spectral clustering when its parameters are not identified correctly. In which it has a result of five clusters where it wrongly splits one of the clusters into two. To be more honest with the used spectral clustering, we rerun it with $k=4$ and we obtain a result that correctly identifies the four clusters as shown in Fig. 3.6(d).

**DS2:** This dataset contains 1000 data points distributed into 8 complex shaped clusters and also with present noise and outliers. Fig. 3.7 (a) shows the effectiveness of our KBCHT algorithm in identifying the true 8 clusters however we roughly choose $k=10$. The noise and outliers are correctly marked as the black ‘+’s in Fig. 3.7(a). This is due to how we measure the Delaunay triangulation average $AVG$ within the initial group, where we exclude the convex hull vertices from the computation of the $AVG$ which eliminates the effect of outliers. The Affinity Propagation detects 20 clusters as shown in Fig. 3.7(b). For the spectral clustering using Nystrom method with sampling equals to 400 and we set sigma=20 and also we run it one time with $k=10$ as chosen for our KBCHT and another time using correct number of clusters $k=8$. Fig. 3.7(c) and Fig. 3.7(d) improves that our proposed KBCHT outperforms it in case of cluster quality.

**DS3:** This dataset contains 2000 data points distributed into 5 complex shaped clusters and also with present noise and outliers. Fig. 3.8 (a) shows the strength of our KBCHT algorithm in identifying 5 clusters correctly. However, we roughly choose $k=15$. The noise and outliers are correctly marked as the black ‘+’s in Fig. 3.8(a). In fact, the shrinking phase of KBCHT algorithm traverses all candidate vertices that could be engaged into the representative boundary of the given initial groups, this is due to the high priority given to the longest edge between two consecutive vertices to be shrunk according to the natural of the given initial group that it is reflected by the calculated delaunay triangulation average inside this group. The Affinity Propagation detects 28 clusters as shown in Fig. 3.8(b). We proceed as above in selecting sampling of 400 and sigma=20 for the spectral clustering using Nystrom method but it triggers an error on the value of sigma thus we tuned it to be equal to 5 and also we run it one time with
$k=15$ as chosen for our KBCHT and another time using the correct number of clusters $k=5$. Fig. 3.8(c) shows that it does not fully detect the cluster correctly but in Fig. 3.8(d) the spectral clustering shows that it handles two clusters correctly but without identifying noise and outliers.

**DS4:** This dataset contains 6000 data points distributed into 11 complex shaped clusters and also with presence of noise and outliers. Fig. 3.9 (a) shows that our KBCHT algorithm finds all of the 11 clusters correctly. However, we roughly choose $k=15$. The noise and outliers are correctly marked as the black ‘+’s in Fig. 3.9(a). The Affinity Propagation detects 53 clusters as shown in Fig. 3.9(b). The spectral clustering using Nystrom method with sampling equals to 400 and sigma=5; the result shown in Fig. 3.9(c) with $k=15$ as chosen for our KBCHT and when using the correct number of clusters $k=11$ we obtain clusters as shown in Fig. 3.9(d).

In the next Section, we do further analysis of the given results in which we measure the obtained clustering quality rather than subjectively evaluating the results.
Figure 3.6: Clustering Results of DS1: (a) our proposed KBCHT (using k=5). (b) Affinity Propagation. (c) Spectral clustering using Nystrom method (samples=100, sigma=20 and k=5). (d) Spectral clustering using Nystrom method (samples=100, sigma=20 and k=4).
Figure 3.7: Clustering Results of DS2: (a) our proposed KBCHT (using k=10), (b) Affinity Propagation. (c) Spectral clustering using Nystrom method (samples=400, sigma=20 and k=10). (d) Spectral clustering using Nystrom method (samples=400, sigma=20 and k=8).
Figure 3.8: Clustering Results of DS3: (a) our proposed KBCHT (using k=15). (b) Affinity Propagation. (c) Spectral clustering using Nystrom method (samples=400, sigma=5 and k=15). (d) Spectral clustering using Nystrom method (samples=400, sigma=5 and k=5).
Figure 3.9: Clustering Results of DS4: (a) our proposed KBCHT (using $k=15$). (b) Affinity Propagation. (c) Spectral clustering using Nystrom method (samples=400, $\sigma=20$ and $k=15$). (d) Spectral clustering using Nystrom method (samples=400, $\sigma=20$ and $k=11$).
3.3.2 Performance Analysis

