Improving a Particle Swarm Optimization-based Clustering Method

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Improving a Particle Swarm Optimization-based Clustering Method

A thesis
Submitted to the Graduate Faculty of the University of New Orleans
in partial fulfillment of the requirements of the degree of
Master of Science in
Engineering
Electrical

By
Sharif Shahadat
B.Sc. Ahsanullah University of Science and Technology
May 2017
Dedication

I am grateful to the Almighty creator for the infinite blessings in my life, which led to the completion of my Master’s thesis. I dedicate this work to my loving parents and my beloved brother, whose prayers and guidance helped me get through this journey. It was due their constant support that made me reach the destination for this journey.
Acknowledgements

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I would like to thank Dr. Vesselin Jilkov, and Dr. Ebrahim Amiri, for accepting the request to serve as members in my thesis committee. Their sincere advices help shape the thesis in the best way possible. I would also like to thank my friends and colleagues, especially Monir Hossain and Sumaiya Iqbal for their invaluable support throughout my studies.

I would like to thank my parents and my brother for their guidance throughout my life. Words are not capable of conveying my feelings towards them.

Finally, I would like to express my deepest gratitude to the University of New Orleans for providing the necessary tools and environment for this work. I recognize that this research would not have been possible without the resources made available by the University.
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Abstract

This thesis discusses clustering related works with emphasis on Particle Swarm Optimization (PSO) principles. Specifically, we review in detail the PSO clustering algorithm proposed by Van Der Merwe & Engelbrecht, the particle swarm clustering (PSC) algorithm proposed by Cohen & de Castro, Szabo’s modified PSC (mPSC), and Georgieva & Engelbrecht’s Cooperative-Multi-Population PSO (CMPSO). In this thesis, an improvement over Van Der Merwe & Engelbrecht’s PSO clustering has been proposed and tested for standard datasets. The improvements observed in those experiments vary from slight to moderate, both in terms of minimizing the cost function, and in terms of run time.

Keywords: Clustering, Particle Swarm Optimization, Metaheuristics, Data Mining
Chapter 1
Introduction

1.1 Motivation

Large datasets are getting more common in the recent years. As a result, it is becoming exceedingly difficult for human analysts to interpret the patterns in large datasets. Cluster analysis or data clustering come to rescue in this regard by offering a large number of vital techniques for the field of pattern recognition. The objective of data clustering is to divide a large amount of data into groups (clusters) based on certain similarity criteria. Clustering finds applications in many fields, including data mining, decision making, bioinformatics, machine learning, object recognition, signal processing, and image segmentation [1–3]. This thesis introduces a novel data clustering algorithm based on Particle Swarm Optimization (PSO).

1.2 Background

Everitt (1974) collected the following two definitions of a cluster [4]:

“A cluster is an aggregation of points in the test space such that the distance between any two points in the cluster is less than the distance between any point in the cluster and any point not in it.”
“Clusters may be described as connected regions of a multi-dimensional space containing a relatively high density of points, separated from other such regions by a region containing a relatively low density of points.”

Clustering can help discover inherent structures in unlabeled data. Clustering algorithms attempt to group data and reveal these structures by optimizing a certain objective function. Nevertheless, a global optimal solution to a problem might not be always found. Moreover, because of the unsupervised nature of clustering algorithms, it is not always guaranteed that the choice of the objective function is appropriate for the problem at hand. For dimensions higher than three, the solutions found from clustering may often be far away from the optimal solution.

1.3 Complexity of Clustering Problem

Although for simple scenarios, the process of clustering might appear to be fairly straightforward, for more general cases, it can be a significantly complex process. Figure 1.1(a) presents an example in which it is difficult to determine appropriate clusters as well as the number of clusters which may suitably reveal the data structure. On the other hand, in figure 1.1(b), it is apparent that four clusters are present. The data-driven nature of various applications increases difficulty in designing a universal algorithm that can accurately and efficiently discover the clusters in the provided data. Also, another factor is the unsupervised nature of the data sets. These kinds of problems encouraged the creation of various types of clustering algorithms such as evolutionary and metaheuristics algorithms. An Evolutionary Algorithm (EA) is a population-based algorithm which is inspired by natural evolution or
Figure 1.1: Complexity of clustering

the collective behavior of natural self-organizing system (e.g. ant colony, bee colony, bird flocking, fish schooling etc.). Specifically, EAs, when used for clustering, have been shown to provide near-optimal solutions for unsupervised data sets. A large number of algorithms have been created using EAs to solve clustering problems [5–7].

1.4 Clustering Algorithms

According to [1], there are two main approaches to clustering: partitional clustering and hierarchical clustering. Partitional clustering [8] produces various partitions and keeps evaluating those partitions using a certain condition (for example, the minimum sum of squares). Hierarchical clustering [9] creates a hierarchy of clusters based on a given dataset using specific criteria. For example, if vehicles represent a class, public transportation vehicles would be a subclass. Furthermore, sedans would be a subclass of public transportation vehicles and different brands are different subclasses).
Hierarchical clustering, on the other hand, does not assume the number of clusters beforehand and can be applied to any dataset. However, hierarchical clustering algorithms do not perform well for larger datasets because due to the amount of time and memory complexity requirements. Although various clustering algorithms exist, the selection of an appropriate algorithm depends on the particular problem.

1.4.1 \textit{k}-means

The \textit{k}-means algorithm is the common choice for clustering large datasets, owing to its low complexity and high execution speed. The main objective of \textit{k}-means is to associate every data point in the dataset to a prototype point (centroid). Essentially, the prototype point acts as a representative of its cluster. Although, the algorithm is simple and efficient, the drawback of \textit{k}-means is that it does not work well with non-convex datasets. Furthermore,
the number of clusters, \( k \), has to be specified before execution of the algorithm. The choice of \( k \) usually has a deep impact on the performance of the algorithm. The \( k \)-means algorithm consists of the following steps:

1. Initialize the number of centroids, \( k \), randomly. In some versions of the algorithm the initial centroids are chosen from the dataset

2. Determine the distance between each data point and each one of the \( k \) centroids

3. Assign each point to the cluster whose centroid is the closest one

4. Update the positions of centroids towards the center of gravity of their respective clusters

5. Repeat steps 2, 3 and 4 until the centroid positions change less than a user-defined threshold

The objective function for the \( k \)-means algorithm is the following:

\[
J = \sum_{j=1}^{k} \sum_{i=1}^{n} ||x^{(j)}_i - c_j||^2
\]  

In equation 1.1, \( k \) is the number of clusters, \( n \) is number of data points, and the term \( ||x^{(j)}_i - c_j||^2 \) represents the distance of the \( i \)-th data point, which has been assigned to the \( j \)-th cluster, from the \( j \)-th centroid.

1.4.2 \( k \)-means++

Initialization plays a crucial role in \( k \)-means algorithm [9]. Also, there is high possibility that the algorithm can get stuck on a local minimum. An example of non-spherical data
clustering using \(k\)-means is shown in figure 1.2.

The \(k\)-means++ algorithm proposed by Arthur and Vassilvitskii [10] starts by picking the \(k\) centers one at a time. After that, choosing each point at random with probability proportional to its squared distance from the already chosen centers.

### 1.4.3 Swarm Intelligence

The collective behavior of ants, bees, fishes, birds and other species, or what is usually called Swarm Intelligence (SI), has been proven a fascinating topic for researchers for quite some time. Usually, SI shows a structured order which is integrated into the system of species. The way the swarms move also appears to possess some characteristics of a single entity [11].

The swarm’s behavior may consist of the following properties:

1. Homogeneity: Every swarm member has the same type of behavior.

2. Locality: Each swarm member’s motion is influenced by other closely located members.
3. Collision Avoidance: Swarm members attempt to avoid collision with other nearby swarm members.

