

Dynamic Graph Neural Evolution: An Evolutionary Framework

Integrating Graph Neural Networks with Adaptive Filtering

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Abstract

This paper proposes an innovative optimization framework, Dynamic Graph Neural Evolution (DGNE), integrating Graph Neural Networks (GNNs) with Evolutionary Algorithms (EAs). Building on the foundation of Graph Neural Evolution (GNE), DGNE introduces a dynamic filtering mechanism and adaptive Gaussian sampling functions to dynamically adjust the population distribution during the optimization process, achieving a balance between global exploration and local exploitation. By emphasizing high-frequency information in the early stages to enhance population diversity and low-frequency information in the later stages to promote convergence, DGNE effectively optimizes the search process. Experiments were conducted on the CEC2017 benchmark suite across 30, 50, and 100 dimensions, comparing DGNE with advanced algorithms (LSHADE, LSHADE_cnEpSin, MadDE, SaDE, EA4eig) and classic algorithms (DE and CMA-ES). Statistical analyses using the Wilcoxon rank-sum test and Friedman mean rank test demonstrate that DGNE achieves the best average ranking in 50 and 100 dimensions and ranks third in 30 dimensions. However, it achieves the highest overall average performance ranking across all dimensions, showcasing its stability and significant advantages in different scenarios. While

its performance in low-dimensional tasks is slightly less competitive compared to high-dimensional ones, DGNE still exhibits strong competitiveness. Additionally, we explored the impact of population size and iteration counts. DGNE was evaluated across population sizes of 20, 30, 50, and 100. The results highlight DGNE's robustness, maintaining competitive rankings across all population sizes, with top rankings for smaller population sizes (20 and 30) and strong results at larger sizes (50 and 100). In terms of iteration counts (500, 1000, and 2000 iterations), DGNE exhibited stable and competitive performance, ranking 1st at 500 iterations and maintaining top-tier rankings even at extended iteration counts. These findings confirm DGNE's adaptability to varying configurations and further validate its effectiveness as a robust optimization framework. Overall, DGNE demonstrates great potential as an optimization method, offering a promising direction for further research and applications in artificial intelligence and optimization fields. Its ability to remain competitive across diverse tasks and configurations underscores its versatility and scalability.

Keywords: Machine Learning, Graph Neural Networks (GNNs), Evolutionary Algorithms (EAs), Dynamic Graph Neural Evolution (DGNE), CEC2017 Benchmark Suite,

1.Introduction

Evolutionary Computation (EC) and Machine Learning (ML) are two key branches of artificial intelligence, offering powerful tools for solving complex problems. EC, inspired by biological evolution, simulates mechanisms like selection, crossover, and mutation to optimize solution spaces. Emerging in the mid-20th century[1-3], EC's independence from mathematical properties like differentiability or convexity makes it highly robust and flexible for high-dimensional, nonlinear, and multi-objective optimization tasks. Classic EC algorithms include the Genetic Algorithm (GA)[4], which mimics natural selection, Differential Evolution (DE)[5], known for its simplicity and efficiency in continuous-space optimization, and Covariance Matrix Adaptation Evolution Strategy (CMA-ES)[6], which enhances performance in high-dimensional problems by modeling solution distributions.

In recent years, several DE improvements have further advanced its performance and adaptability[7-11]. Success-History Based Adaptive Differential Evolution(SHADE) [12]adapts control parameters based on historical successes, improving convergence and global search. SHADE with Linear Population Size Reduction(LSHADE)[13] reduces population size over time, focusing resources on fine-tuning in later stages. Extensions like Ensemble sinusoidal differential covariance matrix adaptation with Euclidean neighborhood (LSHADE_cnEpSin)[14] introduce sinusoidal-based adaptive strategies with Euclidean knowledge sharing and covariance matrix modeling, while LSHADE with semi-parameter adaptation hybrid with CMA-ES (LSHADE_SPACMA)[15] integrates CMA-ES for balancing global exploration and local exploitation, excelling in high-dimensional problems like the CEC 2017 benchmarks. Additionally, Adaptive Differential

Evolution with Optional External Archive(JADE)[16] and Self-Adaptive Differential Evolution(SaDE)[17] further enhance DE with novel mutation strategies, external archives, and self-adaptive strategy selection, boosting performance and adaptability in diverse optimization challenges.

ML is a critical branch of artificial intelligence, aiming to learn patterns and models from data to enable predictions and decision-making on unseen data. Traditional machine learning methods, such as linear regression, support vector machines, and decision trees[18-20], rely heavily on handcrafted features and statistical models. With the growth of data availability and computational power, the introduction of neural networks marked a significant breakthrough in machine learning. By mimicking the structure and information processing mechanisms of the human brain, neural networks enabled the automatic extraction of features from data, establishing a major milestone in the evolution of machine learning. In the 21st century, neural networks advanced further into deep learning, which utilizes multilayered network architectures to learn complex data representations. This advancement has led to revolutionary progress in areas such as speech recognition, image processing, and natural language processing.

The success of deep learning can be attributed to a series of classic architectures. For instance, Convolutional Neural Network (CNN)[21] have demonstrated exceptional performance in computer vision tasks, with models like Residual Network (ResNet)[22] overcoming the training challenges of deep networks. Recurrent Neural Network (RNN)[23] and their variants, such as Long Short-Term Memory(LSTM)[24] and Gated Recurrent Unit(GRU)[25], excel in handling sequential data and time-series tasks. Meanwhile, the Transformer architecture[26] has redefined natural language processing through its attention mechanism, giving rise to groundbreaking models like BERT[27] and GPT[28-30], which are widely applied across various domains. Additionally, Graph Neural Networks (GNNs)[31-34] have emerged to handle non-Euclidean data structures, such as social networks and knowledge graphs, showing immense potential in applications like recommendation systems and molecular modeling. These innovations have propelled deep learning to new heights in data analysis and artificial intelligence applications, continuously pushing the boundaries of what is achievable with machine learning.

