# HI-PMK: A Data-Dependent Kernel for Incomplete Heterogeneous Data Representation

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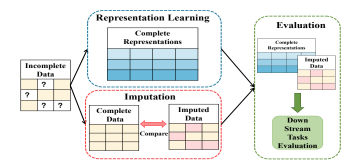
Abstract. Handling incomplete and heterogeneous data remains a central challenge in real-world machine learning, where missing values may follow complex mechanisms (MCAR, MAR, MNAR) and features can be of mixed types (numerical and categorical). Existing methods often rely on imputation, which may introduce bias or privacy risks, or fail to jointly address data heterogeneity and structured missingness. We propose the Heterogeneous Incomplete Probability Mass Kernel (HI-PMK), a novel data-dependent representation learning approach that eliminates the need for imputation. HI-PMK introduces two key innovations: (1) a probability mass-based dissimilarity measure that adapts to local data distributions across heterogeneous features (numerical, ordinal, nominal), and (2) a missingness-aware uncertainty strategy (MaxU) that conservatively handles all three missingness mechanisms by assigning maximal plausible dissimilarity to unobserved entries. Our approach is privacy-preserving, scalable, and readily applicable to downstream tasks such as classification and clustering. Extensive experiments on over 15 benchmark datasets demonstrate that HI-PMK consistently outperforms traditional imputation-based pipelines and kernel methods across a wide range of missing data settings. Code is available at: github.com/echoid/Incomplete-Heter-Kernel

## 1 Introduction

Missing data is a common challenge in real-world, data-driven applications, caused by factors such as data collection errors, survey non-responses, or system malfunctions [3]. These missing values can occur across all data types—numerical, categorical, or heterogeneous—making data analysis more complex and degrading machine learning model performance, often leading to biased or sub-optimal outcomes. Existing methods for handling missing data fall broadly into two categories: imputation-based approaches and representation learning approaches (see Figure 1).

Imputation-based methods estimate missing values using observed data, creating imputed complete datasets for downstream analysis. These methods are often preferred due to their intuitive evaluation process, where imputed data can be directly compared to the original complete dataset. However, they face two significant limitations: (i) revealing original data during imputation raises privacy concerns, and (ii) access to complete datasets is often impractical, necessitating reliance on indirect downstream metrics, which can obscure the evaluation of imputation quality.

**Representation learning methods**, in contrast, bypass imputation entirely by learning robust representations directly from incomplete



**Figure 1.** Comparing imputation and representation learning approaches for incomplete data handling.

data for downstream tasks. This approach mitigates privacy risks, reduces computational overhead, and eliminates dependence on complete datasets for evaluation. By focusing on meaningful representations, these methods provide a secure, efficient, and adaptable framework for handling incomplete datasets.

Another major limitation in existing methods is their reliance on assumptions about data types and missing mechanisms. Most traditional techniques are designed for numerical data [14, 40, 10] and extend to heterogeneous datasets through preprocessing steps like one-hot encoding, which increase dimensionality and computational cost [1]. Furthermore, while many methods assume data is Missing Completely at Random (MCAR), real-world data often follows more complex patterns, such as Missing Not at Random (MNAR) or Missing at Random (MAR) [30]. Failing to address these mechanisms can result in biased outcomes [25].

Our contribution: We propose Heterogeneous Incomplete Probability Mass Kernel (HI-PMK)—a novel, data-dependent kernel method that computes pairwise similarities directly from the observed portions of the data. HI-PMK supports both numerical and categorical features, models uncertainty without imputation, and accommodates arbitrary missing data mechanisms (MCAR, MAR, MNAR). Our approach provides a privacy-preserving, representation learning framework that is model-agnostic and readily applicable to downstream tasks such as classification and clustering. Extensive experiments on over 15 benchmark datasets with varying missing rates and mechanisms show that HI-PMK consistently outperforms both imputation-based pipelines and kernel baselines. The proposed method is simple, scalable, and robust-offering a practical alternative to conventional approaches for learning from incomplete heterogeneous data.

