

MIK: Modified Isolation Kernel for Biological Sequence Visualization, Classification, and Clustering

Presented By SARWAN ALI

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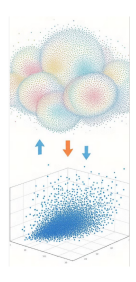
Introduction

- Predicting the impact of amino acid changes on protein function is essential for applications such as disease variant classification and protein engineering
- Computing pairwise sequence similarity becomes more important for
 - $\bullet \ \, {\sf Supervised \ analysis} => {\sf Protein \ function \ prediction} \\$
 - Unsupervised Analysis => Pattern recognition
 - Data visualization
- The t-distributed stochastic neighbor embedding (t-SNE):
 - It is a method for interpreting high dimensional (HD) data by mapping each point to a low dimensional (LD) space (usually two-dimensional)
 - Used for better visualization
 - Dimensionality reduction
- Overall, Sequence analysis is:
 - Crucial for understanding evolutionary relationships
 - Infer the functional and structural properties
 - Identifies disease-causing mutations and drug targets

Background: t-SNE and Dimensionality Reduction

While t-SNE captures overall structure well, it may struggle to preserve local structure efficiently. This limitation prompted our research into alternative approaches

- High-dimensional Data
 - Complex datasets with multiple features
- t-SNE Processing
 - Computes pairwise similarities and reduces dimensionality
- Visualization
 - Represents data in 2D or 3D space for analysis



Background: Traditional methods

- Traditional analytical approaches (Sequence Alignment or Phylogenetic Analysis, etc.) fails,
 - Building a tree out of a large dataset can be difficult
 - Computationally intensive and time-consuming
 - Not general purpose in using for supervised analysis, unsupervised analysis, and visualization

Alternatives

Supervised/Unsupervised Analysis

- Machine Learning (ML)/ Deep Learning (DL) techniques
- Challenges:
 - We need to convert sequences into a format suitable for ML/DL models
- Solution:
 - Sequence Representation (Embeddings)
 - Convert the molecular sequences into a numerical format

Visualization:

Using different kernels and initialization techniques

Motivations and Goals

The vast global spread of pandemics like COVID-19, pushing viral sequence analysis into the "Big Data" realm Motivations:

- Understanding the immune invasion and host-to-host transmission properties of SARS-CoV-2 and its variants
- Knowledge of mutations and variants will help identify transmission patterns - facilitate public health measures
- This will also help in vaccine design and efficacy

Goals:

- High dimensionality data in biological sequences
 - Better low-dimensional Visualization
 - Improve performance and reduce computational cost

Old Before New

Visualization

• t-SNE => Gaussian kernel, Isolation kernel

Supervised Analysis

 Classification => feature engineering, kernel functions, neural networks, LLMs

Unsupervised Analysis

• Clustering => DBSCAN, ...

Research Gap

Visualization

- Better preservation of neighborhood for visualization
- Fast computational time

Supervised Analysis

- Higher predictive performance
- Preserve maximum information in low dimensional embeddings

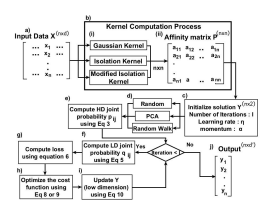
Unsupervised Analysis

- Better grouping and cluster separations
- Preserve maximum information in low dimensional embeddings

Our (High Level) Idea => Visualization

- We propose the Modified Isolation Kernel (MIK), as an alternative to the Gaussian and Isolation kernel
- It is intended to address the existing shortcomings in preserving local and global structures and handling noisy data and outliers
- MIK is evaluated using a variety of initialization techniques
 - Random initialization
 - PCA-based initialization
 - Random walk-based initialization
- The random walk-based initialization for such biological data is not been explored in the literature

