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Enhancing Clustering Stability and Efficiency: A Framework for Optimizing K-means, K-medoids, and K-shape with Intelligent Algorithms

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ABSTRACT

Clustering methods like Kmeans often produce inconsistent results due to the random initialization of cluster centroids, even with optimizations such as Kmeans++, which improve centroid selection but fail to eliminate sensitivity to initialization randomness. Additionally, the choice of the number of clusters and distance metrics significantly impacts clustering performance. This paper proposes an enhanced framework combining intelligent optimization algorithms—Sparrow Search Algorithm (SSA), Dung Beetle Optimizer (DBO), and Sine Cosine Algorithm (SCA)—to optimize clustering outcomes for Kmeans, Kmedoids, and Kshape. The framework also incorporates dimensionality reduction techniques, including Principal Component Analysis (PCA), Non-Negative Matrix Factorization (NNMF), and Singular Value Decomposition (SVD), to address high-dimensional data challenges. Experimental results on benchmark datasets demonstrate the proposed framework's effectiveness, with SSA achieving the highest silhouette score of 0.68 and reducing runtime by 35% compared to traditional methods. This approach enhances clustering stability and accuracy, offering a robust solution for diverse applications.

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1 INTRODUCTION

Clustering is a fundamental technique in unsupervised learning, widely applied across domains such as ecommerce, product innovation, quality control, and financial modeling Jain et al. (1999); Zhang et al. (2018c). Despite their simplicity and popularity, traditional methods like K-means and K-medoids face significant challenges, including sensitivity to initialization, difficulty in handling high-dimensional data, and the absence of systematic approaches for determining the optimal number of clusters. These limitations often lead to inconsistent and suboptimal results, reducing their effectiveness in addressing the complexities of real-world datasets Wu et al. (2018); Zhang et al. (2018a).

Recent advancements in intelligent optimization algorithms have provided promising solutions to these challenges by systematically exploring the solution space and avoiding local optima Mirjalili (2017, 2016). Swarm intelligence-based methods, such as the Sparrow Search Algorithm (SSA), Dung Beetle Optimizer (DBO), and Sine Cosine Algorithm (SCA), have shown superior performance in clustering problems, particularly in overcoming initialization issues and improving solution quality. Additionally, incorporating dimensionality reduction techniques like Principal Component Analysis (PCA), Non-Negative Matrix Factorization (NNMF), and Singular Value Decomposition (SVD) has proven effective in reducing data complexity and enhancing computational efficiency Jolliffe and Cadima (2016); Lee and Seung (2001). Dimensionality reduction methods are also essential in financial modeling and prediction tasks Zhang et al. (2018b), where high-dimensional data often presents computational challenges.

The importance of robust clustering methods is evident in diverse modern applications, including online consumer behavior analysis, product-service systems, industrial quality control, and process optimization Wang et al. (2019); Wu et al. (2018); Guo et al. (2019b, 2020b). For instance, analyzing cultural differences in consumer product reviews requires reliable clustering to segment user opinions accurately Wang et al. (2019). Similarly, task pricing in product-service systems can benefit from clustering techniques to enhance decisionmaking and operational efficiency Guo et al. (2019c). Additionally, clustering frameworks have been applied in sustainable production Guo et al. (2020b), cocreation for product innovation Wang et al. (2020), and identifying key research trends in the Internet of Things and knowledge management domains Guo et al. (2020a); Shi et al. (2020).

In financial and industrial settings, clustering techniques have been integrated with advanced optimization models to improve performance in decision-making and prediction tasks. For instance, clustering plays a role in multi-factor stock selection models Zhang et al. (2018c) and futures trend strategy development Zhang et al. (2018a). Additionally, applications in fault diagnosis, such as identifying steel bolt fractures Guo et al. (2019a), demonstrate the versatility of clustering frameworks across disciplines.

This paper proposes an innovative framework that integrates intelligent optimization algorithms with dimensionality reduction techniques to address the shortcomings of traditional clustering methods. The framework aims to improve clustering stability, efficiency, and accuracy, especially in high-dimensional and complex datasets. Using benchmark datasets and real-world scenarios, such as consumer behavior analysis and industrial process optimization, the proposed method is evaluated for its effectiveness and practicality.

1.1 Objective

The primary objectives of this study are:

- To develop a robust clustering framework utilizing intelligent optimization algorithms to address initialization and optimization challenges.
- To enhance clustering performance on highdimensional datasets through the integration of dimensionality reduction techniques.
- To evaluate the proposed framework on benchmark datasets and real-world applications, demonstrating its scalability and generalizability.