In recent years, the intersection of EC and ML has emerged as a significant direction in artificial intelligence research. Leveraging its powerful global optimization capabilities, evolutionary algorithms have been widely applied to optimize hyperparameters and neural network structures in machine learning models[35, 36]. Conversely, machine learning has significantly advanced evolutionary computation by introducing techniques that enhance its efficiency and adaptability. Methods such as surrogate modeling[37-39]and reinforcement learning[40, 41] have been utilized to guide search processes, reduce computational costs, and dynamically adjust parameters and operators. Studies have revealed a remarkable consistency between evolutionary algorithms and diffusion models[42], large language models(LLMs)[43, 44], and GNNs[45]. Recent studies have

proposed the Graph Neural Evolution (GNE) framework, which models individuals in evolutionary algorithms as graph nodes and leverages frequency-domain filters to effectively balance global exploration and local exploitation[45-47]. This approach has demonstrated exceptional performance in solving complex optimization problems. GNE not only advances the development of evolutionary algorithms but also inspires innovations in GNNs, enhancing global information propagation and mitigating the issue of over-smoothing. As a result, GNE represents a key step forward in the integration of EC and ML, offering significant academic value and practical application potential in addressing challenging optimization problems and advancing intelligent systems. It is worth mentioning that many studies have applied GNNs to solve combinatorial optimization problems[48, 49]. However, these studies differ from GNE in that they focus on spatial-domain-based GNNs, where information is transmitted across the graph, and a loss function is defined for backpropagation to update the weights. In contrast, the GNE framework is based on frequency-domain GNNs, utilizing its mathematical framework to design the evolutionary process, without the need for a loss function or backpropagation to update the weights.

Inspired by recent research on GNE, this paper extends the filter design proposed in GNE into a dynamic filtering mechanism. Specifically, in the early stages of evolution, the filter is designed to enhance the high-frequency information of the population to improve global exploration capability. In the later stages, the filter focuses on amplifying low-frequency information, thereby strengthening exploitation ability and achieving a dynamic balance between exploration and exploitation. Additionally, a new sampling method is designed to further improve the convergence of the population. To evaluate the performance of the proposed algorithm, experiments were conducted on the CEC2017 benchmark suite for dimensions of 30, 50, and 100. The results were compared with several state-of-the-art differential evolution algorithms, including LSHADE[13], LSHADE_cnEpSin[14], which is the winner of the CEC2017 benchmark suite, Multiple Adaptation Differential Evolution (MadDE)[50], SaDE[17], and Four Evolutionary Algorithms with Eigen crossover (EA4eig)[51], which is the winner of the CEC2022 benchmark suite. The experimental results demonstrate that the proposed method shows significant advantages in terms of convergence speed, robustness, and adaptability, particularly excelling in handling complex optimization problems.

The remainder of this paper is organized as follows: In Section 2, we review the research on GNE and propose Dynamic Graph Neural Evolution (DGNE), providing its mathematical formulation. In Section 3, we analyze the performance of DGNE on the CEC2017 benchmark suite and conduct comparative experiments. In Section 4, we summarize the findings of the paper and provide future research directions.

2. Dynamic Graph Neural Evolution (DGNE)

2.1 Review of GNE

GNE is an innovative approach that combines Evolutionary Algorithms (EAs) with GNNs, representing the population through graph structures and leveraging the characteristics of graphs during the optimization process. In GNE, each individual in the population is modeled as a node in the graph, and the similarities between individuals are encoded via the adjacency matrix. This graph-based representation enables GNE to utilize spectral filtering techniques to dynamically balance exploration and exploitation during optimization, achieving robust and efficient optimization.

The first step of GNE is to represent the population as a graph. Each individual X_i in the population is treated as a node in the graph, and the similarity between any two individuals X_i and X_j is calculated based on cosine similarity. The adjacency matrix A is defined as follows:

$$A_{ij} = \frac{Z_i Z_j}{\|Z_i\| \|Z_j\|} \quad (1)$$

where $Z_i = X_i - X_0$ represents the difference vector between individual X_i and the population center X_0 , and A_{ij} represents the similarity between nodes i and j . To normalize the graph, the normalized Laplacian matrix is constructed as:

$$L_{norm} = I - D^{-\frac{1}{2}} A D^{-\frac{1}{2}} \quad (2)$$

where D is the degree matrix, A is the adjacency matrix, and I is the identity matrix. By performing spectral decomposition on the Laplacian matrix L_{norm} , its eigenvalues Λ and eigenvectors U can be obtained:

$$L_{norm} = U \Lambda U^T \quad (3)$$

These spectral components allow the definition of a filter function $g(\Lambda)$, which selectively amplifies or suppresses specific frequency components. Low-frequency components represent the global stability and shared information of the population, while high-frequency components capture local diversity and individual differences within the population. The filtered population can then be expressed as:

$$X_{new} = U g(\Lambda) U^T X \quad (4)$$

In spectral filtering, existing spectral-based graph neural networks typically apply nonlinear transformations φ, ϕ to the original feature matrix X or the filtered feature matrix. These transformations are usually functions similar to Multi-Layer Perceptrons (MLPs). During the training process of graph neural networks, the parameters of these nonlinear transformations are continuously updated using the gradient descent algorithm with a specific learning rate. In this process, the changes in φ and ϕ can be viewed as a resampling operation on the feature matrix in its respective feature space. Through this resampling, features are updated along the optimization

direction with a certain step size, gradually adjusting their distribution to better fit the requirements of specific problems. This enhances the model's representational capability and optimization performance for the target task.