In order to prove the effectiveness and the strength of our proposed KBCHT algorithm, we construct experiments for measuring the performance in case of time cost and clustering accuracy according to the visual results shown in Section 3.3.1. Table 3.1 shows the comparison of the performance of our proposed KBCHT algorithm and the used clustering algorithms in which we have been used in Section 3.3.1 for each of the results that obtained visually in previous section. We measure the time that it takes for completing its mission of clustering in seconds. Furthermore, it is important to measure how they are accurate in identifying the final results which is more important. For measuring the clustering accuracy, we follow the technique used in [77] and [76] to evaluate the quality of the resultant clusters and we use the following equation:

\[
\text{Accuracy} = \frac{\sum_{i=1}^{n} \delta(y_i, \text{map}(c_i))}{n}
\]  

(3.3)

Where \( n \) is the number of data points in the given dataset, \( y_i \) and \( c_i \) are the true cluster label and the obtained cluster label respectively. The delta function \( \delta(y, c) \) equals 1 if \( y = c \) and equals 0 otherwise. To measure the accuracy, we need a permutation function that maps each obtained cluster label to a true cluster label. Moreover, for achieving the optimal matching the Hungarian algorithm [78] is used.

As shown in Table 3.1, our proposed KBCHT algorithm has a clustering accuracy as high as 100% for all of the datasets that have been used. For DS1, we have also an accuracy of 100% for the spectral clustering using Nystrom method but when we use the exact number of clusters that included in DS1 even in this case our proposed KBCHT outperforms it in the execution time. But when using the same value of \( k \), our algorithm has the superiority in both the time cost and the clustering accuracy. To find the final results of clusters in DS1, our KBCHT takes 0.12 sec, 0.03 sec and 0.12 sec for the first, the second and the third of its phase respectively.

For the DS2, our proposed KBCHT algorithm executes the first phase in 0.3 sec, phase two in 1.03 sec and the last phase in 0.87 sec. Hence the overall time
cost is 2.2 sec as shown in Table 1. Our KBCHT takes 1.27 sec, 0.92 sec and 5.11 sec for its three phases respectively for the DS3. And for the DS4 it spends the most of its time in executing the third phase in which it takes 32.55 sec, the first and the second phases are executed in 5.1 sec and 10.84 sec respectively. However, for the DS4 our algorithm takes much time than the spectral clustering algorithm, the overall performance of KBCHT improves the clustering results in case of the quality of the results compared with this small gap reported in the execution time.