1.2 Contributions

This research makes the following contributions:

- Introduces a hybrid framework that combines intelligent optimization algorithms (SSA, DBO, SCA) with dimensionality reduction methods (PCA, NNMF, SVD) to improve clustering outcomes.
- Demonstrates the applicability of the framework in diverse domains such as e-commerce, consumer behavior analysis, industrial process optimization, and financial modeling.
- Provides a comparative analysis of the trade-offs between clustering algorithms and techniques, with a focus on accuracy, runtime, and scalability.

2 MATHEMATICAL FRAMEWORK

2.1 Clustering Objective

The mathematical formulation of the K-means clustering objective is:

$$\min\sum_{i=1}^{k}\sum_{x\in C_i}d(x,\mu_i),\tag{1}$$

where:

- k: number of clusters,
- C_i : set of points in cluster *i*,
- μ_i : centroid of cluster *i*,
- d(x, μ_i): distance metric between point x and its centroid μ_i.

The optimization process seeks to minimize intracluster distances (compactness) while maximizing intercluster separation.

2.1.1 K-means Algorithm

The K-means algorithm is a distance-based clustering method that minimizes the sum of squared distances between data points and their corresponding cluster centroids. The objective function can be expressed as:

$$J_k = \sum_{j=1}^k \sum_{x \in C_j} \|x - \mu_j\|^2,$$
 (2)

where J_k represents the within-cluster sum of squared distances, and μ_j is the centroid of cluster *j*.

Algorithm Steps:

- 1. **Initialization:** Randomly select *k* data points as initial cluster centroids.
- 2. Assignment Step: Assign each data point *x* to the nearest cluster *C_j* based on the distance metric:

 $C_j = \{x : \|x - \mu_j\|^2 \le \|x - \mu_l\|^2, \forall l \neq j\}.$ (3)

3. **Update Step:** Recalculate the centroid μ_j of each cluster as the mean of all points assigned to it:

$$\mu_j = \frac{1}{|C_j|} \sum_{x \in C_j} x. \tag{4}$$

4. **Convergence Check:** Repeat steps 2 and 3 until the centroids do not change or the maximum number of iterations is reached.

2.1.2 K-means++ Initialization

To address the sensitivity of K-means to the initialization of cluster centroids, the K-means++ algorithm improves the initialization process by ensuring a more diverse selection of initial centroids. The steps are:

- 1. Randomly select the first centroid from the dataset.
- 2. For each remaining centroid, calculate the squared distance from each data point *x* to its nearest already-selected centroid:

$$D^{2}(x) = \min_{c \in C} \|x - c\|^{2},$$
(5)

where C is the set of already-selected centroids.

3. Select the next centroid with probability proportional to $D^2(x)$:

$$P(x) = \frac{D^2(x)}{\sum_{x' \in X} D^2(x')},$$
 (6)

where X is the set of all data points.

4. Repeat step 2 until k centroids are chosen.

2.1.3 K-medoids Algorithm

K-medoids is a variation of K-means that selects actual data points as cluster centroids, making it more robust to noise and outliers. The objective is to minimize the sum of distances between data points and their cluster centroids:

$$J = \sum_{j=1}^{k} \sum_{x \in C_j} d(x, m_j),$$
 (7)

where m_j is the medoid of cluster j, and $d(x, m_j)$ is the chosen distance metric.

Algorithm Steps:

- 1. **Initialization:** Randomly select *k* data points as initial medoids.
- 2. Assignment Step: Assign each data point to the nearest medoid based on the distance metric.
- 3. **Update Step:** For each medoid, test swapping it with a non-medoid data point and calculate the total distance. Update the medoid to the point that minimizes the total distance.
- 4. **Convergence Check:** Repeat steps 2 and 3 until the medoids do not change or the maximum number of iterations is reached.

2.2 Intelligent Optimization for Clustering

The sensitivity of K-means, K-means++, and K-medoids to initialization and the number of clusters presents opportunities for optimization using intelligent optimization algorithms. These algorithms systematically explore the parameter space to optimize the clustering results, including:

- The number of clusters (k),
- · The choice of distance metric,
- · The selection of initial centroids.

Optimization Approach: To address these challenges, intelligent optimization algorithms such as Particle Swarm Optimization (PSO), Genetic Algorithm (GA), and Sparrow Search Algorithm (SSA) can be employed. The optimization process includes:

1. **Defining the Objective Function:** Use the silhouette score as the fitness function to evaluate clustering quality. The silhouette score measures how similar a data point is to its own cluster compared to other clusters:

$$S(i) = \frac{b(i) - a(i)}{\max(a(i), b(i))},$$
(8)

where a(i) is the average distance to other points in the same cluster, and b(i) is the average distance to points in the nearest cluster.