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### 2 Related Works

Imputation Methods for Incomplete Data. Conventional techniques—such as mean substitution, MICE, k-NN, and matrix factorization [10, 40, 17, 12]—are simple but struggle with heterogeneous features and often distort downstream distributions. Generative approaches based on VAEs [22, 16, 26], GANs [41, 2], or diffusion models [38, 9] model complex uncertainty, but typically assume homogeneous data, require large training samples, and are not task-aware. Recent empirical studies [46] show that such models often underperform on small tabular datasets. Surveys [23, 37] confirm these limitations, particularly in handling mixed-type features and diverse missingness mechanisms. These challenges motivate imputation-free strategies like representation learning.

Representation Learning for Incomplete Data. Instead of filling in missing values, representation learning directly encodes incomplete inputs into task-specific embeddings. Popular approaches include autoencoders [8] for reconstruction and graph neural networks [45, 7, 21] for propagating partial observations. These methods excel in complex domains, e.g., multimodal [19] adb temporal where structural dependencies aid representation. However, in tabular datasets with mixed types, unstructured missingness, and limited samples, deep models can overfit or generalize poorly. In such scenarios, lightweight methods like probabilistic modeling [17] and similarity-based estimators [29] often remain competitive.

Kernel-Based Approaches for Incomplete Data. Kernel methods offer a non-parametric alternative by modeling similarities in latent space without requiring complete input. Early variants adapt Gaussian or polynomial kernels through observed-feature reweighting or regularization [35, 27, 6, 33, 34], but typically assume numerical inputs and neglect missingness structure. More recent work estimates similarity directly from partial data, including affinity learning with kernel correction [43], low-rank matrix recovery [44], and online estimation [42]. Although effective in spectral clustering or matrix completion, these methods often rely on scaling, kernel tuning, and assume real-valued inputs that limiting their application in mixed-type tabular settings. Our HI-PMK fills this gap by enabling type and uncertainty-aware similarity estimation without such assumptions.

#### 3 Problem Formulations

Let  $X \in \mathbb{R}^{m \times n}$  denote a dataset with m instances and n features, potentially containing missing values. We decompose X into an observed component  $X^{\mathbf{0}}$  and a missing component  $X^{\mathbf{m}}$ , with a binary mask  $M \in \{0,1\}^{m \times n}$  indicating the missingness pattern:  $M_{ij} = 1$  if  $X_{ij}$  is missing, and 0 otherwise. The missingness process is governed by latent parameters  $\Psi$ , which encode factors that influence whether an entry is observed or missing. This relationship can be formalized as a conditional distribution:

$$f(\boldsymbol{M} \mid \boldsymbol{X}, \Psi),$$

where M is potentially dependent on both observed and unobserved parts of X, depending on the missingness mechanism.

The missing mechanism, determined by  $\Psi$ , is typically categorized into the following types:

Missing Completely at Random (MCAR). The probability of a value being missing is entirely independent of both observed ( $X^0$ ) and missing ( $X^m$ ) data. The missingness depends only on  $\Psi$  and can be expressed as:

$$f(\boldsymbol{M} \mid \Psi) \forall \boldsymbol{X}.$$

For example, in a healthcare dataset, some patients may miss follow-up medical tests due to random scheduling conflicts. Here, the observed part  $(\boldsymbol{X^0})$  includes attributes like age and previous test results, while the missing part  $(\boldsymbol{X^m})$  comprises the unrecorded follow-up test results.

**Missing At Random (MAR).** The probability of missingness depends only on the observed data  $X^0$ . This can be expressed as:

$$f(M \mid X^{0}, \Psi) \forall X^{m}$$
.

For instance, in the same healthcare dataset, older patients may be less likely to attend follow-up medical tests. Here, the missingness depends on the observed age  $(X^0)$  but is unrelated to the actual test results  $(X^m)$ .

**Missing Not At Random (MNAR).** The probability of missingness directly depends on the values of the missing data  $X^m$ . This is expressed as:

$$f(M \mid X^{m}, \Psi) \forall X^{o}$$
.

For example, in the same healthcare dataset, patients with very poor test results might avoid follow-up visits due to fear or reluctance. In this case, the missingness depends directly on the test results  $(\boldsymbol{X}^m)$ , making it an MNAR mechanism.