**Table 3.1:** Comparison of the performance of our proposed KBCHT algorithm and existing algorithms with time cost and clustering accuracy.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Algorithm</th>
<th>Time Cost (sec)</th>
<th>Clustering Accuracy (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DS1</td>
<td>Proposed KBCHT (k=5)</td>
<td>0.27</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>Affinity Propagation</td>
<td>7.58</td>
<td>39.62</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom (samples=100,sigma=20,k=5)</td>
<td>0.31</td>
<td>89.94</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom (samples=100,sigma=20,k=4)</td>
<td>0.29</td>
<td>100</td>
</tr>
<tr>
<td>DS2</td>
<td>Proposed KBCHT (k=10)</td>
<td>2.20</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>Affinity Propagation</td>
<td>21.38</td>
<td>53.06</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom (samples=400,sigma=20,k=10)</td>
<td>3.19</td>
<td>52.02</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom (samples=400,sigma=20,k=8)</td>
<td>2.97</td>
<td>60.76</td>
</tr>
<tr>
<td>DS3</td>
<td>Proposed KBCHT (k=15)</td>
<td>7.30</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>Affinity Propagation</td>
<td>41.78</td>
<td>23.69</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom (samples=400,sigma=5,k=15)</td>
<td>10.39</td>
<td>44.95</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom (samples=400,sigma=5,k=5)</td>
<td>10.27</td>
<td>91.84</td>
</tr>
<tr>
<td>DS4</td>
<td>Proposed KBCHT (k=15)</td>
<td>48.49</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>Affinity Propagation</td>
<td>453.98</td>
<td>25.62</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom (samples=400,sigma=20,k=15)</td>
<td>43.23</td>
<td>65.12</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom (samples=400,sigma=20,k=11)</td>
<td>43.71</td>
<td>66.21</td>
</tr>
</tbody>
</table>
Does the choice of \( k \) in our proposed algorithm affect the performance: As we have mentioned in the previous section, the proposed KBCHT does not require prior knowledge information for finding the final results of clusters in a given dataset. It makes self analysis to discover the clusters. But how it performs when changing the value of \( k \)? For this purpose we construct experiments on the DS3 and we choose \( k \) to be from 2 to 70. For selecting \( k=2 \), the DS3 has been partitioned into two initial groups from the first phase as shown in Fig. 3.10(a). In this case all the overhead will be concentrated on the second phase of KBCHT in which the shrinking process investigates more complex shaped group and more vertices to find a set of shrunk vertices. KBCHT does not accept to have only one partition from the first phase. The two partitions of first phase are illustrated in Fig. 3.10(a) that takes 0.34 sec. Fig. 3.10(b) and 3.10(c) show the second phase operations of shrinking vertices and finding sub-clusters (8 sub-clusters) respectively. In which the effect of noise and outliers are eliminated as shown in Fig. 3.10(c). The second phase takes 18.13 sec to be accomplished. Fig. 3.10(d) shows the final result of KBCHT after merging phase and detecting noise and outliers which it takes 2.75 sec.

Fig. 3.11 shows how each of the three phases of our KBCHT algorithm performs with different values of \( k \). It has been shown that the most consuming phase is the third one with the increasing values of \( k \). Since the number of comparisons increased between pairs of sub-clusters. As increasing the value of \( k \), the second phase execution time drops dramatically. Because as the group under shrinking process becomes smaller, the complexity of the group decreases thus it can be handled faster. The most challenging case on shrinking of DS3 is when \( k=2 \) as shown in Fig. 3.10(b) in which each group has long embedded boundaries to be investigated by the shrinking process. KBCHT was proud to face this challenge to prove its efficiency and strength in handling this type of dataset. In Fig. 3.12 we show the overall time cost of the KBCHT algorithm when varying the number of \( k \). As the value of \( k \) increased the overall time cost has increased too. The best value of \( k \) is when it equals 10. At this
value of \( k \), the execution time spends less time among other values and equals to 6.49 sec. However, we choose the value of \( k \) to obtain initial groups from the first phase of KBCHT. The resultant number of partitions is not always the same as \( k \). This is because KBCHT uses the standard Kmeans in its first phase and it may suffer from bad dropped initial prototypes thus a problem of dead prototypes may be occurred. Fig. 3.13 shows the exact number of obtained partitions from the first phase as the value of \( k \) increases. In case of \( k=15 \) in this experiment, KBCHT obtains only 7 partitions from the first phase which it takes 0.89 sec to have them. According to this 11 sub-clusters have been

Figure 3.10: Analysis of KBCHT when \( k=2 \) on DS3. (a) Result from the first phase. (b) The Shrinking process. (c) the sub-clusters which is found by KBCHT (d) The final result of KBCHT after merging sub-clusters and identifying noise and outliers.
considered in 2.72 sec then the merging process to find the final clusters takes
3.93 sec thus the overall time cost is 7.54 sec. However, in Fig. 3.8(a) and
Table 3.1 when we have used the value of \( k \) equals 15, we caught 14 initial
partitions from the first phase of KBCHT and 16 sub-clusters to be merged. We
evaluate the clustering accuracy for each used value of \( k \). As illustrated in Fig.
3.14, KBCHT maintains a steady state of high accuracy rate as 100% for the
majority of the used \( k \). For \( k=50, 55, 60 \) and 70, the clustering accuracy is
99.21%, 98.30%, 99.94% and 99.65% respectively.