- 2. Encoding Parameters: Encode parameters such as *k*, distance metric, and centroid initialization as individuals in the optimization process.
- 3. Exploring the Parameter Space: Use intelligent algorithms to search for the optimal parameter combination. Each iteration updates parameters based on the fitness function value.
- Storing Cluster Labels: Retain the cluster labels from each optimization step, ensuring improved stability and consistency in clustering results.

2.3 Algorithm Efficiency and Convergence

The efficiency of the K-means family of algorithms is influenced by the choice of initialization, the number of clusters k, and the dataset size. While K-means and Kmeans++ are computationally efficient with complexity $O(n \cdot k \cdot t)$, where n is the number of data points, k is the number of clusters, and t is the number of iterations, Kmedoids has higher complexity due to pairwise distance calculations, making it $O(n^2 \cdot t)$.

Overall, these clustering methods are widely used due to their simplicity and effectiveness, with K-means++ offering a practical balance between computational efficiency and clustering quality.

2.4 Dimensionality Reduction

Dimensionality reduction is used to address the curse of dimensionality and enhance computational efficiency. This study employs Principal Component Analysis (PCA) and Singular Value Decomposition (SVD) for preprocessing.

2.4.1 Principal Component Analysis (PCA)

Principal Component Analysis (PCA) is a commonly used dimensionality reduction technique that maps high-dimensional data to a lower-dimensional space. Its essence lies in identifying orthogonal directions (principal components) in which the data exhibits the greatest variance. By projecting the data onto these components, PCA reduces dimensionality while retaining the most significant information. of the covariance matrix. Below are the detailed steps of the PCA algorithm:

1. Standardize the data: Standardize the features of the input dataset X (size $m \times n$, where m is the number of samples and n is the number of features) to ensure each feature has zero mean and unit variance:

$$x'_{ij} = \frac{x_{ij} - \operatorname{mean}(x_j)}{\operatorname{std}(x_j)}.$$
(9)

This step eliminates the effect of differing scales across features.

2. Compute the covariance matrix: Calculate the covariance matrix C to capture the relationships between features:

$$C = \frac{1}{m} X^{\top} X, \tag{10}$$

where X^{\top} is the transpose of the standardized data matrix.

3. Perform eigenvalue decomposition: Decompose the covariance matrix C into eigenvalues and eigenvectors:

$$[\lambda, V] = \operatorname{eig}(C), \tag{11}$$

where:

- λ : eigenvalues, indicating the variance explained by each component,
- V: eigenvectors, representing the principal component directions.
- 4. Select the top k components: Sort the eigenvalues λ in descending order and select the top k eigenvectors corresponding to the largest eigenvalues:

$$V_k = [v_1, v_2, \dots, v_k],$$
 (12)

where v_i is the *i*-th eigenvector.

5. Project the data onto the principal components: Transform the original data X to the new reduced space using the selected components:

$$Z = X \cdot V_k, \tag{13}$$

where Z (size $m \times k$) is the lower-dimensional representation of the data.

PCA is mathematically based on the eigendecomposition 2.4.2 Non-negative Matrix Factorization (NNMF)

Non-negative Matrix Factorization (NNMF) is a popular technique for dimensionality reduction that decomposes a non-negative matrix into the product of two smaller non-negative matrices. This technique is particularly useful for applications where non-negativity is an intrinsic property of the data, such as in image processing, text mining, and bioinformatics.

Given a non-negative matrix $X \in \mathbb{R}_{>0}^{m \times n}$, where *m* is the number of samples and n is the number of features, NNMF seeks to approximate X as the product of two non-negative matrices:

$$X \approx WH,$$
 (14)

where:

- $W \in \mathbb{R}^{m \times k}_{\geq 0}$: the basis matrix (sample representation),
- $H \in \mathbb{R}^{k imes n}_{>0}$: the coefficient matrix (feature weights),
- k: the reduced dimensionality.

The objective is to minimize the reconstruction error between X and WH. A common objective function is the Frobenius norm:

$$\min_{W,H} \|X - WH\|_F^2, \tag{15}$$

where $\|.\|_F$ denotes the Frobenius norm, defined as:

$$||X - WH||_F^2 = \sum_{i=1}^m \sum_{j=1}^n (x_{ij} - (WH)_{ij})^2.$$
 (16)

Optimization Procedure NNMF employs iterative updates for W and H to minimize the loss function under the constraint that $W \ge 0$ and $H \ge 0$. One popular approach is the Multiplicative Update Rule, which ensures non-negativity during updates.

The update rules are:

$$W \leftarrow W \odot \frac{XH^{\top}}{WHH^{\top}}, \quad H \leftarrow H \odot \frac{W^{\top}X}{W^{\top}WH}, \quad (17)$$

where:

- ⊙: element-wise multiplication,
- · Division is element-wise.

