CHARACTERISTICS OF NETWORKS GENERATED BY KERNEL GROWING NEURAL GAS

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ABSTRACT

This research aims to develop kernel GNG, a kernelized version of the growing neural gas (GNG) algorithm, and to investigate the features of the networks generated by the kernel GNG. The GNG is an unsupervised artificial neural network that can transform a dataset into an undirected graph, thereby extracting the features of the dataset as a graph. The GNG is widely used in vector quantization, clustering, and 3D graphics. Kernel methods are often used to map a dataset to feature space, with support vector machines being the most prominent application. This paper introduces the kernel GNG approach and explores the characteristics of the networks generated by kernel GNG. Five kernels, including Gaussian, Laplacian, Cauchy, inverse multiquadric, and log kernels, are used in this study. The results of this study show that the average degree and the average clustering coefficient decrease as the kernel parameter increases for Gaussian, Laplacian, Cauchy, and IMQ kernels. If we avoid more edges and a higher clustering coefficient (or more triangles), the kernel GNG with a larger value of the parameter will be more appropriate.

KEYWORDS

Growing neural gas, kernel method, self-organizing map

1. INTRODUCTION

Today, the amount of data has grown enormously [1, 2]. To efficiently process such massive datasets, vector quantization methods are often used to reduce the number of data points. Thus, vector quantization methods have become increasingly important for handling large datasets.

Self-organizing maps (SOMs) and their alternatives are widely used vector quantization methods commonly applied in various fields such as data visualization, feature extraction, and data classification. These approaches encode a dataset into a set of interconnected units. Kohonen’s SOM is the most popular and widely used of these methods. Kohonen’s SOM creates a network with fixed topology, such as a d-dimensional lattice.

The growing neural gas (GNG) proposed by Fritzke [3] is an alternative to SOM. GNG can flexibly change the network topology during training. GNG can adapt not only the reference vectors but also the network topology to an input data set. GNG can gradually increase the number of neurons and reconstruct the network topology according to the input data. GNG is a useful method for extracting the topology of the input data. Thus, GNG can not only quantize a dataset but also preserve the topology of the dataset as the topology of the network.

The kernel method is useful for projecting data into a high-dimensional feature space. The support vector machine [4] gains good performance for non-linear data, applied kernel method. The kernel Kohonen’s SOM can perform better than Kohonen’s network [5]. However, a kernel
version of GNG has not yet been developed, and the characteristics of networks generated by
kernel GNG remain unknown.

This study aims to develop the kernel GNG and investigate the characteristics of networks
generated by the kernel GNGs with Gaussian, Laplacian, Cauchy, inverse multiquadric (IMQ),
and log kernels. First, the method of the kernel GNG is derived as shown in Sec.3. Second, the
paper shows the feature of networks generated by the kernel GNGs with these kernels in Sec.6.
This paper shows that the kernel GNGs with these kernels can generate networks that effectively
represent input datasets, similar to the original GNG algorithm.

2. RELATED WORK

The best-known and most widely used the SOM is Kohonen’s SOM. The SOM can project
multidimensional data onto a low-dimensional map [6]. The SOM is used in various applications
such as color quantization [7, 8], data visualization [9], and skeletonization [10]. The most
popular SOM is Kohonen’s SOM [11]. However, the network structure generated by Kohonen’s
SOM is static (generally an d-dimensional lattice) [12]. Thus, Kohonen’s SOM cannot flexibly
change the network topology depending on the input dataset.

The growing neural gas (GNG) [3] is a type of SOMs [13] and can find the topology of an input
distribution [14]. The network of the GNG is flexible, and its structure represents the data
structure. GNG has been widely applied to topology learning, such as the extraction of the two-
dimensional outline of an image [15, 16, 17], the reconstruction of 3D models [18], landmark
extraction [19], object tracking [20], anomaly detection [12], and cluster analysis [21, 22, 23].