# 4 Data-Dependent Kernel

Data-dependent kernels enhance similarity computation by incorporating local data distributions into pairwise comparisons [4, 5, 39, 49], enabling better handling of heterogeneous features and capturing nuanced relationships. However, existing methods are not well equipped to operate under incomplete data. To address this, we propose the Heterogeneous Incomplete Probability Mass Kernel (HI-PMK), a novel data-dependent kernel designed to handle both heterogeneity and missingness. In HI-PMK, the H-component models data type diversity, while the I-component captures missingness-aware uncertainty, enabling robust and flexible similarity estimation.

#### 4.1 Probability Mass Kernel (PMK)

The Probability Mass Kernel (PMK) builds upon the concept of  $m_0$ -dissimilarity [4, 28], extending traditional distance metrics by incorporating the local data distribution surrounding the objects being compared. Unlike data-independent kernels, such as Gaussian or Laplacian kernels—where the similarity between two points depends solely on their spatial distance—PMK adjusts similarity based on the density of data in the surrounding region. For example, in sparse regions, two points at the same distance may be considered more similar than in densely populated regions. This adaptability allows PMK to capture complex structures and patterns in data, resulting in more accurate similarity measures.

Let  $X \in \mathbb{R}^{m \times n}$  be a dataset with m instances and n features. The i-th instance can be denoted as a vector  $\mathbf{x}^{(i)} = \langle x_1^{(i)}, x_2^{(i)}, \dots, x_n^{(i)} \rangle$ , where  $i \in [1, m]$ . For a given feature k, let  $R_k(x_k^{(i)}, x_k^{(j)})$  represent the region covering instances  $\mathbf{x}^{(i)}$  and  $\mathbf{x}^{(j)}$  for feature k. Specifically, the size of this region,  $|R_k(x_k^{(i)}, x_k^{(j)})|$ , represents the number of data points whose k-th feature values lie within this range. This count quantifies local data density, enhancing the similarity measure by incorporating contextual information.

The  $m_0$ -dissimilarity between  $\mathbf{x}^{(i)}$  and  $\mathbf{x}^{(j)}$  is calculated as:

$$m_0(\mathbf{x}^{(i)}, \mathbf{x}^{(j)}) = \left(\frac{1}{n} \sum_{k=1}^n \log \frac{|R_k(x_k^{(i)}, x_k^{(j)})|}{m}\right)$$
 (1)

Intuitively, if many data points fall within the region, the two instances are considered more dissimilar, as a dense data distribution surrounds them. Conversely, sparse regions yield small dissimilarity scores.

To efficiently compute  $|R_k(x_k^{(i)}, x_k^{(j)})|$ , each feature k is discretised into b bins, and a pre-computed bin data mass is used. A matrix stores the data masses between all bin pairs for each feature, enabling fast lookups to determine the region size and significantly speeding up the computation process. Finally, the  $m_0$  score is normalized to obtain the PMK:

$$PMK(\mathbf{x}^{(i)}, \mathbf{x}^{(j)}) = \frac{2 \cdot m_0(\mathbf{x}^{(i)}, \mathbf{x}^{(j)})}{m_0(\mathbf{x}^{(i)}, \mathbf{x}^{(i)}) + m_0(\mathbf{x}^{(j)}, \mathbf{x}^{(j)})}.$$
 (2)

This normalization ensures symmetry and self-similarity, aligning PMK with Mercer kernel requirements and enabling its integration with kernel-based learning frameworks.

# 4.2 PMK for Heterogeneous Data

The original PMK was limited to numerical data. To handle mixed data types, it was extended with the **H-Component**, enabling similarities across numerical and categorical features. This enhanced framework is called H-PMK.