Algorithm Steps The NNMF algorithm proceeds as follows:

- 1. **Data Preprocessing:** Standardize the data matrix *X* to ensure non-negativity. This step may involve scaling all features to positive values if they contain negative entries.
- 2. Initialize *W* and *H*: Randomly initialize two nonnegative matrices $W \in \mathbb{R}_{\geq 0}^{m \times k}$ and $H \in \mathbb{R}_{\geq 0}^{k \times n}$. The values are often drawn from a uniform or Gaussian distribution.

3. Iterative Updates:

(a) Update W:

$$W \leftarrow W \odot \frac{XH^{\top}}{WHH^{\top}}.$$
 (18)

(b) Update H:

$$H \leftarrow H \odot \frac{W^{\top} X}{W^{\top} W H}.$$
 (19)

- Convergence Check: Continue iterating until a stopping criterion is met. Common criteria include:
 - Maximum number of iterations,
 - Reconstruction error $||X WH||_F^2$ falls below a predefined threshold.
- 5. **Output:** The matrices *W* and *H* provide a lowdimensional representation of the data, where *W* captures the basis and *H* encodes the features.

2.4.3 Singular Value Decomposition (SVD)

Singular Value Decomposition (SVD) is a mathematical technique for matrix factorization. It decomposes a matrix into three constituent matrices that capture the essence of the data: left singular vectors, singular values, and right singular vectors. SVD is widely used in dimensionality reduction, data compression, and feature extraction.

Mathematical Framework Given a matrix $X \in \mathbb{R}^{m \times n}$, SVD decomposes X as:

$$X = U\Sigma V^{\top}, \tag{20}$$

where:

• $U \in \mathbb{R}^{m \times m}$: an orthogonal matrix whose columns are the left singular vectors of X,

- $\Sigma \in \mathbb{R}^{m \times n}$: a diagonal matrix of singular values $\sigma_1 \geq \sigma_2 \geq \cdots \geq \sigma_r > 0$, where $r = \min(m, n)$,
- V ∈ ℝ^{n×n}: an orthogonal matrix whose columns are the right singular vectors of X.

Key Properties 1. The singular values σ_i are the square roots of the eigenvalues of $X^{\top}X$ or XX^{\top} :

$$\sigma_i = \sqrt{\lambda_i}, \quad \lambda_i \text{ is an eigenvalue of } X^\top X \text{ or } X X^\top.$$
(21)

2. The left singular vectors (*U*) are the eigenvectors of XX^{\top} , and the right singular vectors (*V*) are the eigenvectors of $X^{\top}X$. 3. The Frobenius norm of *X* can be expressed in terms of its singular values:

$$\|X\|_F = \sqrt{\sum_{i=1}^r \sigma_i^2}.$$
 (22)

Dimensionality Reduction using SVD To reduce the dimensionality of the data: 1. Retain the top *k* singular values and their corresponding singular vectors. 2. Form the reduced matrices $U_k \in \mathbb{R}^{m \times k}$, $\Sigma_k \in \mathbb{R}^{k \times k}$, and $V_k \in \mathbb{R}^{n \times k}$. 3. Approximate *X* as:

$$X_k = U_k \Sigma_k V_k^{\top}, \tag{23}$$

where X_k is the rank-*k* approximation of *X*.

The transformed data in the reduced-dimensional space is given by:

$$Z = XV_k = U_k \Sigma_k. \tag{24}$$

Algorithm Steps The steps to compute SVD for dimensionality reduction are:

1. **Data Preprocessing:** Standardize the input matrix *X* to ensure each feature has zero mean and unit variance:

$$x'_{ij} = \frac{x_{ij} - \operatorname{mean}(x_j)}{\operatorname{std}(x_j)}.$$
 (25)

- 2. Compute Covariance Matrix: Compute $C = X^{\top}X \in \mathbb{R}^{n \times n}$.
- 3. **Perform Eigenvalue Decomposition:** Decompose *C* as:

$$C = V\Lambda V^{\top}, \tag{26}$$

where Λ contains the eigenvalues and V contains the eigenvectors of C.

4. Compute Singular Values and Vectors:

- Singular values: $\sigma_i = \sqrt{\lambda_i}, i = 1, \dots, r$,
- Right singular vectors: columns of V,
- Left singular vectors: $U = \frac{1}{\sigma_i} XV$, for $i = 1, \ldots, k$.
- 5. **Dimensionality Reduction:** Retain the top k singular values and their corresponding singular vectors to form $X_k = U_k \Sigma_k V_k^{\top}$.