Workflow



Problem Formulation

- Given dataset $X = \{x_1, x_2, ..., x_n\}$ in \mathbb{R}^d
- Assume a dataset $Y = \{y_1, y_2, ..., y_n\}$
- Objective: Map $X \in \mathbb{R}^d$ to $Y \in \mathbb{R}^{d'}$, such that d' < d
- d' = 2 or 3
- The similarity between points is preserved as much as possible
- Goal: Map points from X to Y such that the probability distribution between P_{ij} and Q_{ij} are as close as possible
- The similarity between a pair of points x_i, x_j in the higher dimensional space is represented by a probability P_{ij}
- The similarly for low dimensional space points y_i, y_j is represented by Q_{ij}

Gaussian Kernel

$$K(x_i, x_j) = exp(\frac{-||x_i - x_j||^2}{2\sigma_i^2})$$

Isolation Kernel

It measures the isolation of a data point from its neighbors

1 Pairwise Squared Distances:

$$D_{ij} = \|x_i - x_j\|^2$$



Isolation Kernel (Contd.)

2 Perplexity and Effective Neighborhood Size:

$$H(P_i) = \log \left(\sum_j P_{ij} \right) + \beta \frac{\sum_j D_{ij} P_{ij}}{\sum_j P_{ij}}$$

- $H(P_i)$ represents the entropy of P_i for the *i*-th point
- $P_{ij} = \exp(-\beta D_{ij})$ represents the similarity between points x_i and x_j

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Isolation Kernel (Contd.)

3 Scaling Parameter (Beta) Adjustment:

$$\beta = \frac{\mathsf{Perplexity}}{\mathsf{distance scaling} \cdot (\mathsf{max}(D_i) + \epsilon)}$$

This beta adjustment is repeated until $H(P_i) \approx \log(\text{Perplexity})$

Isolation Kernel (Contd.)

4 Kernel Matrix Construction:

$$P_{ij} = \frac{P_{ij}}{\sum_{j} P_{ij}}$$

where each row in ${\cal P}$ sums to 1, providing a probabilistic interpretation of the kernel



MIK

The Modified Isolation Kernel incorporates two additional components: a distance scaling factor and weights for each point

• The *distance scaling* factor is computed based on the average pairwise distance between all points in the dataset $X = \{x_1, x_2, \dots, x_n\}$:

$$D_{ij} = \|x_i - x_j\|$$

where D_{ij} is the Euclidean distance between points x_i and x_j . The distance scaling factor s is then calculated as:

$$s = \frac{1}{n(n-1)} \sum_{i=1}^{n} \sum_{j \neq i} D_{ij}$$

This scaling factor is used to normalize the distances, ensuring that the neighborhood sizes are comparable across different datasets

MIK (Contd.)

• Weights Adjustment: After normalizing P_i for each point i, weights w_i are applied to modulate the values in P:

$$P_{ij} = \frac{P_{ij}}{\sum_{j} P_{ij}} \cdot w_i$$

where:

ullet w_i is a weight applied to row i to adjust its influence on the kernel

Weights Computation Using DBSCAN

- The weights are computed based on the density of points around each data point, using the **DBSCAN** (Density-Based Spatial Clustering of Applications with Noise) algorithm.
- The DBSCAN algorithm assigns a weight (or label) to each point according to its neighborhood density. The parameters for DBSCAN include:
 - ullet Epsilon ϵ : the maximum distance between two points to be considered neighbors
 - min samples: the minimum number of points required in a neighborhood for a point to be considered a core point

Weights Computation Using DBSCAN (Contd.)

- ullet Given the dimensionality of the data d, the minimum samples parameter min samples is set to d + 1
- \bullet This ensures that the density estimation accounts for the dimensionality of the dataset, with ϵ chosen based on dataset-specific properties
- The weights w_i for each point x_i are defined as the labels assigned by DBSCAN:

$$w_i = \mathsf{DBSCAN}(x_i)$$

where points labeled as noise by DBSCAN receive a weight of -1.

