ECA-CE: An Evolutionary Clustering Algorithm with Initial Population by Clustering Ensemble

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Abstract: Evolutionary clustering is a type of algorithm that uses genetic algorithms to optimize clustering results. Unlike traditional clustering algorithms which obtain clustering results by iteratively increasing the distance between clusters and reducing the distance between instances within a cluster, the evolutionary clustering algorithm tries to search for the optimal clustering result in the solution space. Not surprisingly, the initial population set in an evolutionary clustering algorithm has significant influence on the final results. To ensure the quality of the initial population, this paper proposed a clustering ensemble-based method, ECA-CE, to do the initial population for the evolutionary clustering algorithm. In ECA-CE, a clustering ensemble method, Hybrid Bipartite Graph Formulation, is applied. Extensive experiments are conducted on 20 benchmark datasets, and the experimental results demonstrate that the proposed ECA-CE is more effective than two evolutionary clustering algorithms F1-ECAC and ECAC in terms of Adjusted Rand index.

1 INTRODUCTION

In most real-world situations, data is unlabeled. Clustering is an unsupervised learning algorithm that takes an unlabeled dataset as input and divides it into a certain number of clusters, where data with similar characteristics are grouped. The ideal clustering result should be that all the clusters generated are independent of each other and data within a cluster are relatively compact.

Traditional clustering algorithms can be mainly divided into three groups, including partition-based, hierarchy-based, and graph-theory-based clustering algorithms. The main idea of the partition-based algorithm is to discover the groupings in the data by optimizing a specific objective function and iteratively improving the quality of partitions (Kang et al., 2019). The shortcoming of partition-based clustering algorithms is that they can easily be trapped in a local optimum. Hierarchy-based clustering algorithms divide data at different levels to form a tree-like structure (Nielsen, 2016). The graph-based clustering algorithms divide data at different levels to form a tree-like structure (Von Luxburg, 2007) construct an undirected graph with similarity weights defined in a matrix and then apply a clustering algorithm to partition the undirected graph.

Different from those traditional methods, evolutionary clustering algorithms utilize genetic algorithms to search for the global optimal clustering results. F1-ECAC (Sainz-Tinajero et al., 2021b) and ECAC (Sainz-Tinajero et al., 2021a) are two recently proposed evolutionary clustering algorithms that achieve good performance. However, one problem of F1-ECAC and ECAC is that the quality of the initial population is unstable, which impacts the final clustering results. If the evolutionary clustering algorithm cannot start with a good initial population, it will require more time to obtain the optimal clustering result, or cannot find that at all.

To alleviate the above problem, this paper proposes an evolutionary clustering algorithm, ECA-CE (Evolutionary Clustering Algorithm with initial population by Clustering Ensemble), whose initial population is produced by a clustering ensemble and employs multiple supervised classifiers to evaluate each individual’s fitness. Our experimental results show that the proposed method achieves better performance than F1-ECAC and ECAC do.

The rest of the paper is organized as follows: Section II summarizes related work. Section III describes ECA-CE. The experimental settings and results for the evaluation of the proposed method are presented in Section IV. Finally, a summary is given in Section V.
2 RELATED WORK

2.1 Traditional Clustering Algorithms

There are several types of traditional clustering algorithms.

1. Partition-based algorithm: For a given initial central point, a heuristic search method is used to iteratively update each cluster and its centroid until the distance between any two instances within a cluster is small enough and the distance between clusters is large enough.

2. Hierarchy-based algorithm: To create clusters in a hierarchical tree-like structure in which the root node corresponds to the entire dataset, and branches are created from the root node to form hierarchical clusters. Mainly there are two types of hierarchy-based clustering algorithms: fused hierarchical clustering and partitioned hierarchical clustering. The fused hierarchical clustering algorithm generates all the nodes in a bottom-up style, while the latter does it in a top-down style.