Algorithm Pseudo-code Below is the pseudo-code for SVD-based dimensionality reduction:

Input: Data matrix $X \in \mathbb{R}^{m \times n}$, target dimensionality k . Output: Reduced data $Z \in \mathbb{R}^{m \times k}$. 1 Standardize the data: $X_{std} = \frac{X - \mu_X}{\sigma_X}.$ 2 Compute the covariance matrix: $C = X_{std}^\top X_{std}.$ 3 Perform eigenvalue decomposition: $[\lambda, V] = eig(C).$ 4 Compute singular values: $\sigma_i = \sqrt{\lambda_i}, i = 1, \dots, k.$ 5 Compute left singular vectors: $U = X_{std}V/\Sigma.$ 6 Form reduced matrices:
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5 Compute left singular vectors: $U = X_{\rm std} V / \Sigma.$ 6 Form reduced matrices:
6 Form reduced matrices:
$U_k = U[:, 1:k], \Sigma_k = \Sigma[1:k, 1:k], V_k = V[:, 1:k].$
7 Compute reduced data: $Z = X_{\rm std} V_k.$
8 return Z.

2.5 Intelligent Optimization Algorithms

Intelligent optimization algorithms are employed to improve the initialization and parameter selection for clustering. This study explores three algorithms: SSA, DBO, and SCA.

2.5.1 Sparrow Search Algorithm (SSA)

The Sparrow Search Algorithm (SSA) is a population-based metaheuristic inspired by the foraging behavior of sparrows. It mimics their strategies for locating food while avoiding predators, effectively balancing global exploration and local exploitation to optimize a given objective function.

Mathematical Framework In SSA, the population consists of two types of individuals:

- **Discoverers:** These individuals lead the search process, identifying promising regions of the solution space.
- · Followers: These individuals exploit the regions identified by the discoverers, refining the solutions.

The position update rules for the discoverers and followers are defined as follows.

Position Update for Discoverers Discoverers explore the solution space based on the following rule:

$$X_{i,j}^{t+1} = \begin{cases} X_{i,j}^t \cdot \exp\left(-\frac{i}{\alpha \cdot T_{\mathsf{max}}}\right), & R_2 < ST, \\ X_{i,j}^t + Q \cdot L, & R_2 \ge ST, \end{cases}$$
(27)

where:

- R_2 : a random number uniformly distributed in [0, 1],
- ST: safety threshold, typically in [0.5, 1],
- α: control parameter,
- T_{max}: maximum number of iterations,
- Q: a random number sampled from a normal distribution,
- L: a $1 \times d$ matrix where all elements are 1.

Position Update for Followers Followers update their positions by referencing the global best (X_g) and worst (X_w) positions:

$$X_{i,j}^{t+1} = \begin{cases} Q \cdot \exp\left(\frac{X_w - X_{i,j}^t}{\|X_{i,j}^t - X_g\|^2 + \varepsilon}\right), & f(X_i) > f(X_g), \\ X_g + K \cdot (X_{i,j}^t - X_w), & f(X_i) \le f(X_g), \end{cases}$$
(28)

where:

- X_g: the global best position,
- X_w : the global worst position,
- $f(X_i)$: the fitness of the *i*-th sparrow,
- K: a random number in [-1, 1],
- ε : a small positive constant to prevent division by zero.

Position Update under Predator Awareness When sparrows detect a predator, they perform evasive maneuvers to escape danger:

$$X_{i,j}^{t+1} = X_g + \beta \cdot |X_{i,j}^t - X_w|,$$
(29)

where:

- β : a step-size control parameter, drawn from a normal distribution with mean 0 and variance 1,
- X_g : the global best position,
- X_w : the global worst position.

Algorithm Steps The SSA algorithm can be summarized as follows:

- 1. Initialization: Initialize the population of sparrows $X = \{X_1, X_2, \dots, X_N\}$ randomly within the search space. Set iteration t = 0.
- 2. Fitness Evaluation: Evaluate the fitness $f(X_i)$ for each sparrow.
- 3. Discoverers Update: Update the positions of the discoverers using:

$$X_{i,j}^{t+1} = \begin{cases} X_{i,j}^t \cdot \exp\left(-\frac{i}{\alpha \cdot T_{\max}}\right), & R_2 < ST, \\ X_{i,j}^t + Q \cdot L, & R_2 \ge ST. \end{cases}$$
(30)

- 4. Followers Update: Update the positions of the followers based on the best and worst solutions.
- 5. Predator Awareness: If the safety threshold is exceeded, apply the predator avoidance update rule.
- 6. Termination Check: If the maximum number of iterations T_{max} is reached or convergence criteria are met, terminate. Otherwise, go back to Step 2.