4. Velocity Matching: Swarm members attempt to match the velocity of other nearby members.

Algorithms such as Particle Swarm Optimization (PSO) and Ant Colony Optimization (ACO) have been developed in the literature [12,13] based on the swarm intelligence of bird flocks and ant colonies.

1.4.3.1 Particle Swarm Optimization

Particle Swarm Optimization (PSO) is an iterative computational method based on SI for optimization of nonlinear functions. The original algorithm was proposed by Kennedy and Eberhart (1995) [12]. PSO mimics the bird flocking behavior to find an optimal solution to a certain optimization problem. The algorithm is initialized with a defined number of particles which represent a set of possible solutions for the problem at hand. Each particle has a random velocity and each moves within the solution space in each iteration. The algorithm considers each particle’s personal best position and global best position after every iteration, in order to determine the particle’s velocity. After the initial proposal of PSO, various types of improved variants emerged, focusing on different applications. PSO has also been modified for the purpose of clustering data, and has showed some promising results compared to other algorithms [14–17].
1.4.3.2 Ant Colony Optimization

Similarly to PSO, Ant Colony Optimization (ACO) is also an iterative computational method for optimization of nonlinear functions, which was proposed by Marco Dorigo (1992) [13]. It is based upon the behavior of ants for seeking a source of food from their colony. Ants leave a trail of pheromone on their way back to the colony after finding food. The ants get attracted to the pheromone and follow the trail in the hope of finding the food source. However, the pheromone trail evaporates quickly, so that the shortest path to the food source gets the highest utilization. This behavior of ants has been put into use as the ACO algorithm for finding the optimal path in a graph space. The concept of ACO has also been used for generating clustering algorithms, some of which have been proved to be working well with different data sets [18–20]

1.4.4 Evolutionary Algorithms

Taking inspiration from nature, EAs use the following general steps to find out the solution to an optimization problem [21,22]

1. Generate a random initial population

2. Evaluate all individuals from the population

3. Choose the best individuals from the population to generate the next generation

4. Create the next generation

5. Repeat steps 2-4 until a stopping criterion is met. The best possible solution or a group of solutions, in the case of multi-objective problems, is chosen
The basic idea can be described in short as “survival of the fittest”. The most popular types of EAs are Genetic Algorithm and Differential Evolution.

1.4.4.1 Genetic Algorithm

Genetic Algorithm (GA) is one of the popular EAs which relies upon nature inspired operations such as selection, crossover, and mutation [23]. Roulette wheel selection [ref] is the widely used selection method which gives every chromosome a space in an imaginary roulette wheel according to their fitness score. It ensures the chromosomes with higher fitness scores are more likely to be selected. Crossover generates children from parent chromosomes, where each child takes one section of each parent’s chromosome. Finally, after selection and crossover, mutation applies a small random change in the new generation of chromosomes, which effectively creates a diversity among the population. The steps followed to generate solutions to an optimization problem using GA can be summarized as follows:

1. Generate initial population
2. Rank and evaluate individual fitness
3. Apply genetic operators such as crossover, mutation or selection on the chromosomes
4. Create population for next generation
5. Repeat steps 2-4 until the best possible solution or a group of solutions, in the case of multi-objective problems, is chosen

A lot of GA variants have been developed serving various optimization problems. Some of the recent ones have been mentioned in [24–26]. Some of the GA variants have also been
used for data clustering and have showed promising results [27–30].

1.5 Summary

Clustering is the process of grouping data or observations into a number of meaningful groups subject to further processing. In this chapter, the general concepts of various clustering algorithms and have been discussed, and some of the fundamental issues have been presented. Although the Thesis mainly concentrates on PSO and $k$-means, some additional algorithms which have been used in data clustering have been presented for information purposes. Each clustering model comes with its own advantages and disadvantages. In particular, the algorithms discussed in this chapter, such as PSO and $k$-means, fall into the class of partitional clustering algorithms. Partitional algorithms are generally more lightweight which make them easily applicable to larger datasets. Some drawbacks of partitional algorithms are the following:

1. They are limited to forming clusters around prototypes

2. The appropriate number of prototypes is subjective and, for some algorithms, has to be inferred using the appropriate cluster validation index

3. Solutions are subject to sub-optimality as they are dependent on the initial positions of the prototypes

Of course, some of the above may also be drawbacks of non-partitional algorithms.
1.6 Organization of Thesis

This thesis consists of four chapters which are organized as follows:

- **Chapter 1** introduces reader to the concepts of data clustering and discusses the various algorithms available to do clustering.

- **Chapter 2** introduces PSO and summarizes previous work that used PSO for data clustering.

- **Chapter 3** explains the proposed improvements and details the performance comparison of the algorithm presented by Van Der Merwe and Engelbrecht [17].

- **Chapter 4** summarizes the findings of this thesis, discusses its contributions and limitations. Future work has also been discussed in this chapter.

This thesis also contains following two appendices:

- **Appendix A** provides the MATLAB codes used to generate the results.

- **Appendix B** provides the acronyms used throughout the thesis, as well as their definitions.
Chapter 2
Data Clustering and Particle Swarm Optimization

2.1 Particle Swarm Optimization

Prior to discussing the details of PSO, a few terms are introduced in order to explain the inner working mechanism of the algorithm.

- Objective function: An function, \( f(x) \), that is to be optimized with respect to \( x \) within certain constraints.

- Search Space: A single or multi-dimensional space in which solutions are found based on the objective function, and the constraints.

- Particle: A member of the swarm, associated with a position vector, a velocity vector and a personal best vector.

- Swarm: A set of particles.

- Position: The location of a particle \( x \) within the search space, which represents a possible solution to the problem at hand. The position is updated using the following
equation,

\[ x(t + 1) = x(t) + v(t + 1) \] (2.1)

where, \( v \) is the velocity vector, and \( t \) is the iteration number.

- **Velocity:** The speed with which a particle moves within the search space. The velocity, \( v \), specifies the particle's course of movement. The velocity is updated using the following equation,

\[ V_i(t + 1) = \omega * V_i(t) + \phi_1 * (X_{pbest}(t) - X_i(t)) + \phi_2 * (X_{gbest}(t) - X_i(t)) \] (2.2)

Where,

\( \omega \) is the inertia weight

\( \phi_1 \) & \( \phi_2 \) is the accelerating factor

\((X_{pbest}(t) - X_i(t))\) is the cognitive term

\((X_{gbest}(t) - X_i(t))\) is the social term

- **Personal Best:** The position, \( p_{best} \), at which a particle achieved the best (maximum or minimum) objective function till a particular iteration. Each particle has a memory of its own personal best.

- **Global Best:** The best position vector, \( g \), out of all personal bests in the whole swarm.

The pseudocode of the PSO algorithm is presented next:
Algorithm 1 Basic PSO

1: for each particle do
    Randomly Initialize particle and velocity
2: end for
3: while maximum iterations or minimum error criteria is not attained do
4:     for each particle do
5:         Find the particle with the best fitness $p$ in the swarm
6:         Calculate particle velocity according to the equation 2.2
7:         Apply the velocity limit
8:         Update particle position according to the equation 2.1
9:         Apply the position limit
10:        Update velocity and position
11:     end for
12: end while
13: return $p$

The objective of PSO is to improve the position vectors and find a global best $g$ where

$$g = \text{argmin}(f(x)).$$

### 2.2 PSO and Data Clustering

An early PSO-based clustering algorithm was proposed by Van Der Merwe and Engelbrecht in 2003 [17]. In this algorithm, a data vector, $y$, is represented as an $M$-dimensional column vector. The $i$-th particle is represented by a group of column vectors, $x_i = \{p_{i1}, p_{i2}, ..., p_{ic}\}$, where $p_{in}$, $n = 1, 2, ..., n_c$, is a potential centroid vector, and $n_c$ denotes the number of centroids. The swarm’s initial position can be either determined by $k$-means or can be chosen randomly.