• These weights can then be incorporated into the Isolation Kernel to emphasize regions with higher or lower point densities

DBSCAN-based Weights

$$w_i = \begin{cases} \text{cluster label of } x_i & \text{if } x_i \text{ is in a dense region} \\ -1 & \text{if } x_i \text{ is labeled as noise} \end{cases}$$

 This approach helps to adjust the kernel by accounting for both the average distances and the density-based clustering within the dataset

Kernel Comparison

Isolation Kernel:

$$P_{ij} = \frac{\exp(-\beta D_{ij})}{\sum_{j} \exp(-\beta D_{ij})}$$

Modified Isolation Kernel with Distance Scaling and Weights:

$$P_{ij} = \frac{\exp(-\beta D_{ij}^{\text{scaled}})}{\sum_{j} \exp(-\beta D_{ij}^{\text{scaled}})} \cdot w_{i}$$

Where:

- $oldsymbol{\circ}$ eta is adjusted iteratively to match the perplexity
- $D_{ij}^{\text{scaled}} = D_{ij}$ · distance-scale adjusts the distance, and w_i applies optional point-specific weights

t-SNE Evaluation Metrics

• Neighborhood Agreement (NA):

$$NA = 1 - \frac{2}{N(N-1)} \sum_{i=1}^{N} \sum_{j \neq i} \left| \frac{d_{ij}^{H} - d_{ij}^{L}}{d_{ij}^{H} + d_{ij}^{L}} \right|$$
 (1)

Trustworthiness (TW) :

$$TW = 1 - \frac{2}{N \cdot k \cdot (2N - 3k - 1)} \sum_{i=1}^{N} \sum_{j \in R_k} (R_{ij} - R_{ij}^L)$$
 (2)



Classification Evaluation Metrics

- We use Support Vector Machine (SVM), Naive Bayes (NB), Multi-Layer Perceptron (MLP), K-Nearest Neighbour (KNN), Random Forest (RF), Logistic Regression (LR), and Decision Tree (DT) classifiers
- We use average accuracy, precision, recall, weighted, and ROC area under the curve (AUC) as evaluation metrics for measuring the goodness of classification algorithms

Clustering Evaluation Metrics

- Silhouette Coefficient [1]
- Calinski-Harabasz Index [2]
- Davies-Bouldin Index [3]

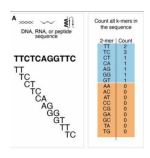
Dataset

Dataset	Seg.	Classes	Sequence Length			Detail	
Dataset	Jeq.		Max	Min	Mean	- Betain	
Protein Subcellular [4]	5959	11	3678	9	326.27	The unaligned protein sequences having information about subcellular locations.	
GISAID [5]	7000	22	1274	1274	1274.00	The aligned spike sequences of the SARS-CoV-2 virus having the information about the Lineage of each sequence.	
Nucleotide [6]	4380	7	18921	5	1263.59	Unaligned nucleotide sequences to classify gene family to which humans belong	

Embedding Methods

- Spike2Vec [7]
- Spaced k-mer [8]
- PWM2Vec [9]

Spike2Vec



k-mers spectrum ¹

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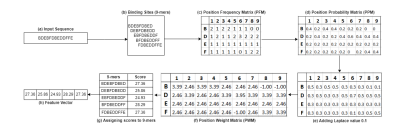
¹https://www.sciencedirect.com/science/article/pii/\$2001037024001703

Spaced k-mers

Gapped 3-mers for Sequence: ADCEFGHIK

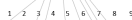
	Original k-mer	Gap at 1st	Gap at 2nd	Gap at 3rd
1	ADC	-DC	A-C	AD-
2	DCE	-CE	D-E	DC-
3	CEF	-EF	C-F	CE-
4	EFG	-FG	E-G	EF-
5	FGH	-GH	F-H	FG-
6	GHI	-HI	G-I	GH-
17	HIK	-IK	H-K	HI-