3. Graph-based algorithm: Each data point in a dataset is represented as a node, the distance between two data points is modeled by an edge between the two nodes with a weight. In this way, the clustering problem can be transformed into a graph partition problem, and the clustering results can be obtained by applying some partition criteria such as minimum cut on the graph.

In this study, we choose a typical method from each category as the baseline for performance comparison. Also they are used as base clustering algorithms for the clustering ensemble.

a) K-means: It is a well-known partition-based method with multiple variants, such as k-medoids, k-medians, etc. K-means is a centroid-based clustering algorithm. This method tries to shorten the average Euclidean distance from each data point to the centroid of the cluster through iterative membership change. Since K-means intends to build sufficiently tight clusters, the algorithm is likely to fall into local optima and is exceptionally sensitive to noisy data points.

b) BIRCH: It constructs a cluster feature tree (CFT) for the dataset. Since the algorithm uses a tree model, then it is well-interpretable and can solve non-spherical problems that cannot be solved by K-means. However, the construction of the tree structure leads to high time complexity. Therefore, BIRCH may not be able to handle high-dimensional data.

c) Spectral Clustering: Spectral Clustering is a graph-based algorithm. Spectral Clustering requires a similarity matrix between instances to work. It is effective in dealing with sparse datasets. Due to the use of dimensionality reduction, it is a more appropriate solution for high-dimensional datasets than BIRCH. However, as Spectral Clustering relies on similarity matrices, different similarity matrices may yield different clustering results.

Algorithm 1: HBGF.

Require: dataset \( D = \{X_1, X_2, \ldots, X_n\} \), \( X_i = \{x_{i1}, x_{i2}, \ldots, x_{im}\} \) \( \triangleright \) \( n \) is the number of instances and \( m \) is the number of features.

graph partitioning algorithm package \( L(SPEC \ (Shi \ and \ Malik, \ 2000) \ or \ METIS \ (Karypis \ and \ Kumar, \ 1998)) \)

1: \( V = D \cup C \); \( \triangleright \) \( C \) contains all clusters \( C = \{e_j | 1 \leq j \leq k^*\} \), \( k^* \) is the number of clusters
2: \( E = \emptyset \);
3: for \( i = 1, \ldots, n \) do:
4: for \( j = 1, \ldots, k^* \) do:
5: if \( v' \in v^i \) then: \( \triangleright \) \( v' \) is an instance. \( v^j \) is a cluster
6: \( E = E \cup \{e_{ij}\} \) \( \triangleright \) add edge \( e_{ij} = (v^i, v^j) \);
7: \( w_{ij} = 1 \) \( \triangleright \) Set equal weights for \( e_{ij} \);
8: end if
9: end for
10: end for
11: \( G = (V, E) \);
12: \( \sigma = L(G) \); \( \triangleright \) Apply the specified graph partitioning algorithm package on \( G \)

2.2 Evolutionary Clustering Algorithm

The evolutionary clustering algorithm generally applies the genetic search method to try to find the optimal results. In order to solve the density-related clustering problem, Zhang et al. (Zhang et al., 2013) proposed an evolutionary clustering algorithm based on DBSCAN (Ester et al., 1996), and applied the time smoothing degree penalty framework in the calculation process. Sainz Tinajero et al. (Sainz-Tinajero et al., 2021a) proposed a single objective evolutionary clustering algorithm (ECAC) based on a supervised classifier, which uses random functions to initialize the population of solutions. After performing selection, crossover, and mutation operations, a new population is obtained. For the evaluation of a clustering solution, we transform the solution to a classification problem, in which each instance is composed of all the original attributes with a label corresponding to a specific cluster. In order to let a classifier
work, they divide all the instances into two partitions: Training and testing partitions. The training part is used to train the classifier, and then the trained model is applied to the testing part. Its AUC index is used as the fitness metric of the clustering results. Multiple supervised classifiers are used to make the evaluation results more reliable. The best solution is kept in the iterative process. The iteration stops when the ending condition, either a predefined number of iterations or a given AUC value, is satisfied. The author subsequently proposed F1-ECAC (Sainz-Tinajero et al., 2021b), a variant of ECAC, which uses F1 score to replace AUC as the evaluation index.