In [17], $k$-means was used to initialize the particles. After initialization, the particles obtained from $k$-means where added to a population of particles obtained randomly. Then, the algorithm applies generic PSO to find out which particle has the best distances from the
data points.

They evaluated PSO and the Hybrid clustering algorithm during the experiments. The algorithms were compared against $K$-means on six classification problems in [17]. Their clustering results were compared using three quality measures namely quantization error, intra-cluster distances and inter-cluster distances. The classification problems were composed by two bi-dimensional artificial and four well-known datasets.

The results of this experiments showed that the algorithms of [17], in general, are better than $k$-Means. Both proposed algorithms converge to lower quantization error in the first artificial problem. The Hybrid clustering algorithm has the smallest quantization error for all proposed datasets. Also, the Hybrid algorithm presents the smallest Inter-cluster Distances.

The pseudocode of the standard PSO data clustering algorithm presented in [17] is presented below:

<table>
<thead>
<tr>
<th>Algorithm 2</th>
<th>Standard Data Clustering PSO</th>
</tr>
</thead>
<tbody>
<tr>
<td>1: <strong>for</strong> each particle <strong>do</strong></td>
<td>randomly initialize particle and velocity</td>
</tr>
<tr>
<td>2: <strong>end for</strong></td>
<td></td>
</tr>
<tr>
<td>3: <strong>while</strong> maximum iterations/minimum error is not attained <strong>do</strong></td>
<td></td>
</tr>
<tr>
<td>4: <strong>for</strong> each particle <strong>do</strong></td>
<td></td>
</tr>
<tr>
<td>5: <strong>for</strong> each data point <strong>do</strong></td>
<td></td>
</tr>
<tr>
<td>6: Calculate distance between all data and centroids</td>
<td></td>
</tr>
<tr>
<td>7: Assign each data point to the closest centroid</td>
<td></td>
</tr>
<tr>
<td>8: <strong>end for</strong></td>
<td></td>
</tr>
<tr>
<td>9: Update global and local best positions</td>
<td></td>
</tr>
<tr>
<td>10: <strong>end for</strong></td>
<td></td>
</tr>
<tr>
<td>11: <strong>for</strong> each particle <strong>do</strong></td>
<td></td>
</tr>
<tr>
<td>12: Update velocity and position</td>
<td></td>
</tr>
<tr>
<td>13: <strong>end for</strong></td>
<td></td>
</tr>
<tr>
<td>14: <strong>end while</strong></td>
<td></td>
</tr>
<tr>
<td>15: <strong>return</strong> $p$</td>
<td></td>
</tr>
</tbody>
</table>
2.3 Particle Swarm Clustering

Cohen and de Castro proposed an improved way of clustering using PSO, and they named it Particle Swarm Clustering (PSC) [16]. The algorithm is evaluated with benchmark datasets and results are compared with that of standard unsupervised clustering algorithms. In PSC, PSO is modified to work with clustering and hence differs from PSO in a sense that unlike PSO, where each particle in the space leads to a potential solution and finally encodes the whole solution, PSC considers each particle as representation of clusters and thus encodes part of the solution. The performance evaluation measure in PSC differs from the fitness quality function used in the general PSO. Rather it initializes particles and then move these particles into such regions which shall represent natural clusters. Equation (2.3) represents a modified form of PSO and is the math equation used in PSC.

\[ V_i(t+1) = \omega V_i(t) + \phi_1 (X_{pbest}(t) - X_i(t)) + \phi_2 (X_{gbest}(t) - X_i(t)) + \phi_3 (y_j - x_i(t)) \]  

(2.3)

where,

\( v_i(t) \) is Particle’s previous position

\( p_i(t) - x_i(t) \) is the cognitive term

\( g_j(t) - x_i(t) \) is the social term

\( y_j(t) - x_i(t) \) is the self-organizing term

The particle swarm clustering algorithm has five main input parameters which are given in the following pseudo code. These are 1) the dataset which is going to be clustered, 2)
maximum number of iterations, 3) number of particles, 4) maximum velocity, and 5) initial value of the learning parameter.

```
1: procedure PSC
2:   for each particle do
3:     Randomly Initialize particle and velocity
4:   end for
5: while maximum iterations or minimum error criteria is not attained do
6:     Find the particle with the best fitness \( p \) in the swarm
7:   end while
8: return \( p \)
9: end procedure
```

Initially, the parameters of velocity and particles (cluster’s centroids) are randomly chosen, with maximum bound on the velocity values already defined. Then the distances between nearest data items are calculated and data items with minimum distance are grouped together to form clusters. After each iteration, the velocity, distance and centroids of clusters are updated until convergence.

The authors evaluated the algorithm using the Ruspini dataset which has 75 components in a two-dimensional space. The dataset was first normalized into the range of \([0,1]\). Then, the parameters were initialized and the PSC algorithm run for the specified number of iterations, which clustered the whole dataset correctly into four groups. It was shown that there were 20, 23, 17 and 15 data items into four clusters, respectively. The number of iterations were 150, and the maximum velocity was set to 0.001. The number of iterations, the maximum velocity, and the number of particles affected the overall performance of the process. Also, in a case where there are as many particles as data items, each particle represented an individual cluster. PSC was also evaluated on the Yeast dataset which has
29 data items, and the clustering performance is compared with \( k \)-means clustering. It was shown that PSC clusters the data more efficiently and addresses the problem of stagnation which other algorithms had.

Another concern that PSC handled is that data is sparse, and we often encounter cases where data items do not belong to any cluster. To handle this problem, PSC moves such data items in the direction of the cluster with the maximum number of data items. Apart from the number of iterations and maximum velocity parameters, the social and cognitive terms also affected clustering. So, instead of randomly initializing these terms, values were changed in a fixed interval as follows:

\[
\phi_1, \phi_2 \in [0.1, 2.05] \text{ and } \phi_3 \in [0.005, 1]
\]

To determine the effect of varying the above parameters in their specified range, all the other parameters are kept constant as given below:

- Number of particles were fixed to 8
- Maximum velocity was initialized to 0.01
- Number of iterations were initialized to 200
- Inertia weight \( \omega = 0.95 \)
- social \( \phi_1 \) and cognitive \( \phi_2 \) terms were fixed to 0
- Self-organizing terms \( \phi_3 = 0.005 \)

After running PSC, it was observed that many of the particle got stuck to their positions and become stationary, which affected the clustering performance. On the other hand, when
the social and cognitive terms were also introduced in the PSC algorithm, clustering was improved. Thus, it can be concluded that only introducing the self-organizing term and fixing social and cognitive terms to 0 leads to poor clustering results whereas introducing all of the three terms results in better clustering performance. This indicates that the social and cognitive terms contribute significantly in the clustering process. It was also concluded that the fixed initialization of all terms leads to poor clustering as opposed to random initialization.

2.4 Cooperative-Multipopulation Data Clustering PSO

Population-based data clustering algorithms employed in dynamic environments, lose the diversity of the population and the outdated memory of the individuals. Georgieva and Engelbrecht in [31] proposed a new particle swarm optimization alternative for clustering in dynamic environments, named cooperative-multipopulation data clustering PSO.

Most of the dynamic population-based clustering algorithms are variations or improvements of the static population-based algorithm of [17]. The new clustering algorithm of [31], cooperative-multipopulation data clustering PSO, is actually a combination of two algorithm named Multi-swarm Data Clustering PSO [32] and Cooperative Data Clustering PSO [33].

From the Multi-swarm Data Clustering PSO [32], the new algorithm takes the repulsion and anti-convergence methods. The repulsion method is the re-initialization of a swarm which is too close to another swarm. The anti-convergence method which re-initializes a swarm whose convergence radius is too small. The re-initialization method is the same as
proposed in [34], where a proportion of the population is placed in new random positions of the search space.