The kernel method is often used for nonlinear separations. The most famous application of the
kernel method is the support vector machine [4, 24]. Many researchers have used the kernel
method to improve the performance of various methods. The kernel k-means [25] partitions the
data points into a higher dimensional feature space and can partition a dataset non-linearly.
Kohonen’s SOM has also been kernelized [26, 27, 28]. The kernel Kohonen’s SOM shows better
performance. However, the kernel GNG is not yet proposed.

3. KERNEL GROWING NEURAL GAS

The kernel growing neural gas (kernel GNG) is a modified version of the GNG that uses a kernel
function. The kernel GNG projects the dataset into a higher dimensional feature space using a
non-linear function and converts it into a network. The kernel trick allows the kernel GNG to
learn the topology of the input without the need for direct projection of the data into feature
space.

The kernel GNG consists of a set of units connected by a set of unweighted and undirected edges.
Each unit i has a weight \( w_i \in \mathbb{R}^d \) corresponding to a reference vector in the input space and a
cumulative error \( E_i \). Given a data point from the dataset \( X = \{ x_1, \ldots, x_n, \ldots, x_N \} \), where \( x_n \in \mathbb{R}^d \)
at each iteration, the kernel GNG updates the unit and the network.

Consider a data point \( x_n \), a unit weight \( w_i \), and a nonlinear mapping function \( \phi(\cdot) \) that maps \( x_n \)
and \( w_i \) to \( \phi(x_n) \) and \( \phi(w_i) \) in feature space. The dot product of the two points, \( \phi(x_n) \) and
\( \phi(w_i) \), is denoted as \( K(x_n, w_i) = \phi(x_n)^T \phi(w_i) \), where \( K(\cdot, \cdot) \) is the kernel function.
The winning unit $s_1$ of the kernel GNG is the one closest to an input data point $x_n$ in the feature space. The criterion to identify $s_1$ is the squared distance between $x_n$ and the weight $w_i$ of unit $i$ in the feature space. The squared distance $D^2(x_n, w_i)$ in the feature space is defined as follows:

$$D^2(x_n, w_i) = \| \phi(x_n) - \phi(w_i) \|^2 = K(x_n, x_n) - 2K(x_n, w_i) + K(w_i, w_i).$$

The kernel GNG identifies a winning unit by minimizing the squared distance between the mapped point and the mapped weight using the above equation. The winning unit $s_1$ with respect to an input $x_n$ is thus obtained by

$$s_1 = \arg\min_i D^2(x_n, w_i).$$

After that, the weight of the winning unit $w_{s_1}$ is updated according to the following rule:

$$w_{s_1}(t + 1) = w_{s_1}(t) - \varepsilon_{s_1} \frac{1}{2} \frac{\partial}{\partial w_{s_1}} D^2(x_n, w_{s_1}),$$

where $t$ is the iteration index and $\varepsilon_{s_1}$ is the learning rate of the winning unit $s_1$. This equation is based on gradient descent to minimize the squared distance $D^2(x_n, w_{s_1})$. Therefore, the update equation for the weight $w_{s_1}$ in the kernel GNG is as follows

$$w_{s_1}(t + 1) = w_{s_1}(t) - \varepsilon_{s_1} \frac{1}{2} \left( \frac{\partial}{\partial w_{s_1}} K(w_{s_1}, w_{s_1}) - \frac{2}{\partial w_{s_1}} K(w_{s_1}, x_n) \right).$$

This equation is consistent with that of the kernel SOM update rule proposed by [26]. This method eliminates the need to maintain the transformed weights and the transformed data points, allowing direct updating of the weights in the input space without updating the high-dimensional weights in the feature space.

Five kernel functions are used in this study, including Gaussian, Laplacian, Cauchy, inverse multiquadric (IMQ), and log kernels. Table 1 shows the kernelized $D^2(x, w)$ and $\frac{\partial}{\partial w} D^2(x, w)$. The code for the kernel GNG is openly available on GitHub (https://github.com/KazuhiroFujita/KernelGNG).