Similarly, in evolutionary algorithms, the nonlinear transformations φ and ϕ can be implemented as resampling operations of the current population along the optimization direction. This process progressively adjusts the population's distribution through resampling, making it more concentrated in high-fitness regions and thereby improving optimization results. Based on this idea, GNE introduces an evolutionary algorithm combined with graph neural networks, where the overall population update process is described by the following equation:

$$X_{new} = \phi(Ug(\Lambda)U^T\varphi(X)) \quad (5)$$

In Eq. (5), φ and ϕ serve as resampling functions applied before and after the filtering operation. By adding resampling functions before and after filtering, GNE further optimizes the population distribution, achieving a more efficient optimization process. This dual transformation leverages the characteristics of graph neural networks, not only extracting global stability and local diversity from the population during spectral filtering but also dynamically adjusting the population distribution to enhance the search capability and adaptability of the evolutionary algorithm. The core idea of GNE is to integrate the spectral filtering and resampling mechanisms of graph neural networks into evolutionary algorithms. Through spectral decomposition and resampling operations, it dynamically balances exploration and exploitation within the population. This approach allows the population to fully leverage global and local information from the graph during the optimization process, achieving robust and efficient optimization.

2.2 Mathematical Model of Dynamic Graph Neural Evolution (DGNE)

DGNE builds on the foundation of GNE by introducing dynamic filtering mechanisms and adaptive Gaussian sampling functions. These mechanisms enable DGNE to dynamically adjust the population distribution during the optimization process via spectral filtering and adaptive resampling strategies, thereby balancing exploration and exploitation.

2.2.1 Construction of the Dynamic Filter

At each iteration, DGNE constructs a dynamic filter to balance the contributions of high-frequency and low-frequency components in the spectral domain. High-frequency components capture local diversity, while low-frequency components reflect global stability. The weights of the high- and low-frequency components are dynamically adjusted based on the current iteration t , with the weight definitions as follows:

$$w_h(t) = 0.8 - 0.5 \left(\frac{t}{T}\right)^2, w_l(t) = 0.2 + 0.5 \left(\frac{t}{T}\right)^2 \quad (6)$$

The expressions for the low-frequency filter and high-frequency filter are given as:

$$\Lambda_l = 2e^{-10\lambda^2}, \Lambda_h = 2 - 2e^{-10\lambda^2} \quad (7)$$

The final expression for the dynamic filter $g(\Lambda)$ is:

$$g(\Lambda) = w_l \Lambda_l + w_h \Lambda_h \quad (8)$$

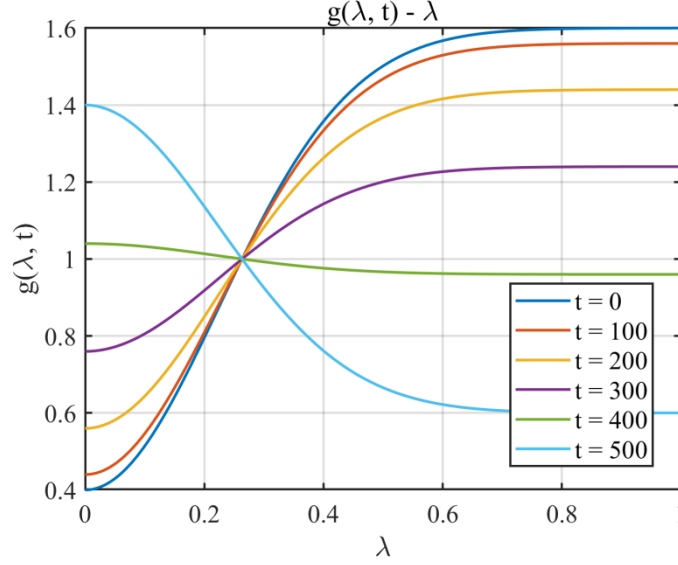


Fig.1 Schematic Diagram of the Dynamic Filter

By amplifying or suppressing specific frequency components, this filter dynamically adjusts the population distribution to balance global exploration and local exploitation. During the early iterations of the algorithm, the weight of the high-frequency components is higher, enhancing the local diversity of the population and promoting broad exploration of the solution space. In the later iterations, the weight of the low-frequency components increases, strengthening global stability and guiding the population to converge toward the optimal solution. **Fig. 1** illustrates the schematic diagram of the dynamic filter, showcasing its adaptation over iterations at steps 0, 100, 200, 300, 400, and 500. The diagram depicts the transition of the filter's shape, where high-frequency components are assigned greater weights in the early stages, gradually shifting to emphasize low-frequency components in later stages. This transition reflects the dynamic balance between exploration and exploitation, effectively guiding the optimization process through different phases.

2.2.2 Adaptive Sampling Functions

DGNE introduces two sampling functions, φ and ϕ , which optimize the population distribution before and after spectral filtering, respectively. Before spectral filtering, φ is used to generate new candidate solutions around the better-performing individuals via a Gaussian distribution. Its sampling formula is as follows:

$$\varphi(X) \sim N(\mu_1, \sigma_t^2) \quad (9)$$

where $\mu_1 = \frac{1}{k} \sum_{i=1}^k X_i$ represents the mean of the top k individuals sorted by fitness, $\sigma_t^2 =$

$0.0005e^{-16\frac{t-T}{T^2}}$ is the variance of the distribution, which decreases dynamically with the number of iterations to focus the search. Here, we set k as $[0.2N]$, where N represents the number of individuals in the population.

$$X^f = Ug(\Lambda)U^T\varphi(X) \quad (10)$$

where U and Λ are the eigenvectors and eigenvalues of the normalized Laplacian matrix L_{norm} , $g(\Lambda)$ is the dynamic filter. After spectral filtering, ϕ generates new candidate solutions by sampling around the better-performing individuals from the combined population before and after filtering. Its sampling formula is as follows:

$$\phi(X) \sim N(\mu_2, \sigma_t^2) \quad (11)$$

where the combined population $X^c = [X, X^f]$, $\mu_2 = \frac{1}{k} \sum_{i=1}^k X_i^c$ represents the mean of the top k individuals in the combined population sorted by fitness. Here, we set k as $[0.2N]$, where N represents the number of individuals in the population.