PWM2Vec



Sliding Window

BFEFEBDE**BFDBEDDFF**DBBDDEFED



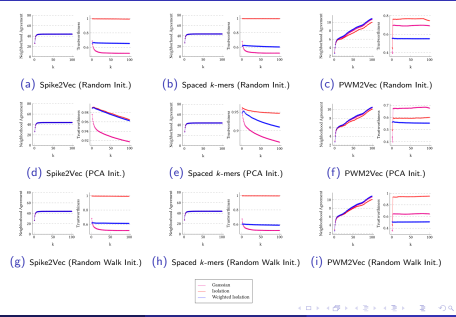
- B 3.39 2.46 3.39 3.39 2.46 2.46 2.46 -1.00 -1.00
- D 2.46 3.39 2.46 2.46 3.39 3.95 3.39 3.39 3.39
- F 2.46 2.46 2.46 2.46 -1.00 3.39 3.39 3.39

Absolute score = 3.39 + 2.46+...+ 3.39 = 28.28

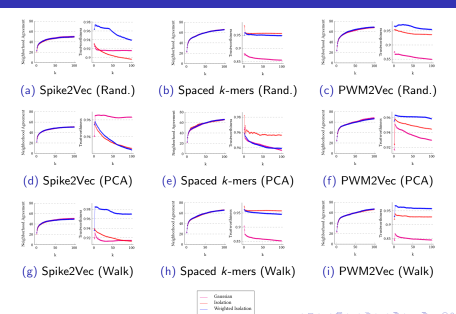
Initialization Methods

- Random Initialization
- PCA-based Initialization
- Random Walk-based Initialization

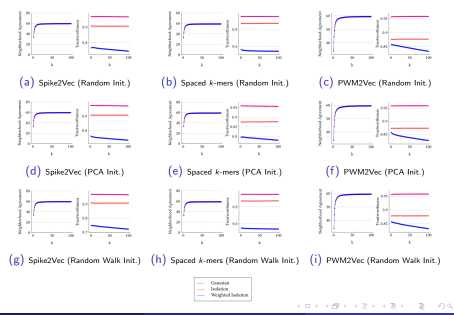
Results - Nucleotide Dataset



Results - GISAID Dataset



Results - Protein Subcellular Dataset



Classification Results - Protein Subcellular Dataset

Kernel	Embeddings	Algo.	Acc. ↑	Prec. ↑	Recall ↑	F1 (Weig.) ↑	F1 (Macro)	ROC AUC ↑	Train Time (sec.) ↓
Gaussian	PWM2Vec	SVM NB MLP KNN RF LR DT	0.5207 0.3876 0.4640 <u>0.5634</u> 0.5252 0.5085 0.3668	0.5190 0.4438 0.4424 0.5671 <u>0.6064</u> 0.4830 0.3702	0.5207 0.3876 0.4640 0.5634 0.5252 0.5085 0.3668	0.5110 0.3965 0.4487 0.5550 0.4835 0.4654 0.3677	0.3947 0.3206 0.2990 <u>0.4567</u> 0.3176 0.2709 0.2597	0.6630 0.6376 0.6196 0.6950 0.6215 0.6106 0.5978	9.7228 0.0554 9.8790 0.1219 6.8668 0.4873 0.6648
Isolation	Spike2Vec	SVM NB MLP KNN RF LR DT	0.6493 0.2701 0.7796 <u>0.8546</u> 0.8345 0.2450 0.6913	0.6550 0.3615 0.7727 <u>0.8595</u> 0.8538 0.0600 0.6973	0.6493 0.2701 0.7796 <u>0.8546</u> 0.8345 0.2450 0.6913	0.6487 0.2285 0.7752 0.8522 0.8069 0.0964 0.6919	0.4977 0.2426 0.6105 <u>0.7464</u> 0.5968 0.0358 0.5203	0.7326 0.6714 0.7917 <u>0.8622</u> 0.7712 0.5000 0.7430	9.7372 <u>0.0455</u> 11.8214 0.1707 5.9505 0.2048 0.6885
Modified Isola- tion (Ours)	Spaced k-mers	SVM NB MLP KNN RF LR DT	0.8697 0.4010 0.8937 0.6119 0.8982 0.2204 0.7265	0.8767 0.5894 0.8971 0.6231 0.8947 0.2584 0.7377	0.8697 0.4010 0.8937 0.6119 <u>0.8982</u> 0.2204 0.7265	0.8712 0.3824 0.8949 0.5914 0.8867 0.0811 0.7301	0.7327 0.3661 <u>0.7847</u> 0.4010 0.7125 0.0337 0.5701	0.8663 0.7481 0.8907 0.6817 0.8443 0.5003 0.7793	1.1591 <u>0.0364</u> 10.0736 0.1299 6.3723 0.2525 1.0005