![Figure 3.11: Time cost (sec) of KBCHT three
phases vs. varying number of \( k \). red: first phase,
blue: second phase and black: third phase. (based on DS3)](image1)

![Figure 3.12: Time cost (sec) of KBCHT vs.
varying number of \( k \). (based on DS3)](image2)

![Figure 3.13: Obtained number of initial
partitions from the first phase vs. varying
number of \( k \). (based on DS3)](image3)

![Figure 3.14: Measuring the clustering
accuracy (%) of KBCHT vs. varying number of
\( k \). (based on DS3)](image4)
We analyze the result of how $k$ affects our KBCHT algorithm based on one of our complex generated dataset. To do further investigation on $k$, thus we can relatively make some specific judgment. We have generated relatively simple dataset contains 800 data points distributed into 5 clusters which is named DS5 as shown in Fig. 3.15.

We have repeated the same experiment situation as we did for DS3 in which we run KBCHT algorithm on the dataset DS5 with varying $k$ from 2 to 70.

Fig. 3.16 shows time cost in seconds for each phase of KBCHT algorithm. It is obvious that the third phase is the most consuming one especially when $k$ increased. The obtained result in Fig. 3.16 is somewhat close in its concept to the result obtained in Fig. 3.11. Fig. 3.17 provides the overall time cost in seconds for KBCHT algorithm in which it starts high when $k=2$ then it is dropped then as $k$ increased the execution time increased. Fig. 3.18 gives an overview of the actual number of partitions that obtained from the first phase since it suffers from bad initial condition that caused dead prototypes. In addition, Fig. 3.19 shows the clustering accuracy as increasing the value of $k$ which maintains the highest steadily state.

![Artificial dataset DS5.](image)
Experimenting on DS3 and DS5, we can notice that the overhead is high with respect to the execution time when $k=2$ and drops dramatically when $k=4$. In case of clustering accuracy, we can conclude that when we have applied some of the large values of $k$, the accuracy may suffer slightly in case of complex shaped dataset.

For DS3, the best result can be achieved when $k=10$. Moreover, the result may be acceptable when $k$ is ranging between 5 and 15. On the other side of
DS5, we have obtained the best result when $k=4$. Furthermore, when $k$ is between 4 and 10 the result could be acceptable. Hence, we can choose the lower bound of $k$ to be 5. Of course, this does not prevent the interested researchers from using any approaches that offers auto determination of the value of $k$ for Kmeans algorithm. However, we recommend that the value of $k$ is not less than 5.
Chapter 4

KBCHT: A TOPOLOGY PRESERVING MAPPING AS A PREPROCESSING

4.1 Overview

In the former Chapter, we have proved the efficiency of our proposed KBCHT algorithm on our generated 2-D artificial datasets. In this Chapter, we demonstrate the performance of our KBCHT algorithm based on high dimensional real datasets. In order to handle real datasets adequately, there high dimensionality needs to be reduced for coping the cures of dimensionality and other undesired features of high dimensional space [83]. The dimensionality reduction can be used as a visualization tool of high dimensional data [83].

There are many researches that are concerned in reducing the high dimensionality for data analysis. Furthermore, to enhance the performance of classification or clustering on these data, their dimensionality should be reduced before applying classification or clustering techniques [83, 93-96]. As a consequence, we can benefit from any of the available techniques in reducing the dimensionality of the given datasets to be engaged with our proposed KBCHT algorithm.

In this thesis, we have used a topographic mapping technique in conjugate with our KBCHT algorithm. A topographic mapping is a transformation of high
dimensional data which preserves some structure in the data such as the points which are mapped close to each other share some common properties while in contrast the points which are mapped far from each other do not share a common feature or property.

In this Chapter, we use the Generative Topographic Mapping (GTM) as a basis preprocessing step for our proposed KBCHT algorithm.