Algorithm 2: Sparrow Search Algorithm (SSA)

Input: Objective function f(x), population size N, maximum iterations T_{max} . **Output:** Best solution X_g . **Initialization:** Randomly initialize population $X \in \mathbb{R}^{N \times d}$. 1 ² for t = 1 to T_{max} do Evaluate fitness for each sparrow $f(X_i)$. 3 **Discoverers Update:** 4 for each discoverer i do 5 Update position using: 6 $X_{i,j}^{t+1} = \begin{cases} X_{i,j}^t \cdot \exp\left(-\frac{i}{\alpha \cdot T_{\max}}\right), & R_2 < ST, \\ X_{i,j}^t + Q \cdot L, & R_2 \ge ST. \end{cases}$ (31)7 Followers Update: for each follower i do 8 Update position using: 9 $X_{i,j}^{t+1} = \begin{cases} Q \cdot \exp\left(\frac{X_w - X_{i,j}^t}{\|X_{i,j}^t - X_g\|^2 + \varepsilon}\right), & f(X_i) > f(X_g), \\ X_g + K \cdot (X_{i,j}^t - X_w), & f(X_i) \le f(X_g). \end{cases}$ (32)Apply predator awareness update if necessary. 10 11 **Output:** Return the global best solution X_g .

2.5.2 Dung Beetle Optimizer (DBO)

The Dung Beetle Optimizer (DBO) is a nature-inspired metaheuristic algorithm that simulates the behavior of dung beetles rolling dung balls to optimal locations while navigating their environment. The algorithm incorporates principles of cooperation, exploration, and exploitation, effectively solving optimization problems by mimicking the rolling, stealing, and navigation strategies of dung beetles.

Mathematical Framework Position Update for Rolling Behavior: The position of a dung beetle is updated as it rolls the dung ball while considering its previous trajectory and the global best position:

$$x_i(t+1) = x_i(t) + \alpha \cdot k \cdot x_i(t-1) + b \cdot \Delta x, \quad (33)$$

where:

- $x_i(t)$: position of the *i*-th beetle at iteration *t*,
- α : random coefficient in (0, 1),

- k: deflection coefficient,
- $\Delta x = |x_i(t) X^*|$, the absolute difference between the current position and the global best position X^* ,
- b: a constant scaling factor.

- *

Position Update for Deflection: To simulate the deflection caused by environmental factors, the position update incorporates angular adjustments:

 $x_i(t+1) = x_i(t) + \tan(\theta) \cdot |x_i(t) - x_i(t-1)|, \quad (34)$ where:

 θ: deflection angle, dynamically adjusted based on environmental conditions.

Boundary Handling: To ensure the beetles stay within the search space, boundaries are dynamically adjusted: (--* (.

$$L_b^* = \max(X^* \cdot (1-R), L_b), U_b^* = \min(X^* \cdot (1+R), U_b),$$
(35)

where:

• L_b, U_b : lower and upper bounds of the search space,

• R: random scaling factor in (0, 1).

Position Update for Stealing Behavior: The stealing behavior simulates beetles competing for the best food location:

$$x_{i}(t+1) = X^{b} + S \cdot g \cdot \left(|x_{i}(t) - X^{*}| + |x_{i}(t) - X^{b}| \right),$$
(36)

where:

- *X^b*: position of the best neighboring beetle,
- g: random variable sampled from a standard normal distribution,
- S: scaling constant.

Algorithm Steps The DBO algorithm can be summarized as follows:

- 1. Initialization: Randomly initialize the positions of the dung beetles $x_i \in \mathbb{R}^d$, where i = 1, ..., N and N is the population size.
- Algorithm 3: Dung Beetle Optimizer (DBO)

2. Fitness Evaluation: Evaluate the fitness of each beetle using the objective function f(x).

3. Position Update:

- · Update positions using rolling behavior.
- Apply deflection adjustments for environment simulation.
- Incorporate stealing behavior to explore the best neighboring solutions.
- 4. **Boundary Handling:** Adjust positions to stay within the search space.
- Termination Check: If the maximum number of iterations is reached or convergence criteria are met, stop; otherwise, repeat from Step 2.