Classification Results - GISAID Dataset

Kernel	Embeddings	Algo.	Acc. ↑	Prec. ↑	Recall ↑	F1 (Weig.) ↑	F1 (Macro)	ROC AUC ↑	Train Time (sec.) ↓
Gaussian	Spike2Vec	SVM NB MLP KNN RF LR DT	0.7397 0.1564 0.7696 0.7678 0.7872 0.7174 0.7756	0.6998 0.6631 0.7308 0.7725 <u>0.7809</u> 0.6514 0.7702	0.7397 0.1564 0.7696 0.7678 0.7872 0.7174 0.7756	0.7009 0.2226 0.7388 0.7634 <u>0.7762</u> 0.6686 0.7668	0.3060 0.1312 0.4025 0.5223 <u>0.5342</u> 0.2447 0.5106	0.6541 0.5745 0.7111 0.7560 0.7637 0.6226 0.7550	3.4177 0.0866 8.2361 0.1034 2.8961 1.2611 0.2956
Isolation	Spaced k-mers	SVM NB MLP KNN RF LR DT	0.4972 0.0276 0.6499 0.6490 0.7029 0.4883 0.6379	0.3674 0.2183 0.6187 0.6460 <u>0.6777</u> 0.2385 0.6457	0.4972 0.0276 0.6499 0.6490 0.7029 0.4883 0.6379	0.3814 0.0271 0.6276 0.6356 0.6799 0.3205 0.6404	0.1007 0.0314 0.1884 0.2063 <u>0.3469</u> 0.0298 0.2828	0.5331 0.5203 0.5922 0.5892 0.6488 0.5000 0.6329	11.9771 0.1053 14.1174 0.1331 6.2961 0.6894 0.8870
Modified Isola- tion (Ours)	PWM2Vec	SVM NB MLP KNN RF LR DT	0.6820 0.5952 0.6741 0.6559 0.7103 0.4748 0.6709	0.6764 0.6176 0.6552 0.6493 <u>0.6955</u> 0.2255 0.6752	0.6820 0.5952 0.6741 0.6559 0.7103 0.4748 0.6709	0.6762 0.5962 0.6584 0.6453 0.6983 0.3057 0.6717	0.3117 0.2017 0.2493 0.2222 0.3612 0.0293 0.3168	0.6446 0.6008 0.6194 0.5966 0.6619 0.5000 0.6514	12.7645 0.1153 13.1291 0.1637 12.8492 0.6317 1.6763