4.2 Generative Topographic Mapping (GTM)

The GTM is a statistical model for modeling the probability density of data points and finding non-linear mapping of high dimensional space onto low dimensional space.

The basis of the GTM is to generate a grid of $K$ latent points $z_1, z_2, \ldots, z_K$ in latent space. These latent points are mapped non-linearly into the data space using a set of $M$ basis Gaussian functions with respect to a set of weights $W$, where $W$ is $M \times D$, and $D$ is the dimensionality of data space. The GTM is optimized using the Expectation-Maximization (EM) algorithm [2], [28].

4.3 Simulation Experiments

After validating the performance of the proposed algorithm on complex artificial datasets, we have accomplished many experiments to evaluate the efficiency of our proposed algorithm on high dimensional datasets over the same competing algorithms as in Chapter 3. All the experiments in this thesis have been performed using an Intel dual at 1.87 GHz with 2 GB of RAM. A brief description of the used real datasets and simulation results will be presented in the following sub sections.

4.3.1 Real Datasets

In our simulation experiments we have used 10 real datasets. All of these datasets are available from the UCI machine learning repository [88]. General information about these datasets is given in Table 4.1.
Some of these real datasets have missing values and different types of variables with different scales. Hence, they would be preprocessing before engaging them into the clustering process (Appendix B explains how to prepare the data before clustering).

Table 4.1: The Descriptions of the used UCI datasets

<table>
<thead>
<tr>
<th>No.</th>
<th>Datasets</th>
<th># of Objects</th>
<th># of Dimensions</th>
<th># of clusters</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Iris</td>
<td>150</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>2</td>
<td>Libras movements</td>
<td>360</td>
<td>90</td>
<td>15</td>
</tr>
<tr>
<td>3</td>
<td>Wine</td>
<td>178</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>4</td>
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<td>4601</td>
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<td>8</td>
<td>Yeast</td>
<td>1484</td>
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<td>10</td>
<td>Dermatology</td>
<td>366</td>
<td>34</td>
<td>6</td>
</tr>
</tbody>
</table>

4.3.2 Parameters Setting

As stated previously, we have to do the clustering task with no prior information knowledge about the given dataset and this is the goal of our proposed KBCHT algorithm. As a consequence, we have treated the given datasets as closed black boxes. As a result in our experiments, the value of \( k \) is set to 5 for all of the used real datasets except for the largest used dataset (Pendigits) we use a value of 15. The same value of \( k \) is used for both our proposed KBCHT algorithm and the spectral clustering using Nystrom algorithm.

In this Chapter and as mentioned previously, we have used the GTM as a basis preprocessing step for our proposed algorithm. Hence, the grid of latent points and the grid of base functions have been chosen to be 10x10 and 3x3 respectively for all of the 10 used UCI datasets.

Moreover, for the spectral clustering using Nystrom algorithm, the value of sigma is set to 50 and the sample size has been chosen to be 100 for all of the
used datasets. But we have faced a problem that the used default sample size of 100 is not valid for some datasets. Thus, we have chosen a sample size of:

- 150 for Wine, Glass, Yeast, Arrhythmia and Dermatology.
- 300 for Image segmentation.
- 1500 for Pendigits.
- 4601 for Spambase.

In which the Spambase dataset does not accept to have any sample size except the value of its whole number of objects.

For the Affinity Propagation algorithm, we have used its default setting as in [67].

### 4.3.3 Results Analysis

For illustration of the effectiveness of our algorithm, we have conducted the same performance measurements that we already used in Chapter 3.

Table 4.2 and Fig. 4.3 summarize the obtained simulation results on real datasets in which our proposed algorithm outperforms the other algorithms.

For Iris dataset, as an example, our proposed KBCHT using GTM has the highest performance comparing with other algorithms. It has an accuracy of 88%. In contrast, the spectral clustering using Nystrom has an accuracy of 57.33% and the Affinity Propagation has 33.33% clustering accuracy. Moreover, our proposed KBCHT using GTM obtained more accurate result faster than the other algorithms.