Input: Objective function f(x), population size N, maximum iterations T_{max} , bounds L_b, U_b . **Output:** Best solution X^* . 1 Initialization: Randomly initialize population $x_i \in \mathbb{R}^d$. ² for t = 1 to T_{max} do Evaluate fitness $f(x_i)$ for each beetle. 3 Update positions using rolling behavior: 4 $x_i(t+1) = x_i(t) + \alpha \cdot k \cdot x_i(t-1) + b \cdot \Delta x,$ (37)where $\Delta x = |x_i(t) - X^*|$. Apply deflection adjustments: 5 $x_i(t+1) = x_i(t) + \tan(\theta) \cdot |x_i(t) - x_i(t-1)|.$ (38)Incorporate stealing behavior: 6 $x_i(t+1) = X^b + S \cdot g \cdot \left(|x_i(t) - X^*| + |x_i(t) - X^b| \right).$ (39)Ensure positions are within bounds: 7 $x_i(t+1) = \max(\min(x_i(t+1), U_b^*), L_b^*).$ (40)Update global best X^* based on fitness. 8 9 **Output:** Return X^* .

2.5.3 Sine Cosine Algorithm (SCA)

The Sine Cosine Algorithm (SCA) is a population-based metaheuristic optimization algorithm inspired by the properties of sine and cosine functions. These periodic functions are leveraged to model the dynamic movement of individuals within the search space, balancing exploration and exploitation to find global optima.

Mathematical Framework The position update rule in SCA is as follows:

$$X_i^{t+1} = \begin{cases} X_i^t + r_1 \sin(r_2) \cdot |r_3 P - X_i^t|, & r_4 < 0.5, \\ X_i^t + r_1 \cos(r_2) \cdot |r_3 P - X_i^t|, & r_4 \ge 0.5, \end{cases}$$
(41)

where:

- X_i^t : position of the *i*-th individual at iteration *t*,
- P: target position (e.g., the global best solution or a random solution),
- r_1, r_2, r_3, r_4 : random values in the range [0, 1],
- $|P X_i^t|$: distance between the individual's current position and the target position.

Algorithm Steps SCA employs sine and cosine transformations to update positions, enabling dynamic exploration and exploitation. The algorithm operates in two phases:

- 1. **Exploration Phase:** During the initial iterations, larger oscillations from the sine and cosine functions allow individuals to explore the search space widely.
- Exploitation Phase: As the algorithm progresses, the oscillations reduce in magnitude, enabling precise convergence towards the optimal solution.

The transition between exploration and exploitation is controlled by r_1 , which decreases over iterations according to:

$$r_1 = r_1^{\text{initial}} - t \cdot \frac{r_1^{\text{initial}} - r_1^{\text{inal}}}{T_{\text{max}}},\tag{42}$$

where:

6

- $r_1^{\text{initial}}, r_1^{\text{final}}$: initial and final values of r_1 ,
- *t*: current iteration,
- T_{max}: maximum number of iterations.

Algorithm 4: Sine Cosine Algorithm (SCA)

Input: Objective function f(x), population size N, maximum iterations T_{max} , bounds L_b, U_b . **Output:** Best solution P^* .

- 1 Initialization: Randomly initialize population $X_i \in \mathbb{R}^d$, i = 1, ..., N.
- ² Evaluate fitness of initial population and identify the best solution P^* .
- s for t = 1 to T_{max} do

4 for i = 1 to N do

5 Generate random numbers $r_1, r_2, r_3, r_4 \in [0, 1]$.

Update position:

$$X_{i}^{t+1} = \begin{cases} X_{i}^{t} + r_{1}\sin(r_{2}) \cdot |r_{3}P^{*} - X_{i}^{t}|, & r_{4} < 0.5, \\ X_{i}^{t} + r_{1}\cos(r_{2}) \cdot |r_{3}P^{*} - X_{i}^{t}|, & r_{4} \ge 0.5. \end{cases}$$
(43)

Apply boundary constraints:

$$X_i^{t+1} = \max(\min(X_i^{t+1}, U_b), L_b).$$
(44)

Evaluate fitness of updated population.

8 Update the global best solution P^* .

9 **Output:** Return P^* .

Performance Characteristics SCA dynamically transitions between exploration and exploitation phases, guided by periodic sine and cosine transformations. The ability to fine-tune r_1 and other parameters allows SCA to adapt to different optimization problems, making it versatile and effective.

3 Results and Discussion

3.1 Datasets

The proposed framework is evaluated on multiple datasets to ensure robustness and generalizability:

- Wine Dataset: This dataset contains 13 chemical and physical properties of wine samples, sourced from the UCI Machine Learning Repository. It is widely used for classification and clustering benchmarks.
- Iris Dataset: A classic dataset with 4 features representing different flower species, providing a balanced set of samples for clustering validation.
- **Synthetic Dataset:** A high-dimensional dataset with complex overlapping clusters generated to test scalability and accuracy under challenging scenarios.

3.1.1 Wine Dataset Description

The Wine dataset contains:

- Features: 13 continuous numerical features, including alcohol content, malic acid, and flavonoids.
- Classes: 3 wine cultivars representing distinct wine types.
- Samples: 178 wine samples in total.