<table>
<thead>
<tr>
<th>kernel</th>
<th>$K(x, w)$</th>
<th>$D^2(x, w)$</th>
<th>$\frac{\partial}{\partial w} D^2(x, w)$</th>
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</thead>
<tbody>
<tr>
<td>Gaussian</td>
<td>$\exp\left(-\frac{|x-w|^2}{2\gamma^2}\right)$</td>
<td>$2\left(1 - \exp\left(-\frac{|x-w|^2}{2\gamma^2}\right)\right)$</td>
<td>$-2\frac{x-w}{\gamma^2}\exp\left(-\frac{|x-w|^2}{2\gamma^2}\right)$</td>
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<tr>
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<td>$2\left(1 - \exp\left(-\frac{|x-w|}{\gamma}\right)\right)$</td>
<td>$-\frac{2x-w}{\gamma}|x-w|\exp\left(-\frac{|x-w|}{\gamma}\right)$</td>
</tr>
<tr>
<td>Cauchy</td>
<td>$\frac{1}{1+|x-w|^2/\gamma^2}$</td>
<td>$2\left(1 - \frac{1}{1+|x-w|^2/\gamma^2}\right)$</td>
<td>$-\frac{4\gamma^2}{\gamma^2(1+|x-w|^2/\gamma^2)^2}\frac{x-w}{\gamma}$</td>
</tr>
<tr>
<td>IMQ</td>
<td>$\frac{1}{\sqrt{|x-w|^2 + \gamma^2}}$</td>
<td>$2\left(1 - \frac{1}{c\sqrt{|x-w|^2 + \gamma^2}}\right)$</td>
<td>$-\frac{2}{\gamma^2}|x-w|^2 + \gamma^2)^{3/2}$</td>
</tr>
<tr>
<td>log</td>
<td>$-\log(|x-w|^r + 1)$</td>
<td>$2\log(|x-w|^r + 1)$</td>
<td>$-2d(x-w)|x-w|^r + 1$</td>
</tr>
</tbody>
</table>
3.1. Algorithm of the Kernel GNG

The kernel GNG, based on the same principles as the original GNG algorithm, extracts the network structure from the input data, but uses kernelized equations. The algorithm is formulated as

1. Initialize the network with two connected neurons. Their weights are two randomly selected data points.
2. Randomly select an input data point \( x_n \) from the dataset.
3. Identify the winning unit \( s_1 \), the one closest to \( x_n \), as defined by
   \[
   s_1 = \text{argmin}_i D^2(x_n, w_i).
   \]
   where \( D^2 \) is shown in table 1. At the same time, find the second nearest unit, \( s_2 \).
4. Increment the ages of all edges connected to the winning unit \( s_1 \).
5. Increase the cumulative error \( E_{s_1}(t) \) by the squared distance between the input data point \( x_n \) and the weight of the winning unit \( w_{s_1} \):
   \[
   E_{s_1}(t + 1) = E_{s_1}(t) + D^2(x_n, w_{s_1}).
   \]
6. Adapt the winning unit \( s_1 \) and its neighbors \( j \) to better reflect the input data point \( x_n \) by updating their weights:
   \[
   w_{s_1}(t + 1) = w_{s_1}(t) - \varepsilon_{s_1} \frac{1}{2} \frac{\partial}{\partial w_{s_1}} D^2(x_n, w_{s_1}),
   \]
   \[
   w_j(t + 1) = w_j(t) - \varepsilon_{n} \frac{1}{2} \frac{\partial}{\partial w_j} D^2(x_n, w_j),
   \]
   where \( \frac{\partial}{\partial w} D^2(x, w) \) is in table 1.
7. If the units \( s_1 \) and \( s_2 \) are connected, reset the age of their connecting edge to zero. Otherwise, create an edge between them.
8. Discard any edges whose ages exceed the maximum age \( a_{\text{max}} \). If this leaves any units isolated, remove them.
9. Insert a new unit after every \( \lambda \) iteration:
   - Identify the unit \( q \) with the largest cumulative error \( E_q \).
   - Among the neighbors of \( q \), find the node \( f \) with the largest error.
   - Insert a new unit \( r \) between \( q \) and \( f \) as follows:
     \[
     w_r = (w_q + w_f)/2.
     \]
   - Create edges between neurons \( r \) and \( q \), and between \( r \) and \( f \), while removing the edge between \( q \) and \( f \).
   - Decrease the cumulative errors of \( q \) and \( f \) by multiplying them by a constant \( \alpha \), and initialize the cumulative error of \( r \) to the updated error of \( q \).
10. Multiply all cumulative errors by a constant \( \beta \) to reduce them.
11. Repeat from step 2 until the number of iterations reaches \( T \).
In this study we used $N_{\text{max}} = 100$, $a_{\text{max}} = 50$, $\lambda = 100$, $\alpha = 0.5$, $\beta = 0.995$, $\varepsilon_{s_1} = 0.2$, and $\varepsilon_n = 0.006$. These parameter settings are based on [3].