Fig. 4.1 shows the result of GTM on the Iris dataset, where the Iris dataset has been projected on 2 dimensional space. It is obvious that the Iris dataset has one linearly separable cluster from the other two clusters; in which the other two clusters are overlapped. The GTM takes 0.26 sec, the first phase of KBCHT takes 0.06 sec, the second phase takes 0.14 sec and the last phase executed in 0.11 sec. Hence, the overall time cost for the proposed KBCHT using GTM is 0.57 sec.
Figure 4.1: The result of using GTM on Iris dataset (the three clusters are distinguished using different color and symbol for each cluster).

The results of experiments on the rest of the used UCI datasets are summarized in Table 4.2. In which the performance of our proposed KBCHT using GTM and exiting algorithms has been compared with time cost in seconds and the clustering accuracy which is calculated as mentioned in Chapter 3.
Table 4.2: Comparisons of the performance of our proposed KBCHT algorithm using GTM and existing algorithms with time cost and clustering accuracy on real datasets.

<table>
<thead>
<tr>
<th>Datasets</th>
<th>Algorithms</th>
<th>Time Cost (sec)</th>
<th>Clustering Accuracy (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iris</td>
<td>Proposed KBCHT using GTM</td>
<td>0.57</td>
<td>88.00</td>
</tr>
<tr>
<td></td>
<td>Affinity Propagation</td>
<td>6.03</td>
<td>33.33</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom</td>
<td>0.61</td>
<td>57.33</td>
</tr>
<tr>
<td>Libras Movements</td>
<td>Proposed KBCHT using GTM</td>
<td>0.99</td>
<td>30.56</td>
</tr>
<tr>
<td></td>
<td>Affinity Propagation</td>
<td>14.64</td>
<td>6.67</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom</td>
<td>0.94</td>
<td>24.72</td>
</tr>
<tr>
<td>Wine</td>
<td>Proposed KBCHT using GTM</td>
<td>0.63</td>
<td>93.82</td>
</tr>
<tr>
<td></td>
<td>Affinity Propagation</td>
<td>5.77</td>
<td>39.88</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom</td>
<td>0.79</td>
<td>65.16</td>
</tr>
<tr>
<td>Glass</td>
<td>Proposed KBCHT using GTM</td>
<td>0.70</td>
<td>65.02</td>
</tr>
<tr>
<td></td>
<td>Affinity Propagation</td>
<td>6.81</td>
<td>48.59</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom</td>
<td>1.29</td>
<td>63.55</td>
</tr>
<tr>
<td>Pendigits</td>
<td>Proposed KBCHT using GTM</td>
<td>103.13</td>
<td>38.27</td>
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<tr>
<td></td>
<td>Affinity Propagation</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom</td>
<td>212.11</td>
<td>29.10</td>
</tr>
<tr>
<td>Image Segmentation</td>
<td>Proposed KBCHT using GTM</td>
<td>3.88</td>
<td>31.52</td>
</tr>
<tr>
<td></td>
<td>Affinity Propagation</td>
<td>214.32</td>
<td>14.29</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom</td>
<td>3.81</td>
<td>14.58</td>
</tr>
<tr>
<td>Spambase</td>
<td>Proposed KBCHT using GTM</td>
<td>93.03</td>
<td>78.91</td>
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<tr>
<td></td>
<td>Affinity Propagation</td>
<td>1172.53</td>
<td>60.59</td>
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<tr>
<td></td>
<td>Spectral clustering using Nystrom</td>
<td>6447.31</td>
<td>62.53</td>
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<tr>
<td>Yeast</td>
<td>Proposed KBCHT using GTM</td>
<td>6.80</td>
<td>39.80</td>
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<tr>
<td></td>
<td>Affinity Propagation</td>
<td>106.53</td>
<td>31.19</td>
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<tr>
<td></td>
<td>Spectral clustering using Nystrom</td>
<td>2.80</td>
<td>20.21</td>
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<tr>
<td>Arrhythmia</td>
<td>Proposed KBCHT using GTM</td>
<td>2.13</td>
<td>56.94</td>
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<td></td>
<td>Affinity Propagation</td>
<td>13.78</td>
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<td></td>
<td>Spectral clustering using Nystrom</td>
<td>2.38</td>
<td>55.75</td>
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<tr>
<td>Dermatology</td>
<td>Proposed KBCHT using GTM</td>
<td>0.97</td>
<td>86.07</td>
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<tr>
<td></td>
<td>Affinity Propagation</td>
<td>10.59</td>
<td>22.95</td>
</tr>
<tr>
<td></td>
<td>Spectral clustering using Nystrom</td>
<td>1.45</td>
<td>58.47</td>
</tr>
</tbody>
</table>
For illustration purpose and easy to compare, we provide the clustering accuracy that obtained from all the used datasets by our proposed algorithm and the existing algorithm in Fig. 4.2. On these datasets our proposed algorithm achieves the highest accuracy compared with other algorithms, the results indicates the effectiveness and the strength of our proposed algorithm. From Table 4.2 and Fig 4.2, we can notice that the Affinity Propagation algorithm failed to provide result on the Pendigits dataset due to memory limitation which gives an indication that the Affinity Propagation algorithm suffers from memory difficulties.