DGNE introduces dynamic filtering mechanisms and adaptive Gaussian sampling functions, providing an innovative optimization framework for evolutionary algorithms. The dynamic filtering mechanism adjusts the weights of high- and low-frequency components, enhancing exploration and local diversity in the early stages of the algorithm while improving exploitation and global stability in the later stages. Through two-stage sampling before and after spectral filtering, DGNE ensures that the population distribution maintains diversity while gradually focusing on high-fitness regions. Additionally, by leveraging the spectral properties of graphs, DGNE dynamically adjusts the population structure to capture both global stability and local diversity. **Fig. 2** illustrates the overall framework of the proposed DGNE, which includes two sampling processes and a dynamic filter. The filter transitions from high-frequency to low-frequency filtering, reflecting the shift from enhancing exploration capabilities in the early stages to strengthening exploitation capabilities in the later stages. This structure allows the model to effectively balance exploration and exploitation throughout the iterative process. The pseudocode provided in **Algorithm 1** outlines the implementation of DGNE, detailing how the population is dynamically updated through spectral filtering, resampling functions, and sampling operations. This algorithm serves as a comprehensive guide to the procedural steps involved in DGNE, ensuring clarity and reproducibility of the method.

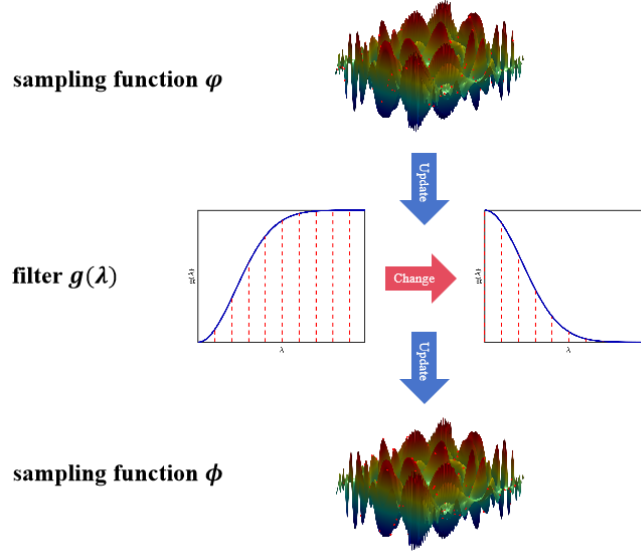


Fig.2 An illustration of the proposed DGNE

Algorithm 1 DGNE

Input: Population size N , Maximum iterations T , Objective Function f , Lower Bound LB , Upper Bound UB , sampling function φ, ϕ , Filter g

Initialization: Population X , optimal solution x_{best} , optimal function value f_{best} , $t = 0$

While $t \leq T$

Update the $\mu_1 = \frac{1}{k} \sum_{i=1}^k X_i$ and $\sigma_t^2 = 0.0005e^{-16\frac{t-T}{2}}$ of sampling function φ

Construct the sampling function φ

Apply $\varphi(X)$ to X for nonlinear transformation.

Calculate the similarity matrix A for the population X

Calculate the normalized Laplacian matrix L using $L = I - D^{-\frac{1}{2}}AD^{-\frac{1}{2}}$

Calculate the Diagonal eigenvalue matrix Λ and the eigenvector matrix U of the L

Update the weights $w_h(t) = 0.8 - 0.5\left(\frac{t}{T}\right)^2$, $w_l(t) = 0.2 + 0.5\left(\frac{t}{T}\right)^2$

Construct the Filter $g(\Lambda) = w_l\Lambda_l + w_h\Lambda_h$

Obtain the updated population X_{new} through $X_{new} = Ug(\Lambda)U^TX$

Obtain the combined population $X^c = [X, X_{new}]$

Update the $\mu_2 = \frac{1}{k} \sum_{i=1}^k X_i^c$ and $\sigma_t^2 = 0.0005e^{-16\frac{t-T}{2}}$ of sampling function ϕ

Construct the sampling function ϕ

Apply $\phi(X_{new})$ to X_{new} for nonlinear transformation.

$t = t + 1$

Update the optimal solution x_{best} and the optimal function value f_{best}

End

3.Experimental Results and Analysis

To comprehensively evaluate the performance of DGNE, extensive experiments were conducted on the CEC2017 benchmark suite, which includes 30 test functions with dimensionalities of 30, 50, and 100. The proposed method was compared against several state-of-the-art and classical evolutionary algorithms, including LSHADE, LSHADE_cnEpSin, MadDE, SaDE, EA4eig, DE, and CMA-ES. To ensure fairness, all algorithms were configured with a population size of 30 and a maximum iteration count of 500. Each algorithm was independently executed 30 times to ensure statistical reliability. The experimental results were analyzed using the Wilcoxon rank-sum test and the Friedman mean rank test to assess the significance and consistency of DGNE's performance. This section is organized as follows: Section 3.1 presents the overall performance comparison of DGNE against competing algorithms; Section 3.2 investigates the impact of varying population sizes on DGNE's performance; and Section 3.3 explores the behavior of DGNE beyond 500 iterations.