### 4. Evaluation Metrics for Kernel GNG Performance and Network Topology

#### 4.1. Evaluation metrics for kernel GNG

The effectiveness of the kernel GNG is evaluated using two different metrics. The first metric, mean square error (MSE), evaluates the average of the squared distances between each input data point and its nearest unit in the input space. This metric is expressed as

$$\text{MSE} = \frac{1}{N} \sum_{n=1}^{N} \min_i \| x_n - w_i \|^2,$$

where $x_n$ is an input data point, $w_i$ is the weight of neuron $i$, and $\| \cdot \|$ is the Euclidean norm.

The second metric, kernel mean square error (kMSE), extends the MSE by measuring the squared distance between the data points and their corresponding weight vectors in the transformed feature space defined by the kernel function. The kMSE is expressed as

$$\text{kMSE} = \frac{1}{N} \sum_{n=1}^{N} \min_i D^2(x_n, w_i).$$

In the above equation, $D^2(x_n, w_i)$ is the distance metric computed in the feature space between the input data point $x_n$ and the weight vector $w_i$. For more information about the kernel distance metric $D^2(x_n, w_i)$, see table 1.

#### 4.2. Network Analysis Metrics for Topology Evaluation

In the field of complex network research, measures are used to study the structure of networks. In this study, two measures are used to examine the generated networks: the average degree and the average clustering coefficient.

The average degree, denoted as $k$, quantifies the average number of edges per node. It is calculated using the following formula

$$k = \frac{1}{N} \sum_{i=1}^{N} k_i,$$

where $k_i$ is the degree (the number of edges) of node $i$.

On the other hand, the average clustering coefficient $C$ indicates how much nodes in the network tend to form connected triangles on average. It is given by
where $c_i$ is the clustering coefficient of node $i$. The clustering coefficient $c_i$ for node $i$ [29] is given by

$$c_i = \frac{2t_i}{k_i(k_i - 1)},$$

where $t_i$ is the number of triangles around $i$ and $k_i$ is the degree of $i$. If $k_i < 2$, $c_i$ is set to zero.

These metrics are derived using NetworkX, a comprehensive Python library tailored for network analysis.

5. EXPERIMENTAL SETTING

For this research, we used several Python libraries, namely NumPy for calculations related to linear algebra, NetworkX for handling network operations and computing coefficients, and scikit-learn for generating synthetic data.

Synthetic and real-world data sets are used to evaluate the characteristics of the network generated by the kernel GNG. The synthetic datasets include Square, Blobs, Circles, Moons, Swiss_roll, and S_curve. Square dataset is constructed using NumPy’s `random.rand` function, which generates two-dimensional data points uniformly distributed between 0 and 1. Blobs dataset is constructed using scikit-learn’s `datasets.make_blobs` function, which uses a Gaussian mixture model of three isotropic Gaussian distributions with default parameters. Circles dataset, created using the `datasets.make_circles` function with noise and scale parameters of 0.05 and 0.5, respectively, contains two concentric circles of data points. The Moons dataset, a distribution mimicking the shape of crescents, was created using the `datasets.make_moons` function with a noise parameter of 0.05. Swiss_roll and S_curve datasets are generated using `datasets.make_swiss_roll` and `datasets.make_s_curve`, respectively. Each synthetic dataset contains 1000 data points. In addition to these synthetic datasets, we also use two-dimensional datasets such as Aggregation [30], Compound [31], Pathbased [32], Spiral [32], D31 [33], R15 [33], Jain [34], Flame [35], and t4.8k [36]. The real-world datasets used are Iris, Wine, Ecoli, Glass, Yeast, Spam, CNAE-9, and Digits from the UCI Machine Learning Repository.