work is called H-PMK. In Equation 1,  $\frac{|R_k(x_k^{(i)},x_k^{(j)})|}{m}$  is the probability mass in the region, denoted as  $P(R_k(x_k^{(i)},x_k^{(j)}))$ , which is calculated by discretizing the values of the numerical features. For categorical features,  $P(R_k(x_k^{(i)},x_k^{(j)}))$  can be computed using probabilities of categorical labels. The computation depends on whether the categorical feature k is **ordinal** (where values follow a natural order, such as size = S, M, L, XL, XXL}) or **nominal** (where values have no inherent order, such as color = Red, Green, Blue, Yellow, White}), as outlined below:

$$P(R_k(x_k^{(i)}, x_k^{(j)})) = \begin{cases} \sum_{z_k = \min(x_k^{(i)}, x_k^{(j)})}^{\max(x_k^{(i)}, x_k^{(j)})} P(z_k), & \text{ordinal } k \\ z_k = \min(x_k^{(i)}, x_k^{(j)}) & & & (3) \\ P(x_k^{(i)} \vee x_k^{(j)}), & \text{nominal } k \end{cases}$$

where  $P(z_k)$  represents the frequency of the feature value  $z_k$  divided by the total number of instances m, and  $P(x_k^{(i)} \lor x_k^{(j)})$  denotes the probability of a feature k having the label of  $x_k^{(i)}$  or  $x_k^{(j)}$ .

## 4.3 PMK for Incomplete Data

The **I-Component** extends PMK to handle incomplete data by addressing missing values under various mechanisms. This enhancement, called I-PMK, ensures robust performance on incomplete datasets.

**Separated Bucket**  $\mathcal{B}_k$  **for Missingness Frequency.** Since PMK relies on probability mass computed from the data distribution, we treat missing values as a distinct category. For each feature k, we introduce a special bucket  $\mathcal{B}_k$  that collects all missing entries in that feature. The size of this bucket, denoted  $|\mathcal{B}_k|$ , reflects the empirical frequency of missingness and is later used to adjust pairwise similarity computations involving missing values.

Adjusting Probability Mass Estimation for Incomplete Entries (Maximizing Uncertainty). To compute  $P(R_k(x_k^{(i)}, x_k^{(j)}))$  when one or both values are missing, we adopt a conservative strategy that reflects the maximum plausible dissimilarity under uncertainty. Specifically, we approximate the probability mass using the largest possible region consistent with the observed data distribution. This design aligns with the principle of Maximizing Uncertainty (MaxU), which treats missing values as potentially taking any value in the feature space.

Only one of them is missing: For any numeric or ordinal feature k, we treat them similarly since numerical data is converted into ordinal through discretization. In this context,  $x_k$  represents the observed value in the pair we are analyzing, while '?' denotes the missing value. The region size  $|R_k(x_k,?)|$  can be estimated by examining the data masses between the bin containing  $x_k$  ( $Bin(x_k)$ ) and the bins representing the first (minimum value) and last (maximum value) in the feature range. We let  $\mathcal{M}_L(x_k)$  and  $\mathcal{M}_R(x_k)$  represent the data masses to the left and right of  $Bin(x_k)$ , respectively, including the mass of  $Bin(x_k)$  itself (as shown in Figure 2). Since '?' denotes a missing value and can correspond to any possible value, we take a conservative approach named Maximize Uncertainty (MU) and assign the maximal possible dissimilarity as follows:

$$|R_k(x_k,?)| = \max(\mathcal{M}_L(x_k), \mathcal{M}_R(x_k)) + |\mathcal{B}_k| \tag{4}$$

If k is a nominal feature,  $|R_k(x_k, ?)|$  is computed based on the frequencies of nominal labels:

$$|R_k(x_k, ?)| = \mathcal{M}(x_k) + \max_{a \in \mathcal{S}_k} \mathcal{M}(a) + |\mathcal{B}_k|$$
 (5)

Here,  $\mathcal{M}(x_k)$  is the frequency of the observed label  $x_k$ , and  $\mathcal{S}_k$  represents the set of all possible nominal labels for feature k. The term  $\max_{a \in \mathcal{S}_k} \mathcal{M}(a)$  accounts for the most frequent label, assuming '?' could correspond to it, resulting in maximal dissimilarity. In both cases, we include the frequency of missing values  $(|\mathcal{B}_k|)$  to ensure that the contribution of missing data is properly accounted for, reflecting their potential to take on any value.