As for the Cooperative Data Clustering PSO [33], this new dynamic population-based algorithm takes each swarm that optimizes only one centroid. The other algorithms [32], [34] and [17] optimizes all centroid in each swarm. In order to be able to optimize all centroids, [33] proposed a context particle. The context particle is based on the best solutions of each swarm. Because of this context particle, the Cooperative Data Clustering PSO [33] is more viable than Multi-swarm Data Clustering PSO [32].

The cooperative-multipopulation data clustering PSO presented by [31] combines Cooperative Data Clustering PSO [33] (because of it’s quality) and Multi-swarm Data Clustering PSO [33] (because of it’s robustness). Furthermore, in order to avoid inaccurate representations of the solution, in [31] added an additional swarm, named the “explorer swarm”. The additional swarm substitutes a swarm that will be reinitialized. During this re-initialization, the re-initialized swarm becomes “explorer swarm” and the previous “explorer swarm” takes the place of the re-initialized swarm in the clustering solution. This additional swarm guarantee that the solution is given by algorithm’s exploration and is not random (given by re-initialization).

The algorithm of the cooperative-multipopulation data clustering PSO is performed using the following steps:

1. Initialize the particles of the $k + 1$ swarms (one additional as ”explorer swarm”).

2. Initialize the context particle.

3. Update each particle’s best position if the new position has less fitness value. The
fitness value is calculated putting the new particle in the context particle. Also, update the swarm best particle if some of the updated particles are better.

4. Update the context particle.

5. Update the position of all particles with the new context particle using standard PSO’s update equations.

6. Verify if two swarms are too close with a minimal inter-cluster distance. Re-initialize one of them in this case and change it by the “explorer swarm”.

7. Verify if a swarm converges too much with a minimal intra-cluster distance. Re-initialize this swarm in this case and change it by the “explorer swarm”.

8. Go to the step 3 until the stopping condition is not reached (max iteration steps). At the end, the resulting context particle is the output of this algorithm.

Then, in [31] an elitist version of their cooperative-multipopulation data clustering PSO algorithm was also proposed. The elitist version only changes the context particle to include new best solutions (not downgrade with re-initialization).

The dataset used for the experimental setup were auto-generated temporal clustering datasets with 8 clusters. Each cluster was changed in a dynamic way, moving patterns from one cluster to another at each interval of change. The algorithms were evaluated on a combination of change frequency between 1 and 5, severities in the same range and dimensions with 3, 8 and 15.

The quality of the algorithms was measured with inter-cluster distance, intra-cluster distance and Ray-turi validity index. The results were averaged over 20 runs where each
algorithm ran for 1000 iterations with 50 particles of swarm size. During the re-initialization, only 10% of the swarm was re-initialized.

The authors compared six data clustering PSO algorithms, the standard [17], the standard with re-initialization, Multi-swarm Data Clustering PSO [32], Cooperative Data Clustering PSO [33], cooperative-multipopulation data clustering PSO and it’s elitist version.

Between six population-based clustering algorithms evaluated on pattern migration datasets, according to the Ray-Turi measure, the elitist cooperative-multipopulation data clustering PSO gave the most optimal solution. However, this algorithm didn’t perform well with the increase of dimension. The cooperative-multipopulation algorithm didn’t present good results considering Ray-Turi and intra-cluster distance measures. But, the elitist version gave better results because it controls the effects of re-initialization on the context particle in a better way.
Chapter 3

Associative Data Clustering PSO

3.1 Improving Swarm Association

An issue with the basic data clustering PSO-based algorithm presented in [17] is the association problem that arises when calculating the square of the Euclidean distance, \( J_i \), between the centroid vectors within the \( i \)th particle, \( x_i \), and those within the global best particle, \( g_{best} \). More specifically, each particle should be represented as a set of centroids, in the sense that the order of centroid vectors within the particle should be of no importance. However, for the purpose of implementing \( J_i \), centroid vectors may be ordered. In [17], there is no indication that a solution to the association problem has been considered. The square distance, \( J_i \), is calculated using the following equation:

\[
J_i = \sum_{j=1}^{n_c} ||x_i^j - g_{best}^j||^2
\]  

(3.1)

where \( n_c \) is the number of centroids, and \( x_i^j \) and \( g_{best}^j \) represent the \( j \)th centroid within the particle and the global best particle, respectively. An example illustrating the aforementioned association problem is shown in Figure 3.1.

In Figure 3.1(a), an example of a particle and a global best particle are shown, each consisting of three centroids. The lines connecting the centroids are used to identify which
centroids are associated with the same particle. The numbers indicate the order of the centroid vector within the particle. In the case that a proper centroid association is not used, it is apparent that the particle will have to undergo a relatively large change in order to match the global best particle. This motion is indicated by the arrows in Figure 3.1(b). Nevertheless, the positions of the centroids within the two particles are in fact close to each other if an appropriate centroid associate association technique is implemented.

3.1.1 Data Clustering PSO with Association

In order to mitigate the association issue, a modification to [17] has been proposed. The overall modified algorithm is presented in Algorithm 3. In particular, the modified step is also described in more detail in Algorithm 4. Based on this modification, the global best solution corresponding to the $i^{th}$ particle is represented as $\hat{g}^i_{\text{best},i}$. Therefore, the equation for calculating the square Euclidean distance becomes as follows:

$$\hat{J}_i = \sum_{j=1}^{n_c} ||x^j_i - \hat{g}^j_{\text{best},i}||^2$$  \hspace{1cm} (3.2)
The algorithm for ADCPSO is shown in Algorithm 3 and the inner algorithm of the modified portion of DCPSO has been shown in Algorithm 4.

**Algorithm 3 Associative Data Clustering PSO**

1: for each particle do
    Randomly Initialize particle and velocity
2: end for
3: while maximum iterations/minimum error is not attained do
4:     for each particle do
5:         for each data point do
6:             Calculate distance between all data and centroids
7:             Assign each data point to the closest centroid
8:         end for
9:         Update global and local best positions
10:     end for
11:     for each particle do
12:         Determine distances between global best centroids and specific particle’s centroids
13:         Rearrange global best, $g_{best}$, centroids according to the distances between centroids
14:     end for
15: end while
16: return $p$

**Algorithm 4 Association Fix for Data Clustering PSO**

1: $n_c$ centroids
2: for each particle do
3:     Initialize all $n_c$ labels to 0
4:     for each centroid in the particle do
5:         if label is 0 then
6:             for each centroid in global best do
7:                 measure distance between particle centroid and global best centroid
8:         end for
9:     end if
10: append global best centroid with smallest distance to new global best
11: Set label to 1 for the chosen global best centroid
12: end for
13: Update velocity and position using the new global best
14: end for
3.1.2 Experimental Setup

A computer with Intel® Core i5-3230M with MATLAB R2016b has been used to simulate all experiments presented in the next section. Several standard datasets have been used in the experiments to evaluate the performance clustering algorithms. The details for the datasets used are mentioned in table 3.1. All of the datasets, except for the artificial dataset, are publicly available at UCI Machine Learning Repository [35].

Table 3.1: Datasets used for evaluation

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Objects</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ruspini</td>
<td>75</td>
<td>2</td>
</tr>
<tr>
<td>Artificial</td>
<td>200</td>
<td>2</td>
</tr>
<tr>
<td>Iris</td>
<td>150</td>
<td>4</td>
</tr>
<tr>
<td>Automotive</td>
<td>240</td>
<td>8</td>
</tr>
<tr>
<td>Sonar</td>
<td>208</td>
<td>60</td>
</tr>
</tbody>
</table>

The following were the parameters for the programs in MATLAB:

- Maximum Clusters: 3
- Particles: 20, 40, 60
- Inertia: 0.99
- Maximum Iterations: 500
- Program Evaluation: 500
- Accelerating factors: 0.5
- Dimensions: 2, 4, 8, 60
### 3.1.3 Results and Comparison

Tables 3.2-11 report the values of the best, average, median, and standard deviation of the global best cost found using 20, 40, and 60 particles, considering 500 runs of the algorithms. A total of five different datasets were used. In general, it can be observed that ADCPSO has achieved a lower cost against the DCPSO. Also, the standard deviation of the ADCPSO was lower than that of DCPSO, which indicates that the performance of DCPSO was more sensitive to particle initialization. Another important point is that the improvement of ADCPSO with respect to DCPSO increases as the dataset dimension increases. Table 3.16 reports the overall percentage improvement of ADCPSO against the DCPSO.