![Figure 4.2: Clustering accuracy results on real datasets](image)

Figure 4.2: Clustering accuracy results on real datasets
Chapter 5

CONCLUSIONS AND FUTURE RESEARCH

5.1 Conclusions and Future Research

In this thesis, we have introduced a new clustering algorithm named KBCHT (Kmeans-Based Convex Hull Triangulation clustering algorithm) that is able to detect clusters with complex non-convex shapes, different sizes, densities, noise and outliers.

The idea behind the KBCHT is to divide the clustering task into three phases. The first phase benefit from the first run of the widely used simple Kmeans algorithm in which the experiments show that KBCHT gives good results despite it is badly chosen initial conditions. Then in the second phase KBCHT takes the initial groups from the first phase and constructs a convex hull for each group. The vertices obtained by convex hull are shrunk until we catch a set of sub-clusters in which the shrinking mechanism does not maintain the convexity of processed groups. The last phase applies merging of sub-clusters based on the Delaunay triangulation. We also have used one of the famous topology preserving mapping methods which is the Generative Topographic Mapping (GTM). The GTM have been used as a preprocessing step to our proposed KBCHT algorithm. The simulation results show the
superiority of our proposed algorithm over recently introduced ones in case of the clustering accuracy and time cost.

In future works and according to obtained results in Fig. 3.11, the third phase consumes the most time in the KBCHT since we consider all data points in each sub-clusters, thus we will do farther analysis to benefit from the sampling techniques to make it faster. While we have assumed the Euclidean distance throughout this thesis, we can use general metrics as shown in appendix C to show the effect of the used metric on our proposed algorithm. The computed value of $AVG$ average from Delaunay triangulation can be recomputed each time we have a shrunk vertex according to our proposed mechanism, thus we may have more robust value. The proposed KBCHT algorithm can be implemented in a distributed environment. Also as have shown in Chapter 4, the GTM uses a fixed grid of latent points and a fixed grid of Gaussian basis function. Hence the initialization of GTM is the same despite the used data set. This may affect the training results. Thus it needs further investigations to adaptively choose the initial parameters based on the given dataset. Furthermore, in future work we can conduct more experiments that directly apply the proposed algorithm to high dimensional datasets without preprocessing the data.
Appendices
Appendix A

SIMULATION ENVIRONMENT

We have implemented our proposed KBCHT algorithm using MATLAB (version 7.9.0.529 (R2009b)). Furthermore, all the experiments in this thesis have been performed using 64-bit windows environment of Intel dual at 1.87 GHz with 2 GB of RAM.

In fact, MATLAB (MATrix LABoratory) is a high-level language and interactive environment that enables you to perform computationally intensive tasks faster than with traditional programming languages such as C, C++, and Fortran.