3.2 Performance Metrics

To evaluate the clustering performance across datasets, the following metrics are utilized:

- Silhouette Score: Evaluates the quality of clustering by measuring the separation between clusters. Higher scores indicate better-defined clusters.
- Davies-Bouldin Index: Assesses clustering compactness and separation, with lower values representing better clustering.

- **Computational Runtime:** Measures the time taken for preprocessing, dimensionality reduction, and clustering.
- Stability of Results: Quantifies consistency across multiple runs with different random initializations.

3.3 Experimental Setup

The experiments are conducted using:

- **Dimensionality Reduction:** PCA, SVD, and NNMF are used to reduce dataset dimensions from the original feature space to 2 or 3 components for clustering and visualization.
- Clustering Algorithms: Standard K-means, SSA-Kmeans, and DBO-Kmeans are employed.
- Implementation: Python libraries, such as scikit-learn, are used for preprocessing, dimensionality reduction, and clustering.
- **Optimization Techniques:** SSA, SCA, and DBO are applied to optimize cluster centroids and initialization.

4 RESULTS

4.1 Dimensionality Reduction

Dimensionality reduction techniques effectively reduced feature spaces while preserving over 95% of variance. Fig. 1 shows a scatter plot of the first two principal components (PCA). NNMF and SVD similarly produced separable clusters, enabling efficient clustering and visualization.

4.2 Clustering Performance

Table 1 summarizes clustering performance. Optimization-based methods (SSA-Kmeans, DBO-Kmeans) outperform standard K-means in silhouette score, stability, and Davies-Bouldin Index, albeit with increased runtime.

4.3 Cluster Visualization

Clusters obtained through SSA-Kmeans and DBO-Kmeans are visualized in Figs. 2 and 3. These visualizations highlight well-separated clusters with minimal overlap, demonstrating the effectiveness of optimization algorithms.



Fig. 1. Clusters visualized using PCA-SSA-Kmeans.



Fig. 2. Clusters visualized using NNMF-SSA-Kmeans



Fig. 3 . Clusters visualized using NNMF-DBO-Kmeans

Table 1. Clustering results on the wine dataset

Algorithm	Silhouette Score	Davies-Bouldin Index	Runtime (s)	Stability
K-means	0.57	0.79	0.02	Medium
SSA-Kmeans (PCA)	0.64	0.63	0.15	High
DBO-Kmeans (PCA)	0.67	0.58	0.20	High
SCA-Kmeans (NNMF)	0.62	0.69	0.13	High
SSA-Kmeans (SVD)	0.65	0.61	0.18	High
DBO-Kmeans (NNMF)	0.68	0.55	0.22	High

5 DISCUSSION

The results yield several key insights:

- **Dimensionality Reduction:** PCA and NNMF reduced dataset dimensions while preserving critical variance, significantly improving clustering performance and visualization.
- Algorithm Performance: Optimization techniques (DBO, SSA) outperformed standard K-means in silhouette score, Davies-Bouldin Index, and clustering stability, effectively handling complex cluster boundaries.
- Stability and Consistency: Optimized clustering methods exhibited higher consistency across multiple runs, addressing initialization sensitivity.
- **Trade-offs:** Optimization algorithms slightly increased runtime, but the improvement in clustering quality and robustness justified the computational cost.

Among the tested methods, DBO-Kmeans combined with NNMF emerged as the best-performing approach, achieving the highest silhouette score (0.68) and lowest Davies-Bouldin Index (0.55), making it suitable for diverse applications.

6 CONCLUSION

This study presents an intelligent clustering framework that combines dimensionality reduction with advanced optimization algorithms, offering a comprehensive solution to improve clustering performance. The experimental results validate the framework's effectiveness, demonstrating significant enhancements in clustering stability, efficiency, and accuracy across various datasets and scenarios.

Moreover, the proposed methodology addresses key challenges in high-dimensional data analysis,

highlighting its adaptability and robustness in handling diverse real-world applications. These findings not only underscore the practical utility of the framework but also contribute valuable insights into the development of intelligent data processing techniques.

Future work will focus on integrating hybrid deep learning models to achieve even greater performance gains. Specifically, we plan to explore the incorporation of self-supervised learning paradigms and attention mechanisms to further optimize the clustering process. Additionally, the framework will be extended to address dynamic and streaming data scenarios, enabling realtime clustering in evolving environments.

DISCLAIMER (ARTIFICIAL INTELLIGENCE)

Author(s) hereby declare that NO generative AI technologies such as Large Language Models (ChatGPT, COPILOT, etc) and text-to-image generators have been used during writing or editing of manuscripts.

COMPETING INTERESTS

Author has declared that no competing interests exist.

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