Additionally, Mardi and Keyvanpour (Mardi and Keyvanpour, 2021) applied the evolutionary clustering algorithm to set the initial points for the K-means algorithm, which they believe can reduce the sensitivity of K-means to abnormal data.

3 PROPOSED ALGORITHM

3.1 ECA-CE

In this section, we describe ECA-CE in detail. We will also discuss Hybrid Bipartite Graph Formulation (HBGF) (Fern and Brodley, 2004), a clustering ensemble algorithm, and how to use HBGF in ECA-CE.

HBGF, a graph-based clustering ensemble algorithm, was proposed by Fern et al. The pseudo-code for the HBGF algorithm is shown in Algorithm 1. It is expected that the clustering results generated by an ensemble of multiple clustering algorithms are more stable.

In ECA-CE, firstly we run three traditional clustering algorithms including the aforementioned K-means, BIRCH, and Spectral Clustering algorithm to obtain three different clustering results. Then HBGF is used to fuse them, and the fused clustering results are taken as an initial solution for the evolutionary clustering algorithm. We think that such an initial solution can help the evolutionary clustering algorithm to find better solutions more quickly. Algorithm 2 shows the process of the ECA-CE algorithm. For a dataset $D = \{X_1, X_2, ..., X_n\}$, $X_i = \{x_{i1}, x_{i2}, ..., x_{in}\}$, where $n$ denotes the number of instances and $m$ denotes the number of features. ECA-CE calculates the individual’s fitness in a supervised manner (Caruana and Niculescu-Mizil, 2006).

Algorithm 2: ECA-CE.

Require: dataset $D$, the population size $\alpha$, number of iterations $\beta$, number of clusters $k$.
1: $\theta = \{g_1, g_2, g_3, ..., g_j, ..., g_n\} = HBGF(D)$;
2: $f_{\text{max}} = 0$;
3: bestCrs = [];
4: for $i = 1, ..., \alpha$ do:
5: \hspace{1em} $\theta_i = \text{random}(\theta)$; $\triangleright$ Using random functions to increase the dissimilarity between chromosomes
6: \hspace{1em} $f(i) = \text{fitness}(\theta_i)$; $\triangleright$ Calculate the fitness of the individual $\theta_i$
7: \hspace{1em} if $f(i) > f_{\text{max}}$ then:
8: \hspace{2em} $f_{\text{max}} = f(i)$;
9: \hspace{2em} bestCrs = $\theta_i$;
10: end if
11: end for
12: for $i = 1, 2, ..., \beta$ do:
13: \hspace{1em} for $j = 1, 2, ..., \alpha$ do:
14: \hspace{2em} $P_j = \text{TournamentSelection}(\alpha)$; $\triangleright$ Use tournament operator to obtain parent chromosomes
15: \hspace{2em} $K_1, K_2, ..., K_\alpha = \text{Single-pointCrossover}(P_1, P_2, ..., P_\alpha)$; $\triangleright$ Generate children using the single-point crossover operator
16: \hspace{2em} $K'_j = \text{Mutation}(K_1, K_2, ..., K_\alpha)$;
17: \hspace{2em} for $j = 1, 2, ..., \alpha$ do:
18: \hspace{3em} $f(j) = \text{fitness}(K'_j)$;
19: \hspace{3em} if $f(j) > f_{\text{max}}$ then:
20: \hspace{4em} $f_{\text{max}} = f(j)$;
21: \hspace{4em} bestCrs = $K'_j$;
22: \hspace{3em} end if
23: \hspace{2em} end if
24: \hspace{1em} end for
25: \hspace{1em} end for
26: return bestCrs;