Table 3.12-15 presents the average execution time and number of iteration to converge required per run. It is apparent that ADCPSO has a disadvantage compared to DCPSO. Although the number of iterations required by ADCPSO in order to converge is in general smaller compared to DCPSO, its overall execution time is higher. The reason is that individual iterations of ASCPSO are more expensive computationally compared to DCPSO. For this reason, we have proposed another improved version of ADCPSO, namely IADCPSO. In particular, IADCPSO exhibits improved execution times, which comes at the expense of additional memory storage requirements. The extra memory storage is simply equal to the one required for storing the personal bests. The IADCPSO algorithm is described in section 3.2.
Table 3.2: Results for Ruspini dataset for the ADPSO algorithm

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Best</th>
<th>Average</th>
<th>Median</th>
<th>Std. Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>2.9782</td>
<td>3.0978</td>
<td>2.9843</td>
<td>0.1691</td>
</tr>
<tr>
<td>40</td>
<td>2.9781</td>
<td>3.0811</td>
<td>2.9806</td>
<td>0.1315</td>
</tr>
<tr>
<td>60</td>
<td>2.9781</td>
<td>3.084</td>
<td>2.9804</td>
<td>0.1323</td>
</tr>
</tbody>
</table>

Table 3.3: Results for Ruspini dataset for the DCPSO algorithm

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Best</th>
<th>Average</th>
<th>Median</th>
<th>Std. Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>2.9782</td>
<td>3.1022</td>
<td>2.986</td>
<td>0.1938</td>
</tr>
<tr>
<td>40</td>
<td>2.9781</td>
<td>3.0865</td>
<td>2.9815</td>
<td>0.1329</td>
</tr>
<tr>
<td>60</td>
<td>2.9781</td>
<td>3.0899</td>
<td>2.981</td>
<td>0.1336</td>
</tr>
</tbody>
</table>

Table 3.4: Results for Artificial dataset for the ADCPSO algorithm

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Best</th>
<th>Average</th>
<th>Median</th>
<th>Std. Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>7.7114</td>
<td>7.809</td>
<td>7.8189</td>
<td>0.0704</td>
</tr>
<tr>
<td>40</td>
<td>7.7114</td>
<td>7.793</td>
<td>7.793</td>
<td>0.067</td>
</tr>
<tr>
<td>60</td>
<td>7.7113</td>
<td>7.7945</td>
<td>7.813</td>
<td>0.0677</td>
</tr>
</tbody>
</table>

Table 3.5: Results for Artificial dataset for the DCPSO algorithm

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Best</th>
<th>Average</th>
<th>Median</th>
<th>Std. Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>7.7115</td>
<td>7.837</td>
<td>7.857</td>
<td>0.0798</td>
</tr>
<tr>
<td>40</td>
<td>7.7115</td>
<td>7.806</td>
<td>7.816</td>
<td>0.0692</td>
</tr>
<tr>
<td>60</td>
<td>7.7114</td>
<td>7.7997</td>
<td>7.814</td>
<td>0.0689</td>
</tr>
</tbody>
</table>

Table 3.6: Results for Iris dataset for the ADCPSO algorithm

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Best</th>
<th>Average</th>
<th>Median</th>
<th>Std. Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>2.8338</td>
<td>2.9043</td>
<td>2.8862</td>
<td>0.07113</td>
</tr>
<tr>
<td>40</td>
<td>2.8272</td>
<td>2.8524</td>
<td>2.8482</td>
<td>0.0169</td>
</tr>
<tr>
<td>60</td>
<td>2.8264</td>
<td>2.8415</td>
<td>2.839</td>
<td>0.0108</td>
</tr>
</tbody>
</table>
Table 3.7: Results for Iris dataset for the DCPSO algorithm

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Cost</th>
<th>Best</th>
<th>Average</th>
<th>Median</th>
<th>Std. Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td></td>
<td>2.8436</td>
<td>3.0371</td>
<td>2.995</td>
<td>0.2245</td>
</tr>
<tr>
<td>40</td>
<td></td>
<td>2.8296</td>
<td>2.9131</td>
<td>2.899</td>
<td>0.0563</td>
</tr>
<tr>
<td>60</td>
<td></td>
<td>2.8316</td>
<td>2.8814</td>
<td>2.873</td>
<td>0.0365</td>
</tr>
</tbody>
</table>

Table 3.8: Results for Automotive dataset for the ADCPSO algorithm

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Cost</th>
<th>Best</th>
<th>Average</th>
<th>Median</th>
<th>Std. Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td></td>
<td>15.6334</td>
<td>16.181</td>
<td>16.075</td>
<td>0.389</td>
</tr>
<tr>
<td>40</td>
<td></td>
<td>15.6081</td>
<td>15.7966</td>
<td>15.7705</td>
<td>0.1221</td>
</tr>
<tr>
<td>60</td>
<td></td>
<td>15.5913</td>
<td>15.7038</td>
<td>15.6946</td>
<td>0.0682</td>
</tr>
</tbody>
</table>

Table 3.9: Results for Automotive dataset for the DCPSO algorithm

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Cost</th>
<th>Best</th>
<th>Average</th>
<th>Median</th>
<th>Std. Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td></td>
<td>15.8136</td>
<td>16.7989</td>
<td>16.65</td>
<td>0.6707</td>
</tr>
<tr>
<td>40</td>
<td></td>
<td>15.68287</td>
<td>16.122</td>
<td>16.08</td>
<td>0.255</td>
</tr>
<tr>
<td>60</td>
<td></td>
<td>15.6323</td>
<td>15.9251</td>
<td>15.903</td>
<td>0.1536</td>
</tr>
</tbody>
</table>

Table 3.10: Results for Sonar dataset for the ADCPSO algorithm

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Cost</th>
<th>Best</th>
<th>Average</th>
<th>Median</th>
<th>Std. Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td></td>
<td>376.176</td>
<td>394.7668</td>
<td>393.0316</td>
<td>10.1231</td>
</tr>
<tr>
<td>40</td>
<td></td>
<td>369.7983</td>
<td>379.4235</td>
<td>378.5748</td>
<td>5.43123</td>
</tr>
<tr>
<td>60</td>
<td></td>
<td>368.4715</td>
<td>374.0484</td>
<td>373.3401</td>
<td>3.9172</td>
</tr>
</tbody>
</table>

Table 3.11: Results for Sonar dataset for the DCPSO algorithm

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Cost</th>
<th>Best</th>
<th>Average</th>
<th>Median</th>
<th>Std. Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td></td>
<td>383.6466</td>
<td>404.0151</td>
<td>403.3845</td>
<td>9.5313</td>
</tr>
<tr>
<td>40</td>
<td></td>
<td>373.7812</td>
<td>386.0607</td>
<td>385.2033</td>
<td>6.0774</td>
</tr>
<tr>
<td>60</td>
<td></td>
<td>370.8911</td>
<td>378.2333</td>
<td>377.9088</td>
<td>3.5355</td>
</tr>
</tbody>
</table>
Table 3.12: ADCPSO run time

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Average time of each iteration (seconds)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ruspini</td>
</tr>
<tr>
<td>20</td>
<td>0.266</td>
</tr>
<tr>
<td>40</td>
<td>0.448</td>
</tr>
<tr>
<td>60</td>
<td>0.586</td>
</tr>
</tbody>
</table>