3.1 Overall Performance Comparison

To validate the performance of the DGNE, experiments were conducted on 30 benchmark functions from the CEC2017 test suite. The experiments were performed with dimensionalities of 30, 50, and 100 to comprehensively evaluate the scalability and optimization effectiveness of DGNE. The proposed method was compared with various high-performance and classical evolutionary algorithms, including LSHADE, LSHADE_cnEpSin, MadDE, SaDE, EA4eig, DE, and CMA-ES. To ensure fairness, the population size was set to 30, and the number of iterations was fixed at 500 for all algorithms. Each algorithm was independently run 30 times to ensure the statistical reliability of the results. Additionally, the Wilcoxon rank-sum test and Friedman mean rank test were employed to perform statistical analyses on the experimental results[52], verifying the significance and consistency of the algorithm's performance.

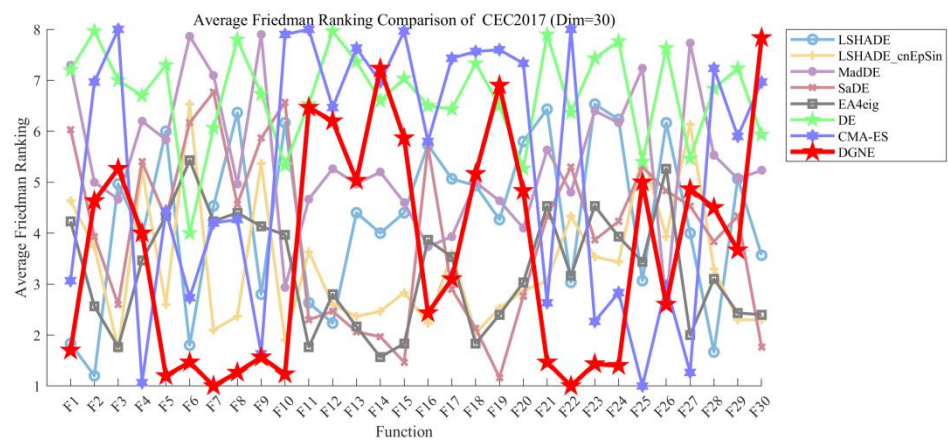
Table 1 Wilcoxon rank sum test statistical results

DGNE VS.	CEC 2017 (Dim=30)	CEC 2017 (Dim=50)	CEC 2017 (Dim=100)
LSHADE	14/6/10	17/4/9	21/2/7
LSHADE_cnEpSin	14/3/13	18/1/11	19/4/7
MadDE	18/3/9	21/4/5	25/1/4
SaDE	14/5/11	19/2/9	22/2/6
EA4eig	13/2/15	16/3/11	22/1/7
DE	23/5/2	27/0/3	28/0/2

CMA-ES	20/6/4	23/0/7	24/0/6
Overall (+/=-)	116/30/64	141/14/55	161/10/39

Table 2 Friedman mean rank test for all test suites

Suites	CEC 2017					
Dimensions	30		50		100	
Algorithms	Ave.	Overall	Ave.	Overall	Ave.	Overall
	Rank	Rank	Rank	Rank	Rank	Rank
LSHADE	4.29	5	4.17	4	3.92	3
LSHADE_cnEpSin	3.38	2	3.25	2	3.71	2
MadDE	5.41	7	5.65	7	6.11	7
SaDE	3.98	4	4.32	5	4.38	5
EA4eig	3.27	1	3.55	3	3.94	4
DE	6.72	8	6.65	8	6.30	8
CMA-ES	5.27	6	5.30	6	5.12	6
DGNE	3.68	3	3.13	1	2.52	1