In all experiments performed, the pre-processing includes normalizing each data point $x_n = (x_{n1}, ..., x_{nd}, ..., x_{nD})$ in a dataset $X = \{x_1, ..., x_n, ..., x_N\}$ using the following formula:

$$x_n = \left(\frac{x_{n1} - \bar{x}_1}{\sigma_1}, ..., \frac{x_{nd} - \bar{x}_d}{\sigma_d}, ..., \frac{x_{nD} - \bar{x}_D}{\sigma_D}\right),$$

where $\bar{x}_d = \frac{1}{N} \sum_{n=1}^N x_{nd}$, and $\sigma_d = \sqrt{1/N \sum_{n=1}^N (x_{nd} - \bar{x}_d)^2}$.

6. RESULTS

In this section, we present four experimental results that demonstrate the effectiveness of kernel GNGs. First, we provide a 2-dimensional visualization of the networks generated by kernel GNGs, illustrating their structure and connectivity. Second, we present the evolution of MSE and kernel MSE over iterations $t$. Third, we explore the dependence of MSE and network structure on
the kernel parameter $\gamma$, revealing the role of this parameter in shaping the network. Finally, we describe the characteristics of the network generated by kernel GNGs.

### 6.1. Visualization of Networks Generated by GNG and Kernel GNGs

Figure 1 shows the networks generated from synthetic datasets by the GNG and kernel GNGs. The kernels used for the kernel GNGs include the Gaussian kernel with $\gamma = 1.8$, the Laplacian kernel with $\gamma = 1.8$, the IMQ kernel with $\gamma = 1.8$, the Cauchy kernel with $\gamma = 1.8$, and the log kernel with $\gamma = 3$. All networks are derived with $\text{END} = 2 \times 10^4$, and the random seed is set to 1. In all cases, the networks generated by kernel GNGs accurately reflect the input topology. However, the Laplacian kernel produces a significantly more complex network structure, but it spreads the units over the input topology. This result shows the ability of kernel GNGs to effectively extract the topology of the dataset.

![Figure 1](image)

**Figure 1.** This figure shows networks generated from synthetic data by the GNG and the kernel GNGs with Gaussian, Laplacian, IMQ, Cauchy, and log kernels. Data points are shown as gray dots, while the network units are shown as black dots, with their positions indicating the reference vectors. Black lines mark the edges of the networks.
6.2. Convergence of MSE and kMSE for Kernel GNGs

Figure 2 shows the convergence patterns of both the MSE of the kernel GNG and the GNG over iterations. For Blobs and Iris, the MSEs corresponding to all kernel GNGs begin to converge at about 10^4 iterations. In contrast, for Wine, the MSE of the kernel GNG using the Laplacian and the log starts to converge at about $2 \times 10^4$ iterations, while the others continuously and slowly decrease even after $2 \times 10^4$ iterations. For Wine, the MSEs of the kernel GNGs are larger than that of the GNG, except when the log kernel is used. For Ecoli, the MSEs associated with the kernel GNGs with the Laplacian, Cauchy, and IMQ kernels converge at $4 \times 10^4$ iterations. The kernel GNGs with Gaussian and logarithmic kernels converge at $2 \times 10^5$ and $10^5$ iterations, respectively. For Ecoli, the convergence values of the kernel GNGs with all kernels are larger than those of the GNG.

Figure 3 shows the convergence behavior of the kMSE over iterations for the kernel GNGs. For Blobs and Iris, the kMSE for all kernel GNGs starts to converge at about $10^4$ iterations. In contrast, for Wine, the kMSE approaches a low value at about $2 \times 10^4$ iterations. For Ecoli, while the kMSE reaches low values at $10^4$ iterations, it exhibits a slow and continuous decline after this iteration. These observations suggest that the kernel GNGs reach appropriately low MSE and kMSE values around $2 \times 10^4$ iterations.