3.2 Chromosome Coding and Population Initialization

First, the individual’s chromosomes (solution) of the genetic algorithm are encoded with integers as shown in Figure 1. For the $i$th individual, its chromosomes $G^i = \{g_1, g_2, g_3, ..., g_j, ..., g_n\}$ can be represented as a vector with $n$ positions where $n$ is the number of the instances of the dataset and $g_j$ denotes the cluster number for the $j$th instance. The initialization procedure for the first generation of the parent individual is as follows: firstly, three different types of clustering algorithms K-means, BIRCH, and Spectral Clustering, are used to generate three clustering results, and then HBGF is used to fuse the three clustering results into one cluster result. At this point, the fused clustering results can only be the chromosome of one indi-
individual. There are three common ways to ensure diversity among the initialized individuals: 1) data perturbation, such as autonomous sampling of bagging; 2) attribute perturbation of the dataset, such as attribute sampling of the random subspace; 3) algorithm parameter perturbation, such as setting different hyperparameters of the base learner. After trying the above three methods in our experiments, we found that neither of them could obtain good results. Therefore, we use K-means, BIRCH, and Spectral Clustering to do the clustering and combine their clustering results by HBGF. After obtaining the first solution $\theta$, we generate more solutions by making some moderate changes to $\theta$: 30% of the chromosomes (instances) randomly chosen and changed clustering groups. The process is shown in Figure 2.

![Figure 1: Results of clustering with five groups and n chromosomes (instances).](image1)

![Figure 2: 30% of the instances are randomly chosen and changed clustering groups.](image2)

### 3.3 Objective Function

The choice of the objective function will affect the speed of convergence of the genetic algorithm and the ability to find an optimal solution. In ECA-CE, we use supervised classifiers to get the fitness of an individual and Figure 3 shows how ECA-CE employs multiple supervised classifiers to get the individual’s fitness. Instead of calculating the similarity between features as in traditional clustering algorithms, the objective function is treated as a supervised learning task. The objective function pseudo-code is shown in Algorithm 3. The calculation of the individual’s fitness of ECA-CE is based on a single objective, which makes it easier to calculate the individual’s fitness and speeds up the convergence of the algorithm.

When using supervised classifiers to calculate the fitness of an individual, the bias of a single classifier can be reduced by having multiple supervised classifiers involved in the calculation at the same time. In this paper, we choose three supervised classifiers.

#### Algorithm 3: Objective function: fitness.

**Require:** dataset $D$, individual chromosomes $\theta$

1. $X_{train}, Y_{train}, Y_{test} = train\_test\_split(D, \theta)$;
2. $res = 0$;
3. for $i = 1, \ldots, c$ do:
   - $model^i = C^i.train(X_{train}, Y_{train})$;
   - $Y_{pred} = model^i.predit(X_{test})$;
   - $res = res + f1\_score(Y_{test}, Y_{pred})$;
4. end for
5. return $res/c$;

A certain number of training sets are chosen to train each supervised classifier, and a test set is evaluated to calculate the F1 score of each supervised classifier on the test set and summed to find the mean value.

### 3.4 Genetic Algorithm Operators

a) Tournament selection: In each round of the tournament operator (Blickle and Thiele, 1995), two or more individuals are randomly selected from the population, and the individual with the greatest fitness becomes the parent.

b) Single-point crossover: In the single-point crossover operator (Deb et al., 1995), a position on the parent chromosome is chosen at random. This position becomes the crossover point. As shown in Figure 4, at the crossover point of the parents’ chromosomes, the left and right chromosomes are swapped with each other to obtain two offsprings, each carrying the genetic information of the parent’s generation.

c) Mutation operator: The mutation operation is generally applied to an offspring produced using selection and crossover operators. Because the mutation operator (Das et al., 2009) generally compromises the genetic information of the offspring, mutations usually occur with very low probability. As shown in Figure 5, using the swap mutation operator, two chromosomes are randomly selected and their values are swapped.