**Both of them are missing:** When both entries are missing, we adopt a conservative strategy to approximate their maximum possible dissimilarity. For numerical or ordinal features, the worst-case assumption is that the two missing values differ maximally, leading to:

$$|R_k(?,?)| = m, (6)$$

where m denotes the total number of instances in the dataset. For nominal features, since the missing entries may belong to two different categories, we estimate the maximal disagreement by summing the frequencies of the two most common categories:

$$|R_k(?,?)| = \max_{a \in S_k} \mathcal{M}(a) + \max_{b \in S_k \setminus \{a\}} \mathcal{M}(b) + |\mathcal{B}_k|,$$
 (7)

where  $S_k$  is the set of possible categories for feature k,  $\mathcal{M}(a)$  denotes the frequency of label a, and  $|\mathcal{B}_k|$  is the number of missing entries in feature k. This formulation ensures that completely missing pairs are penalized with maximal dissimilarity, while preserving consistency across feature types.

# 4.4 Methodology Analysis

**Separate Bucket**  $\mathcal{B}_k$ . The bucket  $\mathcal{B}_k$  stores all missing entries for feature k and plays a key role in capturing implicit patterns under structured missingness. Under MCAR, where missingness occurs

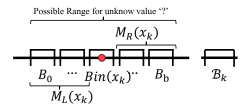
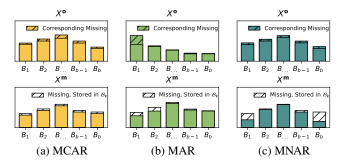


Figure 2. Illustration of probability mass adjustment for missing values in numeric or ordinal feature k. The red dot marks the observed value  $x_k$ ;  $B_0$  and  $B_b$  represent the first and last bins, respectively; and  $\mathcal{B}_k$  denotes the designated bin for missing values.



**Figure 3.** Visualization of mass bucket distributions under different missingness mechanisms. In  $X^0$ , the term *Corresponding Missing* refers to observed values associated with missing entries in  $X^m$ . In  $X^m$ , values are grouped and stored into buckets  $\mathcal{B}_k$ , shows the distribution of missingness.

uniformly at random,  $\mathcal{B}_k$  has limited effect due to the lack of informative structure. However, under MAR and MNAR,  $\mathcal{B}_k$  becomes highly informative. In MAR settings, missingness depends on observed values ( $X^{\mathbf{o}}$ ), which are often correlated with unobserved values ( $X^{\mathbf{m}}$ ). As shown in Figure 3 (b),  $\mathcal{B}_k$  implicitly captures this correlation structure by accumulating contextually similar missing entries. In MNAR cases (Figure 3 (c)), missingness depends directly on the missing values themselves. Since these unobserved entries tend to concentrate around specific value ranges or categories,  $\mathcal{B}_k$  acts as a proxy for these latent patterns. Thus, the inclusion of  $\mathcal{B}_k$  significantly enhances robustness under non-random missing mechanisms.

Maximizing Uncertainty (MaxU). The uncertainty-aware term is grounded in the principle of maximum entropy [18], which assigns the largest plausible dissimilarity when a value is missing. MaxU conservatively models the missing entry as potentially any value in the feature domain  $\mathcal{X}_k$ , leading to the worst-case dissimilarity:  $|R_k(x_k,?)| = \sup_{x' \in \mathcal{X}_k} |R_k(x_k,x')|$ . This strategy implicitly assumes a uniform distribution over all plausible values, avoiding arbitrary assumptions and improving robustness under adversarial or structured missingness. Compared to alternative schemes like Average Uncertainty (AvgU) and Minimum Uncertainty (MinU), MaxU aligns with the minimax principle [15], preserving discriminative power by maximizing entropy. While AvgU assumes a mean-case estimate and MinU favors optimistic matches, both are prone to failure under high sparsity or biased missing patterns. MaxU avoids such pitfalls, reducing imputation bias and maintaining kernel expressiveness across diverse missing mechanisms.

# 4.5 Limitation and Complexity Analysis

The original  $M_0$  similarity [4] computes dissimilarity via binbased histograms, resulting in a preprocessing time complexity of  $O(mnb + nb^2)$ , where m is the number of instances, n the number of features, and b the number of bins. This scaling becomes prohibitive for large or high-dimensional datasets. By contrast, our proposed HI-PMK avoids binning and instead uses precomputed probability masses based on the observed data distribution. The resulting similarity matrix can be computed in  $O(m^2 \cdot n)$  time and stored in  $O(m^2)$  space—matching the complexity of standard similarity measures such as Euclidean or cosine distance. This efficiency enables HI-PMK to scale to moderate-sized datasets while supporting mixed feature types and structured missingness. A detailed runtime analysis is provided in Section 5.3.