Figure 2. Evolution of the MSE over iterations. Subfigures (A) to (D) show the MSE for different data sets: (A) Blobs, (B) Iris, (C) Wine, and (D) Ecoli, respectively. Each data point represents the average kMSE obtained from 10 independent runs, all initialized with random values.
Figure 3. Evolution of the kernel mean square error (kMSE) over iterations. Subfigures (A) to (D) show the kMSE for different data sets: (A) Blobs, (B) Iris, (C) Wine, and (D) Ecoli, respectively. The values shown are the averages derived from 10 independent runs, each initialized with random values.

6.3. Influence of Kernel Parameters on Network Characteristics in kernel GNGs

Figure 4 shows the dependence of the kernel parameters on the MSE, the average degree, and the average clustering coefficient of the networks generated by the kernel GNG with Gaussian, Laplacian, Cauchy, and IMQ kernels. The MSE, average degree, and average clustering coefficient are computed from the network at $4 \times 10^5$ iterations. Interestingly, as the kernel parameters increase, the MSE, the average degree, and the average clustering coefficient decrease. The kernel GNG with the Laplacian kernel tends to generate a network characterized by a higher degree and a higher clustering coefficient than the others.

Figure 5 shows the effect of the kernel parameters on the MSE, the average degree, and the average clustering coefficient of the networks generated by the kernel GNG with log kernel, with the computations ending at the $4 \times 10^5$ iterations. A noteworthy observation is the absence of any discernible dependence of the network features on the kernel parameter of the kernel GNG with log kernel.
Figure 4. Dependence of various network metrics on kernel parameters for networks generated by kernel GNG with Gaussian, Laplacian, Cauchy, and IMQ kernels for Blobs, Iris, Wine, and Ecoli datasets. Subfigures (A), (D), (G), and (J) show the dependence of the MSE on the kernel parameter. Subfigures (B), (E), (H), and (K) show the average degree, while subfigures (C), (F), (I), and (L) show the average clustering coefficient. Each value shown represents an average derived from 10 independent runs, each initialized with random values.
Figure 5. Dependence of various network metrics on kernel parameters in networks generated by kernel GNG with log kernel and GNG for Blobs, Iris, Wine, and Ecoli datasets. Subfigures (A), (D), (G), and (J) show the influence of kernel parameters on the MSE. Subfigures (B), (E), (H), and (K) show the average degree, while subfigures (C), (F), (I), and (L) show the average clustering coefficient. Each displayed value represents the average derived from 10 independent runs, each initialized with random values.

6.4. Comparison of Network Characteristics in kernel GNGs

Table 2 gives a comprehensive overview of the average degree $N_d$ of the networks generated by the GNG and the kernel GNGs with the Gaussian, Laplacian, Cauchy, IMQ, and log kernels. The Gaussian, Laplacian, Cauchy, and IMQ kernels use a $\gamma$ parameter value of 1.8. In contrast, the log kernel uses a $\gamma$ parameter value of 3. Each row represents a dataset, and the corresponding data dimension is documented in the second column, denoted by $D$. For networks generated by the kernel GNG using the Gaussian, Cauchy, and IMQ kernels, $N_d$ is less than or equal to that of GNG. While the Laplacian kernel often produces a larger $N_d$ than the GNG, especially for two-dimensional datasets, it produces a smaller $N_d$ for datasets such as CNAE and Digits. In many cases, the log kernel’s $N_d$ is equal to or smaller than the GNG’s, although it is larger than the GNG’s values for datasets such as Wine, Spam, Glass, Yeast, and Digits.

In parallel, table 3 shows the average clustering coefficient $C$ of the networks generated by the GNG and the kernel GNGs. The values of $C$ for the kernel GNGs using Gaussian, Cauchy, and
IMQ kernels are often less than or equal to those of the GNG. The Laplacian kernel often produces a \( C \) larger than the GNG’s for lower dimensional data. In addition, the log kernel’s \( C \) tends to be higher than the other kernels for data points with larger dimensions.