### 4 EXPERIMENTAL SETUP

In this section, the traditional algorithmic hyperparameters and datasets required for the ECA-CE algorithm are described in detail. All experiments were done on a PC with an Intel Core i7 2.9GHz, 8-core processor, 16GB, and the python programming language.
4.1 Datasets and Evaluation Metrics

For this experiment, we selected 20 datasets from the UCI (Asuncion and Newman, 2007) and Fränti and Sieranoja’s Clustering Benchmark repository (Fränti and Sieranoja, 2018),(Gionis et al., 2007),(Chang and Yeung, 2008),(Veenman et al., 2002),(Jain and Law, 2005). Table I shows the information of them.

All the datasets used are labelled, which are suitable for classification tasks. In this study, we use them to test clustering algorithms. Therefore, we set the number of clusters as the number of classes in each dataset, then compare the similarity between a cluster and its corresponding class. Adjusted RAND Index(Steinley, 2004) is used for the comparison. The larger the ARI value is, the better clustering result we obtain.

Table 1: Information of the datasets used in experiment.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>N. Features</th>
<th>N. Classes</th>
<th>N. Instances</th>
</tr>
</thead>
<tbody>
<tr>
<td>aggregation</td>
<td>2</td>
<td>4</td>
<td>788</td>
</tr>
<tr>
<td>jain</td>
<td>2</td>
<td>2</td>
<td>373</td>
</tr>
<tr>
<td>pathbased</td>
<td>2</td>
<td>3</td>
<td>300</td>
</tr>
<tr>
<td>r15</td>
<td>2</td>
<td>15</td>
<td>600</td>
</tr>
<tr>
<td>spiral</td>
<td>2</td>
<td>3</td>
<td>312</td>
</tr>
<tr>
<td>breast-cancer-wisconsin</td>
<td>30</td>
<td>2</td>
<td>569</td>
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<tr>
<td>breast-tissue</td>
<td>9</td>
<td>6</td>
<td>106</td>
</tr>
<tr>
<td>dermatology</td>
<td>34</td>
<td>6</td>
<td>366</td>
</tr>
<tr>
<td>ecoli</td>
<td>7</td>
<td>8</td>
<td>336</td>
</tr>
<tr>
<td>forest</td>
<td>27</td>
<td>4</td>
<td>523</td>
</tr>
<tr>
<td>glass</td>
<td>9</td>
<td>6</td>
<td>214</td>
</tr>
<tr>
<td>iris</td>
<td>4</td>
<td>3</td>
<td>150</td>
</tr>
<tr>
<td>leaf</td>
<td>14</td>
<td>36</td>
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<td>liver</td>
<td>5</td>
<td>16</td>
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<td>parkinsons</td>
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<td>195</td>
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<td>seeds</td>
<td>7</td>
<td>3</td>
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</tr>
<tr>
<td>segment</td>
<td>19</td>
<td>7</td>
<td>210</td>
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<td>transfusion</td>
<td>4</td>
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<td>wine</td>
<td>13</td>
<td>3</td>
<td>178</td>
</tr>
<tr>
<td>zoo</td>
<td>16</td>
<td>7</td>
<td>101</td>
</tr>
</tbody>
</table>

4.2 Hyperparameter Setting for all Participating Algorithms

Apart from the proposed method, five other algorithms F1-ECAC, ECAC, K-means, BIRCH, and Spectral Clustering are also tested. Among them, F1-ECAC and ECAC are recently proposed evolutionary clustering algorithms and the rest are traditional clustering algorithms. Each algorithm run 10 times and its average performance is calculated for better reliable results. For each dataset, the number of clusters
is set to be the number of categories.

a) Three evolutionary clustering methods ECA-CE, F1-ECAC, and ECAC:

Both ECA-CE and F1-ECAC generate 200 individuals per iteration for 200 iterations, while ECAC generates 20 individuals per iteration for a total of 2000 iterations. When initializing the ECA-CE population, the hyperparameters of the traditional clustering algorithm are as follows:

K-means: the number of times the algorithm is run for different values of the initialization of the center points is 10.