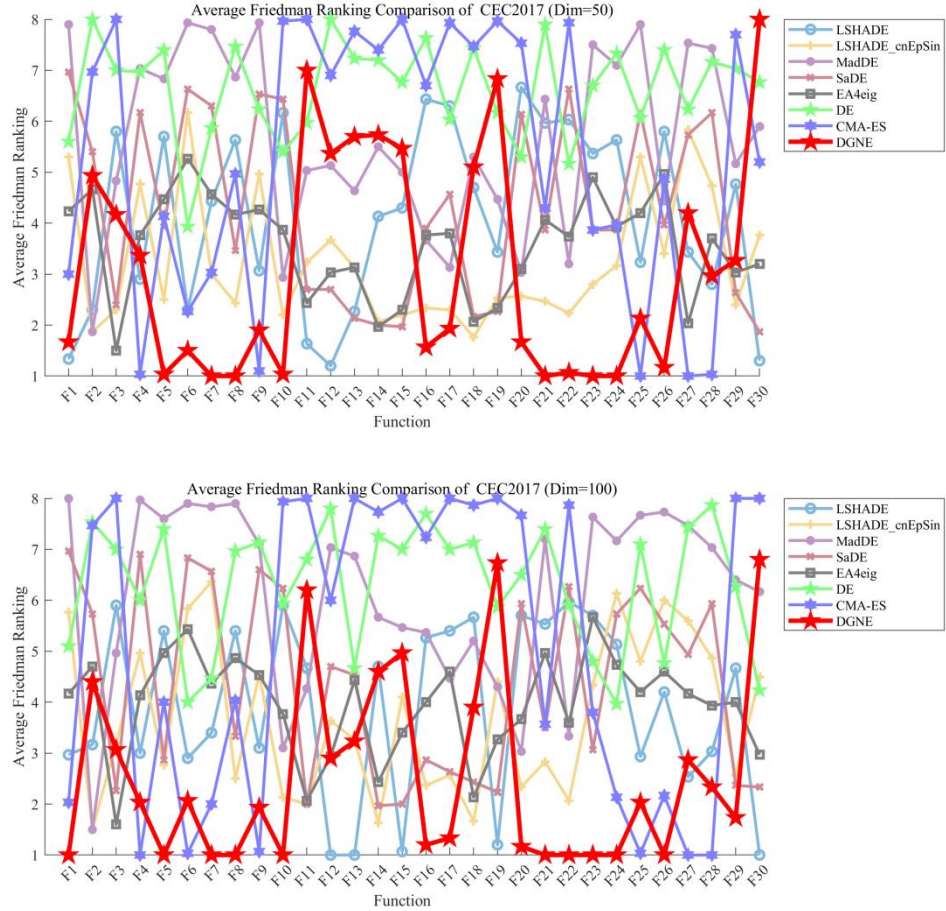


Fig.3 Average Friedman Rankings of DGNE and Competitors on CEC2017

Table 1 presents the results of the Wilcoxon rank-sum test, which evaluates the statistical performance of DGNE against several state-of-the-art and classical evolutionary algorithms on the CEC2017 benchmark suite (dimensions 30, 50, and 100). **Table 2** complements this by providing the Friedman mean rank test results, which rank the algorithms based on their average performance across the same test suite, offering a comprehensive evaluation of overall competitiveness. The results indicate that DGNE performs exceptionally well in medium- and high-dimensional tasks. In **Table 1**, DGNE achieves results of 141/14/55 and 161/10/39 at 50 and 100 dimensions, respectively, demonstrating its strong performance in complex optimization tasks. This trend is further validated in **Table 2**, where DGNE secures average rankings of 3.13 and 2.52 at 50 and 100 dimensions, respectively, ranking first overall in terms of comprehensive performance across different dimensions. For 30-dimensional tasks, DGNE still performs impressively with results of 116/30/64 in **Table 1**. Although it ranks third in **Table 2** with an average rank of 3.68, it is only behind LSHADE_cnEpSin and EA4eig, the champion algorithms of the CEC2017 and CEC2022 test suites, respectively. Notably, the Wilcoxon rank-sum test results show that DGNE's performance at 30 dimensions is very close to these two champion algorithms, highlighting its competitiveness in low-dimensional problems. Among the competitors, LSHADE_cnEpSin, the champion of the CEC2017 test suite, demonstrates reliability and robustness across all dimensions, particularly excelling in

high-dimensional tasks. For example, in **Table 1**, it achieves a result of 19/4/7 at 100 dimensions, and in **Table 2**, it consistently ranks second overall. Similarly, EA4eig, the champion of the CEC2022 test suite, shows its unique strengths in low-dimensional tasks. For instance, in 30-dimensional tasks, it ranks first in **Table 2** with an average rank of 3.27 and achieves 13/2/15 in **Table 1**. However, EA4eig's performance declines as the dimensionality increases, ranking third and fourth in **Table 2** at 50 and 100 dimensions, respectively, indicating its suitability for low-dimensional problems. LSHADE demonstrates adaptability across dimensions, achieving a result of 21/2/7 at 100 dimensions in **Table 1**, and steadily improving its rank in **Table 2**, moving from fifth at 30 dimensions to third at 100 dimensions, showcasing its scalability. SaDE also maintains stable performance, with results such as 14/5/11 at 30 dimensions in **Table 1** and consistently ranking fourth or fifth in **Table 2**. In conclusion, DGNE achieves undisputed first place in comprehensive performance at 50 and 100 dimensions and demonstrates strong competitiveness at 30 dimensions, only slightly behind LSHADE_cnEpSin and EA4eig, the two champion algorithms. Based on the combined results of **Table 1** and **Table 2**, DGNE exhibits outstanding scalability and adaptability, particularly excelling in high-dimensional tasks. Meanwhile, other algorithms such as LSHADE_cnEpSin in high-dimensional tasks and EA4eig in low-dimensional tasks also demonstrate their strengths in specific scenarios. **Fig. 3**, combined with the analysis of the CEC2017 test suite, highlights DGNE's performance across different function types and dimensionalities. DGNE performs exceptionally well on unimodal and multimodal functions (F1–F10) as well as on certain composition functions (F21–F26), particularly in high-dimensional problems, where its dynamic graph structure excels in exploitation and scalability. However, its performance declines on hybrid functions (F11–F15) and the highly complex composition function F30, reflecting its limitations in handling intricate, non-separable landscapes with diverse characteristics. Additionally, DGNE's effectiveness in low-dimensional tasks (e.g., 30 dimensions) is relatively weaker, where simpler algorithms like EA4eig and LSHADE are more efficient in navigating smaller search spaces. Overall, DGNE demonstrates strong potential for high-dimensional and specific problem types, but it requires further enhancement in exploration capabilities and adaptability to tackle complex hybrid functions and low-dimensional tasks effectively.

3.2 Impact of Population Size

As shown in **Table 3**, DGNE was evaluated alongside state-of-the-art algorithms using population sizes of 20, 30, 50, and 100. The experiments were conducted on the CEC2017 benchmark suite, comprising 90 functions across 30, 50, and 100 dimensions (30 functions per dimension). The number of iterations was fixed at 500 for these experiments. DGNE consistently achieved competitive rankings across all population sizes, with average ranks of 3.09 (population size = 20), 3.11 (population size = 30), 3.79 (population size = 50), and 3.33 (population size = 100). Notably, DGNE ranked 1st for population sizes of 20 and 30, 3rd for a population size of 50, and 2nd for a population size of 100. These results highlight DGNE's robustness to variations in population size. Among the competing algorithms, LSHADE_cnEpSin demonstrated strong performance, particularly at larger population sizes. For instance, it ranked 1st at population sizes of 50 and 100,

with average ranks of 2.98 and 2.53, respectively. This suggests that LSHADE_cnEpSin benefits from larger populations, likely due to its enhanced exploration capabilities. EA4eig also performed well, securing 2nd place at population sizes of 20 and 50, with average ranks of 3.41 and 3.41, respectively. However, its performance slightly declined at a population size of 100, where it ranked 3rd. LSHADE and SaDE maintained stable performance across all population sizes, consistently ranking 4th or 5th, indicating their reliability in diverse scenarios. On the other hand, MadDE, DE, and CMA-ES exhibited weaker performance, consistently ranking in the lower tiers across all population sizes. For example, DE ranked 8th in all cases, highlighting its limitations in handling larger populations and more complex optimization tasks.