In summary, Gaussian and Cauchy kernels typically produce networks of equal or reduced complexity compared to the GNG. The IMQ kernel tends to produce simpler networks, the Laplacian kernel produces more complex networks, and the log kernel produces more complex networks, especially for high-dimensional data.

7. CONCLUSIONS

This paper describes the kernel growing neural gas (GNG) and investigates the characteristics of the networks it generates. Several kernel functions are tested, including Gaussian, Laplacian, IMQ, Cauchy, and log kernels. The results show that the reference vectors produced by the kernel GNG match the input dataset, which is confirmed by the sufficiently small mean square error (MSE) values. However, the topology of the network generated by kernel GNG depends on the kernel function parameter \( \gamma \). The average degree and the average clustering coefficient decrease as \( \gamma \) increases for Gaussian, Laplacian, Cauchy, and IMQ kernels.

The choice between kernel GNG and GNG is complex, mainly because the only discernible difference from our results is in the metrics of the network topology. However, if we avoid more edges and a higher clustering coefficient (or more triangles), kernel GNG with a larger value of \( \gamma \) will be more appropriate. Such a feature will be particularly valuable for 3D graphics applications, where the kernel GNG may be able to simplify the mesh structure of polygons for a more efficient rendering.

Table 2: The average degree of a network, \( N_d \), generated by GNG and kernel GNGs

<table>
<thead>
<tr>
<th>dataset</th>
<th>( D )</th>
<th>GNG</th>
<th>Gaussian</th>
<th>Laplacian</th>
<th>Cauchy</th>
<th>IMQ</th>
<th>Log</th>
</tr>
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<tbody>
<tr>
<td>Square</td>
<td>2</td>
<td>4.24</td>
<td>3.97</td>
<td>\textbf{6.58}</td>
<td>4.09</td>
<td>3.92</td>
<td>4.18</td>
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<tr>
<td>Blobs</td>
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<td>\textbf{10.89}</td>
<td>3.91</td>
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<tr>
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<tr>
<td>Moons</td>
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<td>\textbf{6.76}</td>
<td>3.02</td>
<td>2.79</td>
<td>2.93</td>
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<td>4.01</td>
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<td>S_curve</td>
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<td>4.19</td>
<td>3.98</td>
<td>\textbf{4.92}</td>
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<td>3.67</td>
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<td>2.51</td>
<td>2.82</td>
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<td>5.31</td>
<td>5.05</td>
<td>\textbf{6.45}</td>
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</table>

\( D \) indicates the dimension of a data point. The Gaussian, Cauchy, and IMQ kernels are used with a \( \gamma \) parameter set to 1.8, while the logarithmic kernel uses a \( \gamma \) parameter set to 3. The purities are the mean of 10 runs with random initial values. The largest and the smallest values are bold and italic, respectively.
Table 3: Average clustering coefficient of a network generated by GNG and kernel GNGs

<table>
<thead>
<tr>
<th>dataset</th>
<th>$D$</th>
<th>GNG</th>
<th>Gaussian</th>
<th>Laplacian</th>
<th>Cauchy</th>
<th>IMQ</th>
<th>log</th>
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</table>

$D$ indicates the dimension of a data point. The purities are the mean of 10 runs with random initial values. The largest and the smallest values are bold and italic, respectively.

REFERENCES


Kazuhisa Fujita received a Ph.D. degree from the University of Electro-Communications in 2007. He was a Research Associate at the Department of Computer and Information Engineering, National Institute of Technology, Tsuyama College, in 2007-2013. He was an Assistant Professor at National Institute of Technology, Tsuyama College, in 2013-2018. He was also a Visiting Fellow at the University of Electro-Communications in 2011-2022. From 2018, he worked as an Associate Professor at Komatsu University. His current research interests include game-playing artificial intelligence, neural networks, a clustering method, and the free energy principle.