Table 3.13: DCPSO run time

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Average time of each iteration (seconds)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ruspini</td>
</tr>
<tr>
<td>20</td>
<td>0.194</td>
</tr>
<tr>
<td>40</td>
<td>0.301</td>
</tr>
<tr>
<td>60</td>
<td>0.406</td>
</tr>
</tbody>
</table>

Table 3.14: ADCPSO average number of iterations for convergence

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Average no. of iterations to converge</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ruspini</td>
</tr>
<tr>
<td>20</td>
<td>150</td>
</tr>
<tr>
<td>40</td>
<td>109</td>
</tr>
<tr>
<td>60</td>
<td>96</td>
</tr>
</tbody>
</table>

Table 3.15: ADCPSO average number of iterations for convergence

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Average no. of iterations to converge</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ruspini</td>
</tr>
<tr>
<td>20</td>
<td>193</td>
</tr>
<tr>
<td>40</td>
<td>131</td>
</tr>
<tr>
<td>60</td>
<td>112</td>
</tr>
</tbody>
</table>

Table 3.16: Improvement achieved by ADCPSO against DCPSO (%) 

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Best Cost Improvement (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ruspini</td>
</tr>
<tr>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>40</td>
<td>0</td>
</tr>
<tr>
<td>60</td>
<td>0</td>
</tr>
</tbody>
</table>
3.2 Improved ADCPSO (IADCPSO)

3.2.1 Reducing Execution Time

The drawback of ADCPSO is that the program runs take longer than DCPSO due to its higher computational complexity. To remedy this issue, an additional modification to the algorithm has been proposed, so that the association part of the algorithm runs only when there is a change in the global best solution. In order to achieve this objective, all of the updated global best vectors, $\hat{g}_{best,i}^j$, are stored as separate variables. Therefore, there will be one global vector stored per particle. The additional memory requirements equal the memory needed to store the personal best vectors for all particles. Essentially, there is a trade off between memory and execution time.

3.2.2 Results and Comparison

Tables 3.17-19 represent the best cost achieved by IADCPSO, the average time for each iteration, and the average number of iterations needed for convergence. The results demonstrate that IADCPSO exhibits almost the same cost as ADCPSO, while reducing the execution time close to that of the DCPSO algorithm.

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Ruspini</th>
<th>Artificial</th>
<th>Iris</th>
<th>Automotive</th>
<th>Sonar</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>2.978</td>
<td>7.7115</td>
<td>2.832</td>
<td>15.683</td>
<td>373.925</td>
</tr>
<tr>
<td>40</td>
<td>2.978</td>
<td>7.7114</td>
<td>2.828</td>
<td>15.592</td>
<td>370.53</td>
</tr>
<tr>
<td>60</td>
<td>2.978</td>
<td>7.7113</td>
<td>2.827</td>
<td>15.586</td>
<td>368.1</td>
</tr>
</tbody>
</table>
Table 3.18: Execution time for IADCPSO

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Average time of each iteration (seconds)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ruspini</td>
</tr>
<tr>
<td>20</td>
<td>0.161</td>
</tr>
<tr>
<td>40</td>
<td>0.265</td>
</tr>
<tr>
<td>60</td>
<td>0.364</td>
</tr>
</tbody>
</table>

Table 3.19: IADCPSO average number of iterations needed for convergence

<table>
<thead>
<tr>
<th>No. of Particles</th>
<th>Average no. of iterations to converge</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ruspini</td>
</tr>
<tr>
<td>20</td>
<td>149</td>
</tr>
<tr>
<td>40</td>
<td>110</td>
</tr>
<tr>
<td>60</td>
<td>96</td>
</tr>
</tbody>
</table>
Chapter 4

Conclusions and Future Work

4.1 Primary Findings

This research aimed to improve a PSO based data clustering algorithm. One major issue with several existing PSO-based clustering algorithms was identified. The issue was the association of particle centroids with the centroids of the global best particle when computing the cognitive term. We have proposed a modification to the PSO clustering algorithm in [17]. We performed comparisons between the proposed ADCPSO and the existing DCPSO. In summary, ADCPSO showed slight to moderate improvement, in terms of the cost, for datasets with dimensions higher than 3. Also, the proposed algorithm converged faster than the original DCPSO, in terms of the number of iterations. The only drawback was that the execution time per iteration was higher for ACPSO. This was due to the fact that individual iterations required additional computations for performing centroid associations between particles and the global best vector. This concern was also mitigated in this thesis in an second modified algorithm, namely IADCPSO.
4.2 Recommendations for Future Work

As this thesis focused on the use of PSO to cluster datasets, various future work can be done based on this research:

- Applying the association operation between the particles and their personal best, in addition to the association operation between the particles and the global best,

- Studying the scalability of the algorithms,

- Comparing the performance of IADCPSO with additional existing PSO-based clustering algorithms,

- Implementing advanced PSO-based dimensional clustering algorithms, similar to the one proposed in [36]
Bibliography


A.1 Data Clustering PSO

```matlab
for p=1:500

data_set=textread('data_set_sonar.txt');  \% Import Dataset

dataset_subset=0;
n_dimensions=60;
max_iter=500;
vmax=0.01;
w=0.99;

global Y

phi1=0.5;  \% 1.1;
phi2=0.5;  \% 0.8;
phi3=0.005;  \% 0.3;
```
phi4 = 0.06;

% Y = rand(length(Y), 2);
N = max(size(data_set));
Y = zeros(N, n_dimensions);
for i = 1:n_dimensions
    Y(:, i) = data_set(:, i) / max(data_set(:, i));
end
n_centroids = 3;
n_particles = 60;

tic
v = cell(n_particles, 1);

for i = 1:n_particles
    v{i} = vmax * (2 * rand(n_centroids, n_dimensions) - 1);
end

x = cell(n_particles, 1);
for i = 1:n_particles
    for j = 1:n_centroids
        r = ceil(rand(1, 1) * (length(Y) - 1) + 1);
        x{i}(j, :) = Y(r, :);
    end
end

motion_iter = [];
personal_best_cost = linspace(1e99, 1e99, n_particles);
personal_best = x;
global_best_cost = 1e99;
global_best = inf(n_centroids, n_dimensions);

prev_global_best_cost = inf;
iter = 0;
run_flag = 1;

while (iter < max_iter && run_flag == 1)
    run_flag = 0;
    for iter_inner = 1:50
        iter = iter + 1;
        % Particle loop
        for i = 1:n_particles
            % Centroid loop
            for j = 1:n_centroids
                Distance{i}(j,:) = (sum((repmat(x{i}(j,:),[N ... 1])-Y).^2,2))';
            end
        end
        for i = 1:n_particles
            [minimum_dist_temp, closest_centroid_temp] = ...
            min(Distance{i},[],1);
            minimum_dist(i,:) = minimum_dist_temp;
            closest_centroid(i,:) = closest_centroid_temp;
        end
        total_minimum = sum(minimum_dist,2);
    end