Classification Results - Nucleotide Dataset

Kernel	Embeddings	Algo.	Acc. ↑	Prec. ↑	Recall ↑	F1 (Weig.) ↑	F1 (Macro)	ROC AUC ↑	Train Time (sec.) ↓
		SVM	0.3688	0.6767	0.3688	0.2707	0.2116	0.5447	1.0749
		NB	0.1317	0.7136	0.1317	0.1384	0.1435	0.5411	0.0125
	C 1	MLP	0.3857	0.7391	0.3857	0.2926	0.2315	0.5533	3.1893
Gaussian	Spaced	KNN	0.2697	0.3876	0.2697	0.2560	0.2308	0.5560	0.0560
	k-mers	RF	0.4295	0.6904	0.4295	0.3664	0.3188	0.5873	2.0117
		LR	0.3612	0.7508	0.3612	0.2478	0.1825	0.5360	0.1341
		DT	0.4285	0.6625	0.4285	0.3663	0.3198	0.5880	0.2364
		SVM	0.3213	0.3221	0.3213	0.3079	0.2588	0.5728	3.8166
		NB	0.2323	0.4337	0.2323	0.1836	0.2043	0.5560	0.0123
		MLP	0.5275	0.5203	0.5275	0.5192	0.4616	0.6858	7.3494
Isolation	Spike2Vec	KNN	0.5283	0.5340	0.5283	0.5275	0.4976	0.7093	0.0585
		RF	0.7469	0.7539	0.7469	0.7452	0.7345	0.8314	3.4126
		LR	0.3105	0.0965	0.3105	0.1472	0.0677	0.5000	0.0795
		DT	0.6151	0.6161	0.6151	0.6150	0.5866	0.7605	0.3282
		SVM	0.5798	0.5762	0.5798	0.5728	0.5358	0.7273	2.6021
Modified		NB	0.2604	0.3666	0.2604	0.2363	0.2463	0.5700	0.0080
Modified Isola-	C I	MLP	0.6207	0.6208	0.6207	0.6178	0.5734	0.7498	7.1481
tion	Spaced k-mers	KNN	0.5050	0.5093	0.5050	0.5041	0.4755	0.6954	0.0611
	K-mers	RF	0.7481	0.7689	0.7481	0.7455	0.7366	0.8316	4.1538
(Ours)		LR	0.3096	0.1191	0.3096	0.1466	0.0679	0.5001	0.0931
		DT	0.6237	0.6257	0.6237	0.6237	0.5959	0.7644	0.3863

Clustering Results - Protein Subcellular Dataset

Kernel	Embeddings	Algo.	Silhouette ↑	Calinski ↑	Davies ↓
	Spike2Vec	K-means Agglomerative K-Modes	0.101796 0.078076 -0.359683	360.804549 322.784851 1.646458	3.622216 3.585005 1.119053
Gaussian	Spaced k-mers	K-means Agglomerative K-Modes	0.126160 0.122294 -0.309347	428.163587 389.211763 4.658780	3.500538 3.418970 1.068371
	PWM2Vec	K-means Agglomerative K-Modes	0.061646 0.019724 -0.261891	271.209939 229.333382 1.060538	3.586904 4.163685 1.035597
	Spike2Vec	K-means Agglomerative K-Modes	0.015248 0.100906 -0.406924	58.296491 66.592869 0.230669	2.205289 2.267826 1.709086
Isolation	Spaced k-mers	K-means Agglomerative K-Modes	0.894164 0.894826 -0.494872	49.226179 54.900365 3.841355	0.064535 2.231084 2.039936
	PWM2Vec	K-means Agglomerative K-Modes	0.427347 0.892029 -0.382076	57.304634 64.086967 0.146916	2.029489 1.472744 1.543540
Modified	Spike2Vec	K-means Agglomerative K-Modes	0.100461 0.258273 -0.337161	875.4463 856.6747 0.404064	1.249022 1.088766 1.154124
Isola- tion (Ours)	Spaced k-mers	K-means Agglomerative K-Modes	0.298419 0.845733 -0.491851	766.048709 807.127025 48.420757	1.279214 1.059536 2.083230
	PWM2Vec	K-means Agglomerative K-Modes	0.191296 0.295157 -0.358505	626.270386 594.130597 0.255160	1.425956 1.278188 1.504934

Clustering Results - GISAID Dataset

Kernel	Embeddings	Algo.	Silhouette ↑	Calinski ↑	Davies ↓
	Spike2Vec	K-means Agglomerative K-Modes	0.725709 0.728701 -0.724895	2360.1012 2401.6734 84.479174	0.803562 0.897672 1.082510
Gaussian	Spaced k-mers	K-means Agglomerative K-Modes	0.671649 0.697097 -0.526676	1758.287930 1750.801213 70.972746	0.582548 0.535291 1.769435
	PWM2Vec	K-means Agglomerative K-Modes	0.691450 0.660800 -0.520338	1435.243293 1401.089635 94.359037	0.693299 0.818952 2.462654
	Spike2Vec	K-means Agglomerative K-Modes	0.068962 0.597965 -0.594450	122.328499 130.992233 0.137340	0.912677 2.127489 2.041841
Isolation	Spaced k-mers	K-means Agglomerative K-Modes	0.926955 0.935912 -0.624410	85.446835 90.096434 0.397808	1.213278 0.796774 2.425513
	PWM2Vec	K-means Agglomerative K-Modes	0.942670 0.940357 -0.671301	90.830309 91.914837 0.143632	0.032672 0.929066 2.922499
Modified	Spike2Vec	K-means Agglomerative K-Modes	0.062111 0.069198 -0.530450	191.373940 185.944651 1.363172	2.674008 2.621923 1.438185
Isola- tion (Ours)	Spaced k-mers	K-means Agglomerative K-Modes	0.139867 0.134961 -0.461139	689.139460 688.940686 0.695404	1.760740 1.658960 1.294385
	PWM2Vec	K-means Agglomerative K-Modes	0.078648 0.058910 -0.377294	322.821055 312.395730 1.832262	2.010462 2.072378 1.199501