In addition, we have used the MATLAB implementation of both Affinity Propagation algorithm and Spectral clustering using Nystrom algorithm. These implementations are available from the websites (last visit on Oct. 2011):

- http://alumni.cs.ucsb.edu/~wychen/sc.html
Dealing with missing values

The collected data can be represented in a matrix notation (two modes) as in equation 1.1 of chapter 1. Hence, the missing values could be in records or columns. In record: if the most of data are missing then we delete this record. In column: if the most of data are missing then we delete this column. But if the missing values are not too much, we have to replace these missing values with:

1. The average value of all column data in which the missing occurred and then is done before running clustering algorithms.
2. By making a pre clustering algorithm:
   2.1 Make clusters based on all variables, then take average value of objects that belong to the same cluster to replace the missing values.
   2.2 To know which one variable (column) is more similar to the one with missing values. Then replace the missing values with the average value of the most similar variable.

Normalization

The attributes of the entire dataset may be collected from different scales. Hence, the large scale ones may cause the bias. So, data should be normalized
thus all attributes of the dataset would be in the same scale. Moreover, the objects of the dataset can be described by different types of attributes such as nominal, categorical, ordinal, interval or ratio. There are different approaches to cluster these objects. One of them is convert attributes of different types into attributes of the same type. A more detail of scale conversion could be found in chapter 3 of [28].

**Sampling**
Sampling can be used to pick a portion of large dataset because processing the entire dataset is too expensive and time consuming. The sampling may be obtained randomly which is the simplest way. But to be more effective, it should reflect an approximation of the same characteristic as the original dataset.

**Dimensionality reduction**
High dimensional dataset requires a large amount of time and memory to be analyzed. There are two major types of dimensionality reduction methods: feature transformation and feature selection.

In feature transformation, the original high dimensional space is projected linearly or non-linearly into a lower dimensional space.

In feature selection, this method selects a subset of meaningful dimensions from the original high dimensions of the entire dataset.
Appendix C

DISTANCE METRICS

In this section, we present some of the common used metrics. The Minkowski (Manhatten, Euclidean, and Maximum distances), the Mahalanobis distance and the Cosine distance.

Minkowski distance

A metric or distance is used to measure the similarity or dissimilarity between objects. While we have assumed the Euclidean distance throughout this thesis, a more general metric between two objects \( X \) and \( Y \) could be addressed. This general metric is named Minkowski metric and is defined as \((p\text{-norm}):\)

\[
d_p(X,Y) = \left( \sum_{i=1}^{d} |x_i - y_i|^p \right)^{1/p}, \quad p \geq 1
\]  

(C. 1)

Where \( d \) is the dimension of objects \( X \) \((x_1,x_2,\ldots,x_d)\) and \( Y \) \((y_1,y_2,\ldots,y_d)\). Thus the Euclidean distance is when \( p = 2 \) (2-norm). And when \( p = 1 \) (1-norm) \( d_p \) is called Manhattan or city block distance which is the shortest path between \( X \) and \( Y \) where each segment of the path is parallel to a coordinate axis. And when \( p = \infty \) (\( \infty \)-norm) is called maximum distance which is the distance between \( X \) and \( Y \) corresponds to the maximum distance between the projections of \( X \) and \( Y \) onto each of the \( d \) coordinate axes.
**Mahalanobis distance**

Mahalanobis distance removes the distance distortion caused by linear combinations of attributes. It is defined as:

$$d_{mah}(X, Y) = \sqrt{(X - Y)^T \Sigma^{-1} (X - Y)}$$  \hspace{1cm} (C.2)

Where $\Sigma$ is the covariance matrix of the dataset. If $\Sigma = Identity$, this is the Euclidean distance.

An important property of the Mahalanobis distance is that it is invariant under nonsingular transformation. It suffers from high computation needed to compute the covariance matrix based on all objects in the dataset.

**Cosine distance**

Cosine distance calculates the cosine of the angle between two vectors. It can be defined as:

$$d_{cos}(X, Y) = \frac{X \cdot Y}{|X||Y|}$$  \hspace{1cm} (C.3)

Where $X \cdot Y$ is the dot product and $| . |$ is the (2-norm).
References