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prev_global_best_cost = global_best_cost;

for i=1:n_particles
    if total_minimum(i) < personal_best_cost(i)
        personal_best_cost(i) = total_minimum(i);
        personal_bests{i} = x{i};
    end

    if total_minimum(i) < global_best_cost
        global_best_cost = total_minimum(i);
        global_best = x{i};
    end
end

for i=1:n_particles
    v{i} = v{i} ...
        + phi1*(personal_bests{i} - x{i}) + ...
        phi2*(global_best - x{i});
    x{i} = x{i}+w*v{i};
end

w = w*0.99;

if (abs(global_best_cost - prev_global_best_cost) > 0.001)
A.2 Data Clustering PSO with $k$-means

```matlab
for p=1:500
    data_set=textread('data_set_sonar.txt'); % Ruspini Dataset
    dataset_subset=0;
    n_dimensions=60;
    max_iter=500;
    vmax=0.01;
    w=0.99;
    global Y
    phi1=0.5; % 1.1;
    phi2=0.5; % 0.8;
```
phi3=0.005; % 0.3;
phi4=0.06;
N=max(size(data_set));
Y=zeros(N,n_dimensions);
for i=1:n_dimensions
    Y(:,i)=data_set(:,i)/max(data_set(:,i));
end
n_centroids=3;
n_particles=60;
tic
v=cell(n_particles,1);
for i=1:n_particles
    v{i}=vmax*(2*rand(n_centroids,n_dimensions)-1);
end
x=cell(n_particles,1);
for i=1:n_particles
    for j=1:n_centroids
        r=ceil(rand(1,1)*(length(Y)-1)+1);
        x{i}(j,:)=Y(r,:);
    end
end
motion_iter=[];
personal_best_cost=linspace(1e99,1e99,n_particles);
personal_bests= x;
global_best_cost=1e99;
global_best= inf(n_centroids,n_dimensions);

prev_global_best_cost=inf;
iter=0;
run_flag=1;

while (iter<max_iter && run_flag==1)
    run_flag=0;
    for iter_inner=1:50
        iter=iter+1;
        % Particle loop
        for i=1:n_particles
            % Centroid loop
            for j=1:n_centroids
                Distance{i}(j,:)=(sum((repmat(x{i}(j,:),[N ... 1])-Y).^2,2)).';
            end
        end
        for i=1:n_particles
            [minimum_dist_temp, closest_centroid_temp]= ... 
            min(Distance{i},[],1);
            minimum_dist(i,:)=minimum_dist_temp;
            closest_centroid(i,:)=closest_centroid_temp;
        end
        total_minimum=sum(minimum_dist,2);
    end
prev_global_best_cost=global_best_cost;

for i=1:n_particles
    if total_minimum(i) < personal_best_cost(i)
        personal_best_cost(i)=total_minimum(i);
        personal_bests{i} = x{i};
    end

    if total_minimum(i) < global_best_cost
        global_best_cost=total_minimum(i);
        global_best = x{i};
    end

end

for i=1:n_particles
    v{i} = v{i} + phi1*(personal_bests{i} - x{i}) + ...
          phi2*(global_best - x{i});
    x{i} = x{i}+w*v{i};
end

w=w*0.99;

if (abs(global_best_cost-prev_global_best_cost)>0.001)
    c_iter=iter;
    run_flag=1;
end
A.3 Associative DCPSO

```matlab
end
end
time=toc;
for i=1:3
    [indices, global_best, SumD] = kmeans(Y,3,'start',global_best);
global_best_cost_kmeans=sum(SumD);
end
```
phi2=0.5;  \% 0.8;
phi3=0.005;  \% 0.3;
phi4=0.06;
N=max(size(data_set));
Y=zeros(N,n_dimensions);
for i=1:n_dimensions
    Y(:,i)=data_set(:,i)/max(data_set(:,i));
end

n_centroids=3;
n_particles=60;

tic
v=cell(n_particles,1);
for i=1:n_particles
    v{i}=vmax*(2*rand(n_centroids,n_dimensions)-1);
end

x=cell(n_particles,1);
for i=1:n_particles
    for j=1:n_centroids
        r=ceil(rand(1,1)*(length(Y)-1)+1);
        x{i}(j,:)=Y(r,:);
    end
end
motion_iter=[];

personal_best_cost=linspace(1e99,1e99,n_particles);

personal_bests= x;

global_best_cost=1e99;

global_best= inf(n_centroids,n_dimensions);

prev_global_best_cost=inf;

iter=0;

run_flag=1;

while (iter<max_iter && run_flag==1)

run_flag=0;

for iter_inner=1:50

iter=iter+1;

% Particle loop

for i=1:n_particles

% Centroid loop

for j=1:n_centroids

Distance{i}(j,:)=(sum((repmat(x{i}(j,:),[N ...

1])-Y).^2,2)).';

end

end

for i=1:n_particles

[minimum_dist_temp, closest_centroid_temp]= ...

min(Distance{i},[],1);

minimum_dist(i,:)=minimum_dist_temp;

closest_centroid(i,:)=closest_centroid_temp;

end
total_minimum=sum(minimum_dist,2);

prev_global_best_cost=global_best_cost;

for i=1:n_particles
    if total_minimum(i) < personal_best_cost(i)
        personal_best_cost(i)=total_minimum(i);
        personal_best {i} = x{i};
    end

    if total_minimum(i) < global_best_cost
        global_best_cost=total_minimum(i);
        global_best = x{i};
    end
end

gtemp=global_best*0;

for i=1:n_particles
    label = zeros(1,n_centroids);
    for j=1:n_centroids
        d1=inf;
        for c=1:n_centroids
            if (label(c)==0)
                d=sum((x{i}(j,:)-global_best(c,:)).^2);
                if (d<d1)
                    ctemp=c;
                    d1=d;
                end
            end
        end
    end
end
end
end
gtemp(j,:) = global_best(ctemp,:);
label(:,ctemp) = 1;
end

v{i} = v{i} ...
    + phi1*(personal_bests{i} - x{i}) + phi2*(gtemp - ...
x{i});
x{i} = x{i} + w*v{i};
end
w = w*0.99;

if (abs(global_best_cost - prev_global_best_cost) > 0.001)
c_iter = iter;
run_flag = 1;
end
end
end
time = toc;
end
A.4 ADCPSO with $k$-means

for $p=1:500$

    data_set=textread('data_set_sonar.txt');
    dataset_subset=0;
    n_dimensions=60;
    max_iter=500;
    vmax=0.01;
    w=0.99;
    global Y

    phi1=0.5;  \% 1.1;
    phi2=0.5;  \% 0.8;
    phi3=0.005;  \% 0.3;
    phi4=0.06;
    N=max(size(data_set));
    Y=zeros(N,n_dimensions);
    for $i=1:n\_dimensions$
        Y(:,i)=data_set(:,i)/max(data_set(:,i));
    end
    n_centroids=3;


n_particles=60;

tic

v=cell(n_particles,1);

for i=1:n_particles
    v{i}=vmax*(2*rand(n_centroids,n_dimensions)-1);
end

x=cell(n_particles,1);
for i=1:n_particles
    for j=1:n_centroids
        r=ceil(rand(1,1)*(length(Y)-1)+1);
        x{i}(j,:)=Y(r,:);
    end
end

motion_iter=[];
personal_best_cost=linspace(1e99,1e99,n_particles);
personal_bests=x;
global_best_cost=1e99;
global_best=inf(n_centroids,n_dimensions);

prev_global_best_cost=inf;
iter=0;
run_flag=1;
while (iter<max_iter & run_flag==1)
run_flag=0;
for iter_inner=1:50
    iter=iter+1;
    % Particle loop
    for i=1:n_particles
        % Centroid loop
        for j=1:n_centroids
            Distance{i}(j,:)= (sum((repmat(x{i}(j,:),[N ... 1])-Y).^2,2)).';
        end
    end
    for i=1:n_particles
        [minimum_dist_temp, closest_centroid_temp] = ...
        min(Distance{i},[],1);
        minimum_dist(i,:) = minimum_dist_temp;
        closest_centroid(i,:) = closest_centroid_temp;
    end
    total_minimum = sum(minimum_dist,2);
    prev_global_best_cost = global_best_cost;
    for i=1:n_particles
        if total_minimum(i) < personal_best_cost(i)
            personal_best_cost(i) = total_minimum(i);
            personal_bests{i} = x{i};
        end
        if total_minimum(i) < global_best_cost
            global_best_cost = total_minimum(i);
        end
    end
global_best = x{i};

end

gtemp=global_best*0;

for i=1:n_particles

    label = zeros(1,n_centroids);
    for j=1:n_centroids
        d1=inf;
        for c=1:n_centroids
            if (label(c)==0)
                d=sum((x{i}(j,:)-global_best(c,:)).^2);
                if (d<d1)
                    ctemp=c;
                    d1=d;
                end
            end
        end
    end
    gtemp(j,:)=global_best(ctemp,:);
    label(:,ctemp)=1;
end

v{i} = v{i} ...
\[ x(i) = x(i) + \phi_1 \cdot (\text{personal_bests}_i - x(i)) + \phi_2 \cdot (\text{gtemp} \ldots x(i)); \]
\[ x(i) = x(i) + w \cdot v_{\{i\}}; \]