Overall, DGNE's ability to adapt to varying population sizes while maintaining top-tier performance underscores its robustness and scalability. While LSHADE_cnEpSin and EA4eig also demonstrated strong adaptability, DGNE's consistent ranking across different configurations makes it a highly competitive choice for optimization tasks.

Table 3 Friedman Average Rankings of Different Algorithms with Varying Population Sizes

population size	20		30		50		100	
Algorithms	Ave. Rank	Overall Rank	Ave. Rank	Overall Rank	Ave. Rank	Overall Rank	Ave. Rank	Overall Rank
LSHADE	4.23	4	4.13	4	4.39	5	4.21	4
LSHADE_cnEpSin	3.83	3	3.44	2	2.98	1	2.53	1
MadDE	5.63	7	5.72	7	5.45	7	5.60	7
SaDE	4.55	5	4.23	5	3.91	4	4.26	5
EA4eig	3.41	2	3.59	3	3.41	2	3.74	3
DE	5.89	8	6.56	8	6.95	8	7.24	8
CMA-ES	5.36	6	5.23	6	5.13	6	5.11	6
DGNE	3.09	1	3.11	1	3.79	3	3.33	2

3.3 Behavior Beyond 500 Iterations

In **Table 4**, we extended the evaluation to include iteration counts of 500, 1000, and 2000, with the population size fixed at 30. The rankings were based on the CEC2017 benchmark suite with 90 functions (30 functions per dimension). DGNE demonstrated strong performance across all iteration counts, achieving average rankings of 3.11 (500 iterations), 3.45 (1000 iterations), and 3.47 (2000 iterations). DGNE ranked 1st at 500 iterations and 2nd at 1000 and 2000 iterations, underscoring its stability and competitiveness even as the number of iterations increased.

Among the competing algorithms, EA4eig emerged as a strong performer at higher iteration counts, ranking 1st at 1000 and 2000 iterations with average ranks of 3.09 and 3.17, respectively. This suggests that EA4eig benefits from extended exploration time, particularly in complex optimization tasks. LSHADE_cnEpSin also maintained competitive performance, ranking 2nd at 500 iterations and 3rd at 1000 and 2000 iterations. Its consistent performance across different iteration counts

highlights its robustness in balancing exploration and exploitation. LSHADE and SaDE showed stable performance, with LSHADE ranking 4th at 500 iterations and 5th at 1000 and 2000 iterations, with average ranks of 4.13, 4.62, and 4.59, respectively. SaDE demonstrated consistent performance, ranking 5th at 500 iterations and 4th at 1000 and 2000 iterations, with average ranks of 4.23, 4.29, and 4.46, respectively. On the other hand, MadDE, DE, and CMA-ES exhibited weaker performance. DE consistently ranked 8th across all iteration counts, with average ranks of 6.56, 6.61, and 6.42, highlighting its limitations in handling extended optimization processes. MadDE showed slight improvements, moving from 7th at 500 iterations to 6th at 2000 iterations, with average ranks of 5.72, 5.14, and 4.85. Similarly, CMA-ES improved from 6th at 500 iterations to 7th at 2000 iterations, with average ranks of 5.23, 4.78, and 4.95. Despite these minor improvements, both MadDE and CMA-ES remained in the lower tiers, indicating their challenges in maintaining competitiveness over longer optimization processes.

These experiments confirm DGNE's adaptability to different configurations, further validating its effectiveness as a robust optimization framework. The results demonstrate that DGNE maintains consistent performance across varying iteration counts, making it a reliable choice for a wide range of optimization tasks. While EA4eig and LSHADE_cnEpSin also demonstrated strong performance at higher iteration counts, DGNE's ability to remain competitive across all scenarios highlights its versatility and scalability. **Table 4** Friedman Average Rankings of Different Algorithms with Varying Iteration Counts

Table 4 Friedman Average Rankings of Different Algorithms with Varying Iteration Counts

iterations	500		1000		2000	
Algorithms	Ave. Rank	Overall Rank	Ave. Rank	Overall Rank	Ave. Rank	Overall Rank
LSHADE	4.13	4	4.62	5	4.59	5
LSHADE_cnEpSin	3.44	2	4.03	3	4.09	3
MadDE	5.72	7	5.14	7	4.85	6
SaDE	4.23	5	4.29	4	4.46	4
EA4eig	3.59	3	3.09	1	3.17	1
DE	6.56	8	6.61	8	6.42	8
CMA-ES	5.23	6	4.78	6	4.95	7
DGNE	3.11	1	3.45	2	3.47	2

3.4 Analysis of the Exploration and Exploitation

The balance between exploration and exploitation is a crucial concept in evolutionary computation. The exploration phase aims to search a broader solution space, while the exploitation phase focuses on refining the search to better approximate the optimal solution. Properly balancing these two phases is essential for enhancing the robustness of the algorithm[53]. In this section, we calculate

the percentages of exploration and exploitation using Eq. (12) and Eq. (13), respectively. **Fig.4** illustrates the evolution of exploration and exploitation proportions in the population, calculated using the following equations:

$$Exploration(\%) = \frac{Div(t)}{Div_{max}} \times 100 \quad (12)$$

$$Exploitation(\%) = \frac{|Div(t) - Div_{max}|}{Div_{max}} \times 100 \quad (13)$$

$$Div(t) = \frac{1}{dim} \sum_{d=1}^{dim} \frac{1}{n} \sum_{i=1}^n |median(x_d(t)) - x_{id}(t)| \quad (14)$$

The parameter $Div(t)$ denotes the dimension diversity measure and Div_{max} represents the maximum diversity achieved throughout the entire iteration process, which is calculated in Eq.(14). The results show that the exploration proportion is high in the early stages and decreases rapidly after 200 iterations, while the exploitation proportion is low initially and increases significantly after 200 iterations.