#### 5 Experimental Results

We evaluate HI-PMK on both clustering and classification tasks. While PMK was initially designed for clustering, our results show that HI-PMK generalizes well to classification, highlighting its robustness across downstream applications. Additional experiments are available in the Supplementary Material [47]. Code: github.com/echoid/Incomplete-Heter-Kernel

# 5.1 Experimental Setup

#### 5.1.1 Datasets.

We evaluate two types of datasets from the UCI Machine Learning Repository<sup>1</sup>, as summarized in Table 1. (i) Real-World Datasets with Synthetic Missingness: We introduce MCAR, MAR, and MNAR patterns at varying missing rates into 10 fully observed datasets to systematically evaluate robustness under controlled conditions. (ii) Real-World Incomplete Datasets: Six naturally incomplete datasets with unknown mechanisms (possibly a mixture of mechanisms), reflecting practical challenges in real-world settings.

Dataset Summary											
Dataset	N	Ord	Nom	Num	С	M.R.					
Complete Datasets											
Adult	48,842	1	7	6	4	-					
Australian	69,014	0	8	6	2	-					
Banknote	1,372	0	0	5	2	-					
Breast	2,869	4	5	0	2	-					
Car	1,728	6	0	0	4	-					
Heart	303	0	8	5	5	-					
Sonar	208	0	0	60	2 4 5 2 2 5	-					
Spam	4,601	0	0	57	2	-					
Student	649	11	16	2		-					
Wine	4,898	0	0	12	2	-					
Incomplete Datasets											
Hepat	155	0	13	6	2	5.67%					
Horse	368	13	ĺ	8	2	23.80%					
Kidney	400	2	10	12	2	10.54%					
Mammo	961	4	0	1	2	3.37%					
Pima	768	0	0	8	2	12.24%					
Wiscon	699	9	0	0	2	0.25%					

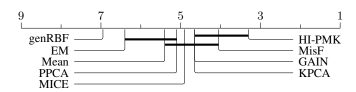
Table 1. N = instances, Ord/Nom/Num = ordinal/nominal/numerical, C = classes, M.R. = missing rate.

Synthetic Incomplete Data Generation. We introduced missingness into complete datasets under three standard mechanisms. MCAR removes values uniformly at random, independent of features or values. MAR introduces missingness based on observed features, following the strategy in [24]. MNAR depends on unobserved values: for numerical features, extreme values (e.g., high/low percentiles) are more likely to be missing; for ordinal features, boundary categories have higher missing rates; for nominal features, certain categories are assigned greater missing probabilities.

https://archive.ics.uci.edu/

Dataset	Hepat		Horse		Kidney		Mammo		Pima		Wiscon	
Metric	ACC	NMI	ACC	NMI	ACC	NMI	ACC	NMI	ACC	NMI	ACC	NMI
Mean	0.8129	0.0015	0.8398	0.0078	0.9775	0.0067	0.8148	0.0959	0.7735	0.0111	0.9628	0.7295
MICE	0.8129	0.0021	0.8506	0.0024	$\overline{0.9700}$	0.0074	0.8241	0.0959	0.7722	0.0910	0.9614	0.7427
EM	$\overline{0.8000}$	0.0025	0.8398	0.0079	0.9500	0.0045	0.8200	0.0911	0.7696	0.0211	0.9614	0.7387
MisF	0.8065	0.0014	0.8262	0.0078	0.9650	0.0023	0.8200	0.0959	0.7722	0.0169	0.9628	0.7335
GAIN	0.8274	0.0012	0.8501	0.0071	0.8525	0.0160	0.8106	0.0932	0.7462	0.0596	0.9642	0.7387
genRBF	0.7935	0.0772	0.6304	0.0166	0.6250	0.0132	0.5140	0.0002	0.6510	0.0052	0.5564	0.4505
KPCA	0.7935	0.0159	0.6850	0.0583	0.6250	0.0056	0.8127	0.0117	0.6510	0.0092	0.9500	0.5186
PPCA	0.8000	0.0015	0.8506	0.0540	0.9625	0.0081	0.8241	0.0959	0.7722	0.0909	0.9628	0.7427
Simp	-	0.0019	-	0.0001	-	0.0424	_	0.0951		0.0556		$\overline{0.0000}$
Gow	-	0.1968	-	0.1280	-	0.3899	-	0.2970	-	0.1069	-	0.6939
HI-PMK	0.8065	0.2001	0.8506	0.1066	0.9875	0.6663	0.8241	0.3271	0.7735	0.1374	0.9700	0.7585