BIRCH: the threshold for merging a new sub-cluster with the closest sub-cluster is 0.5.

Spectral Clustering: Constructing similarity matrices using RBF kernels.

For all three methods, the probability of crossover is set to 95%, and the probability of chromosome mutation is set to 5%. They use the same method for evaluation.

b) Other traditional clustering algorithms

For K-means, the maximum number of iterations is set to 300. The algorithm is run 10 times and the average performance of them is calculated. For BIRCH, the threshold for merging new sub-clusters with the closest sub-cluster is set to 0.5 and the number of CF sub-clusters per node is set to 55. For Spectral Clustering, similarity matrices are constructed by RBF kernels.

4.3 Experimental Results

In this section we present the experimental results of the experiment. Firstly, we look at the convergence effect of the objective function of ECA-CE using two different population initialization methods i.e., the clustering ensemble algorithm initialized population and the randomly initialized population. ECA-CE searches the individual with the largest fitness in each generation as the best clustering result in that generation. As shown in Figure 6, when using the aggregation dataset, the figure shows the maximum fitness of the clustering ensemble initialization population and random initialization population algorithms in each iteration. When the initial population algorithm of the clustering ensemble has about 100 iterations, the fitness converges to 0.93. The evolutionary clustering algorithm using a random method to initialize the population shows no sign of convergence in 200 iterations, and the maximum fitness is 0.66 in the 200th iteration. HBGF and Randomization: We compared the quality of clustering results using different population initialization methods. As shown in Table 2, HBGF initialization outperforms randomization initialization on 16 out of 20 datasets in terms of ARI, which demonstrates the effectiveness of using a clustering ensemble for population initialization.

Secondly, we compare the difference in clustering quality between ECA-CE and two other evolutionary clustering algorithms, F1-ECAC, and ECAC. The difference in clustering quality between ECA-CE and other traditional clustering algorithms is also presented in Table 3. As shown in Table 3, ECA-CE is doing better than ECAC on 18/20 datasets. Especially on three datasets zoo, R15, and aggregation, the ARI
Table 4: ARI for ECA-CE and baseline algorithms.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>ECA-CE</th>
<th>F1-ECAC</th>
<th>ECAC</th>
<th>K-means</th>
<th>BIRCH</th>
<th>SC</th>
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</tbody>
</table>

In Table 4, the ARI of ECA-CE, ECAC, F1-ECAC, and the three traditional clustering algorithms are shown for all tested datasets. ECA-CE is the best in 10 of the 20 datasets, which is followed by K-means with 4, BIRCH with 3, F1-ECAC with 2, and ECAC with only 1.

4.4 Analysis of the Results

In Friedman’s test (Chatfield and Mander, 2009), the lower the average ranking of the algorithm, the better its average performance. The average ranking of ECA-CE (2.2) over the 20 datasets is shown in Table 4 to be lower than the other five algorithms, thus demonstrating the superiority of ECA-CE over the other algorithms on the experimental datasets.

5 SUMMARY

In this paper, we have proposed an evolutionary clustering algorithm ECA-CE that uses a clustering ensemble to initialize population and employs multiple supervised classifiers to evaluate the fitness scores of individuals. To ensure the diversity of the initial population, after using a clustering ensemble to obtain a solution, 30% of chromosomes. Multiple supervised classifiers are then used to jointly assess the fitness of the generated individual when designing the objective function. Furthermore, it was experimentally verified that ECA-CE outperforms the two recently proposed evolutionary clustering algorithms: F1-ECAC and ECAC in terms of ARI.

In our future work, we would investigate the use of other clustering ensemble methods to initialize the population to ensure the generation of good genetic parents. We also plan to apply ECA-CE to some more practical clustering tasks.

REFERENCES