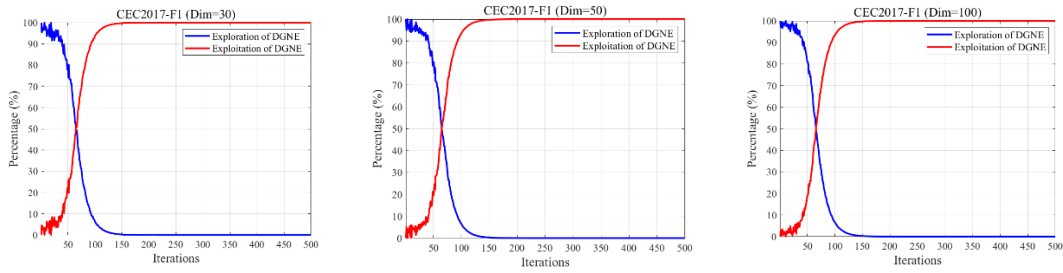


Fig.4 Exploration and Exploitation Analysis of DGNE

Fig.5 illustrates the low-frequency and high-frequency filter functions. It can be observed that the low-frequency filter function decreases as the eigenvalue λ increases, while the high-frequency filter function increases with λ . **Fig.1** shows the dynamically weighted filter function. As the iterations progress, the filter gradually transitions from a high-frequency-biased filter to a low-frequency-biased filter. Remarkably, this transition corresponds well with the exploration-to-exploitation shift observed in **Fig.4**, further validating the design of DGNE.

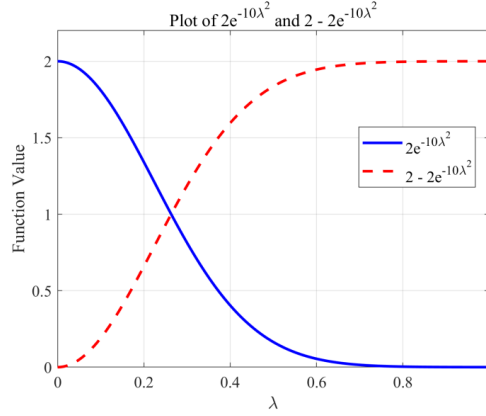


Fig.5 Illustration of Low-Frequency and High-Frequency Filter Functions

4. Conclusion

This paper proposes an innovative optimization framework, DGNE, which integrates GNNs with EAs. By introducing a dynamic filtering mechanism and adaptive Gaussian sampling functions, DGNE achieves an effective balance between global exploration and local exploitation during the optimization process. By dynamically adjusting the population distribution, DGNE demonstrates significant advantages in solving complex optimization problems, as validated by experiments on the CEC2017 benchmark suite. The experimental results show that DGNE outperforms or matches state-of-the-art algorithms such as LSHADE, LSHADE_cnEpSin, MadDE, SaDE, EA4eig, as well as classic algorithms like DE and CMA-ES, in terms of convergence speed, robustness, and adaptability. DGNE achieves the best average rankings in 50 and 100 dimensions and ranks second in 30 dimensions, highlighting its scalability and outstanding performance in high-dimensional optimization tasks. Furthermore, DGNE effectively leverages the spectral properties of graphs and dynamically balances exploration and exploitation, making it particularly effective in addressing large-scale and complex problems.

However, DGNE also has some limitations. In low-dimensional tasks, while DGNE remains competitive, there is still room for improvement, especially in enhancing its exploration capability in the early stages of the algorithm. Additionally, DGNE shows weaker performance on certain hybrid and highly complex composition functions, suggesting areas for further refinement in handling highly diverse and rugged landscapes. Moreover, the impact of population size and iteration count further influences DGNE's performance. DGNE has demonstrated robust adaptability across varying population sizes, consistently achieving top rankings with population sizes of 20, 30, 50, and 100. The framework ranked 1st for smaller populations (20 and 30), 2nd for a population size of 100, and 3rd for 50, underscoring its scalability in different configurations.

Similarly, DGNE's performance across different iteration counts (500, 1000, and 2000) further confirms its stability and competitiveness. The results demonstrate that DGNE remains consistently competitive, ranking 1st at 500 iterations and 2nd at higher iteration counts. This adaptability is a key strength of DGNE, making it a versatile optimization tool in various scenarios. Future research could focus on enhancing DGNE's exploration capabilities to further improve its performance on challenging multimodal and hybrid functions. This could include exploring the use of alternative filter functions, sampling functions, and adaptive sampling mechanisms. Moreover, optimizing its computational efficiency in low-dimensional tasks could improve its versatility. Exploring other advanced mechanisms, such as multi-objective optimization, parallelization, and integration with other machine learning techniques, could further expand DGNE's applicability in a wider range of optimization and artificial intelligence challenges.

In conclusion, DGNE offers a robust, scalable, and efficient optimization framework, demonstrating significant potential in advancing the integration of evolutionary algorithms and graph neural networks, and providing a promising direction for solving complex real-world problems. Its ability to maintain top-tier performance across varying population sizes and iteration counts, combined with its adaptability and scalability, makes DGNE a highly competitive choice for optimization tasks.

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