Clustering Results - Nucleotide Dataset

Kernel	Embeddings	Algo.	Silhouette ↑	Calinski ↑	Davies ↓
	Spike2Vec	K-means Agglomerative K-Modes	0.845873 0.849057 -0.616155	100.442214 121.928330 0.680251	0.099883 0.097623 12.304512
Gaussian	Spaced k-mers	K-means Agglomerative K-Modes	0.849962 0.851768 -0.710487	110.207433 122.335197 0.595153	0.097384 0.096128 12.299829
	PWM2Vec	K-means Agglomerative K-Modes	0.906833 0.907149 -0.783775	1216.9008 1242.8203 19.695747	0.950717 1.207521 1.991363
	Spike2Vec	K-means Agglomerative K-Modes	0.266999 0.142199 -0.398666	85.054216 95.157338 0.214001	2.373702 2.552974 1.951997
Isolation	Spaced k-mers	K-means Agglomerative K-Modes	0.059934 0.045584 -0.361089	79.041871 86.883693 0.390651	2.562520 2.917264 1.595999
	PWM2Vec	K-means Agglomerative K-Modes	0.493470 0.928548 -0.535966	63.606950 93.383507 6.971815	1.599440 0.510479 1.929305
Modified	Spike2Vec	K-means Agglomerative K-Modes	0.326686 0.217707 -0.527625	16050.3293 15024.6706 6.309529	0.781717 1.035089 1.787152
Isola- tion (Ours)	Spaced k-mers	K-means Agglomerative K-Modes	0.445578 0.414100 -0.618379	4740.6237 4529.5243 0.269577	0.496505 0.507057 2.518221
	PWM2Vec	K-means Agglomerative K-Modes	0.906386 0.906386 -0.486018	85715.7221 85715.7222 0.072697	0.201059 0.201059 2.195824

Runtime

Kernel	Protein Subcellular	GISAID	Nucleotide
Gaussian	89.12 sec.	6.69 sec.	1.81 sec.
Isolation	135.81 sec.	103.50 sec.	29.80 sec.
MIK	3.34 sec.	2.91 sec.	1.02 sec.
MIK vs. Gaussian % improvement	96.25%	56.50%	43.64%
MIK vs. Isolation % improvement	97.54%	97.18%	96.57%

Recommended Initialization strategies

Dataset	Best Performing	Worst Performing	
Protein Subcellular	Random Walk	Random	
GISAID	Random Walk	Random	
Nucleotide	Random Walk	Random	

 Recommendation for initialization method based on the summary of performance on different datasets.

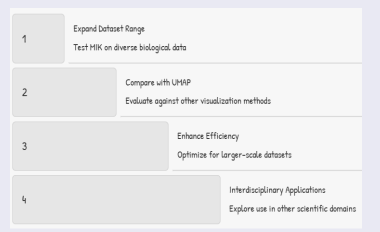
Key Findings and Implications

- MIK as an alternative to the Gaussian kernel, which is built upon the concept of the Isolation Kernel
- MIK uses adaptive density estimation
- Several initialization techniques were evaluated.



Future Directions

While our research demonstrates the potential of MIK, there are several avenues for future work



Thank You

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