\text{end}

\[ w = w \cdot 0.99; \]

\textbf{if} (abs(\text{global_best_cost} - \text{prev_global_best_cost}) > 0.001)

\[ \text{c_iter} = \text{iter}; \]
\[ \text{run_flag} = 1; \]
\text{end}

\text{end}

\text{end}

\text{time} = \text{toc};

\textbf{for} \ i = 1:3

\[ [\neg, \text{global_best}, \text{SumD}] = \text{kmeans}(Y, 3, '\text{start'}, \text{global_best}); \]
\[ \text{global_best_cost}_k\text{means} = \text{sum}(\text{SumD}); \]
\text{end}

\text{end}

\section*{A.5 Improved ADCPSO}
for p=1:500

data_set=textread('data_set_sonar.txt');
dataset_subset=0;
n_dimensions=60;
max_iter=500;
vmax=0.01;
w=0.99;
global Y

phi1=0.5; % 1.1;
phi2=0.5; % 0.8;
phi3=0.005; % 0.3;
phi4=0.06;
N=max(size(data_set));
Y=zeros(N,n_dimensions);
for i=1:n_dimensions
    Y(:,i)=data_set(:,i)/max(data_set(:,i));
end

n_centroids=3;
n_particles=60;
v = cell(n_particles, 1);

for i = 1 : n_particles
    v{i} = vmax * (2 * rand(n_centroids, n_dimensions) - 1);
end

x = cell(n_particles, 1);
for i = 1 : n_particles
    for j = 1 : n_centroids
        r = ceil(rand(1, 1) * (length(Y) - 1) + 1);
        x{i}(j,:) = Y(r,:);
    end
end

motion_iter = [];
personal_best_cost = linspace(1e99, 1e99, n_particles);
personal_bests = x;
global_best_cost = 1e99;
global_best = inf(n_centroids, n_dimensions);
global_bests = x;

prev_global_best_cost = inf;
iter = 0;
run_flag = 1;
while (iter < max_iter && run_flag == 1)
    run_flag = 0;
    for iter_inner = 1 : 50
iter=iter+1;

% Particle loop
for i=1:n_particles

% Centroid loop
for j=1:n_centroids
    Distance{i}(j,:)=sum((repmat(x{i}(j,:),[N ... 1])-Y).^2,2)).';
end
end

for i=1:n_particles
    [minimum_dist_temp, closest_centroid_temp]= ... 
        min(Distance{i},[],1);
    minimum_dist(i,:)=minimum_dist_temp;
    closest_centroid(i,:)=closest_centroid_temp;
end

total_minimum=sum(minimum_dist,2);

best_updated=0;
prev_global_best_cost=global_best_cost;
for i=1:n_particles
    if total_minimum(i) < personal_best_cost(i)
        personal_best_cost(i)=total_minimum(i);
        personal_bests{i} = x{i};
    end
    if total_minimum(i) < global_best_cost
        global_best_cost=total_minimum(i);
        global_best = x{i};
    end
end
best_updated=1;

end

end
gtemp=global_best*0;

for i=1:n_particles
  if best_updated==1
    label = zeros(1,n_centroids);
    for j=1:n_centroids
d1=inf;
      for c=1:n_centroids
        if (label(c)==0)
          d=sum((x{i}(j,:)-global_best(c,:)).^2);
          if (d<d1)
            ctemp=c;
            d1=d;
          end
        end
      end
gtemp(j,:)=global_best(ctemp,:);
    label(:,ctemp)=1;
  end
  global_best{i}=gtemp;
end

v{i} = v{i} ...
\begin{verbatim}
phi1*(personal_best{i} - x{i}) + ...
phi2*(global_best{i} - x{i});

x{i} = x{i}+w*v{i};

end

w=w*0.99;

if (abs(global_best_cost-prev_global_best_cost)>0.001)

c_iter=iter;

run_flag=1;

end

end

end

time=toc;
end
\end{verbatim}

A.6 Particle Swarm Clustering

\begin{verbatim}
%% Initialization

data_set=textread('data_set_sonar.txt');

max_it=150;

vmax=0.01;

n_part=3;

w=0.95;

n_cluster_per_particle=3;
\end{verbatim}
global Y

phi1=0; \% 1.1;
phi2=0; \% 0.8;
phi3=0.005; \% 0.3;
phi4=0.06;

Y = [data_set(:,1)/max(data_set(:,1)) ... 
     data_set(:,2)/max(data_set(:,2))];
N=max(size(data_set));
x=rand(n_cluster_per_particle*n_part,2); \% usually every particle ...
   xi initialized at random
c = reshape(x,1, []);

v=vmax*(2*rand(n_part*n_cluster_per_particle,2)-1); \% at random, vi ...
   in [-vmax,vmax]
distYX=linspace(1e99,1e99,n_part);
pI=[0 0];
pG=[0 0];
win_counter=zeros(max(size(x)),1);
t=1;

while t < max_it
    win_counter=zeros(n_part*n_cluster_per_particle,1);
    for i = 1 : N \%for each data
        for j=1:n_part*n_cluster_per_particle
            distYX(j) = dist(Y(i,:),x(j,:))';
end

distMatrix(:,i)=distYX;
for k=1:n_cluster_per_particle
  particle=mat2cell(distMatrix,[3 3 3], N);
end

I = find(distYX==min(distYX));
win_counter(I)= win_counter(I)+1;

f= @(x) sum(sum(bsxfun(@minus,Y,x).^2)); % distance function
if f(x(I,:)) < f(pI)
  pI = x(I,:);
end
if f(x(I,:)) < f(pG)
  pG = x(I,:);
end

v(I,:) = min(vmax, max(-vmax, v(I,:) + phi1*(pI - x(I,:)) ...
  + phi2*(pG - x(I,:)) + phi3*(Y(i,:) - x(I,:))));
x(I,:)=x(I,:)+w*v(I,:);
end

% for each empty particles, redirect them to the winning particles
[max_count]=max(win_counter);
for i = 1 : n_part*n_cluster_per_particle
  if(win_counter(i)==0)
    v(i,:) = min(vmax, max(-vmax, w*v(i,:) + ...
      phi4*(x(max_count,:) - x(i,:))));
    x(i,:) = x(i,:) + v(i,:);
  end
end
end

end

\( w = 0.95 \times w; \)

\( t = t + 1; \)

end

disp(x)
Appendix B

Acronyms

Following is the list of acronyms used in this thesis. Acronyms are ordered alphabetically.

ACO  Ant Colony Optimization
ADCPSO  Associative Data Clustering Particle Swarm Optimization
DCPSO  Data Clustering Particle Swarm Optimization
EA  Evolutionary Algorithm
GA  Genetic Algorithm
IADCPSO  Improved Associative Data Clustering Particle Swarm Optimization
PSC  Particle Swarm Clustering
PSO  Particle Swarm Optimization
SI  Swarm Intelligence
Vita

Sharif Shahadat was born on October 19, 1990, in Dhaka, Bangladesh. The author completed his bachelor’s degree in Electrical and Electronic Engineering from Ahsanullah University of Science and Technology, Dhaka, Bangladesh in 2012. He finished his Masters in Electrical Engineering from the University of New Orleans in May 2017 with a cumulative GPA of 3.60. The author worked with his advisor Dr. Dimitrios Charalampidis for his master’s thesis. His research interests are in Data clustering, Particle Swarm Optimization and Metaheuristics.