**Table 2.** NMI scores for clustering tasks on incomplete datasets and Classification accuracy for incomplete datasets. The bold values indicate the best-performing models, while the underlined values represent the second-best performance for each dataset.



**Figure 4.** Critical Difference (CD) Diagram showing the ranking of models based on their F1 scores across all datasets. Smaller ranking value represent better performance.

# 5.1.2 Benchmark Methods for Comparison

Imputation-Based Methods. We include representative imputation techniques: Mean/Mode Imputer as a simple baseline, MICE [40] for its iterative inference, EM [10, 20] for likelihood-based estimation, and MisF [36], a tree-based approach tailored for mixed-type data. Although recent studies [11, 32, 46] question the efficacy of deep models on tabular data, we also evaluate GAIN [41], a generative method representative of deep learning-based imputers.

**Kernel Representation Methods.** We assess genRBF [34], which directly models similarity in the presence of missing data, and two classical kernel learning methods—KPCA [31, 48] and PPCA [13]—which rely on prior imputation (via MICE) and highlight the limitations of conventional kernels when applied to heterogeneous or incomplete datasets.

**Clustering Methods.** We include Simp and Gow [28], two similarity-based clustering methods designed for heterogeneous data. To enable clustering on incomplete datasets, we first apply MICE imputation to ensure complete inputs.

# 5.1.3 Evaluation Metrics

For clustering, we applied K-means with the number of clusters K set to the ground-truth number of classes. Performance was evaluated using Normalized Mutual Information (NMI) and Adjusted Rand Index (ARI), where ARI accounts for chance agreement. Each experiment was repeated five times, and results were averaged for stability.

For classification, we used Support Vector Machines (SVM) with RBF kernels. On complete datasets, we used standard RBF kernels; for incomplete binary-class datasets, RBF kernels were computed over similarity matrices. Methods like genRBF and HI-PMK, which rely on pairwise similarities, were evaluated using SVMs with precomputed kernels. We performed 5-fold cross-validation, tuning hyperparameters (e.g., C, kernel width) via nested inner 5-fold CV. Fi-

nal models were trained on the full training set and evaluated using F1 score and accuracy.

# 5.2 Analysis of Results

Incomplete Data with Unknown Missing Mechanism. Table 2 presents classification accuracy and clustering NMI on real-world incomplete datasets. As expected, Simp and Gow, being clustering-only methods, are not applicable to classification. HI-PMK consistently achieves the best or near-best performance, demonstrating its robustness under unknown and mixed missing mechanisms. Although these datasets contain relatively low missing rates, HI-PMK maintains strong results without relying on imputation or prior knowledge of the missingness pattern.

Complete Data with Synthetic Missing Mechanism. Figure 5 reports F1 scores across varying missing rates and mechanisms, with Table 3 showing detailed results at 20% missingness. HI-PMK consistently ranks among the top performers under all mechanisms. It is particularly effective on both numerical datasets and mixed-type datasets, confirming its adaptability. Unlike most baselines, its performance remains stable even as missingness increases. HI-PMK also generalizes well to high-dimensional datasets like *Sonar* and *Spam*. The Critical Difference (CD) diagram in Figure 4 ranks methods based on F1 scores, with HI-PMK achieving the top rank overall, followed by MisF and GAIN.

## 5.3 Scalability

We assess the runtime scalability of HI-PMK under varying sample sizes, feature dimensions, and missing rates using synthetic datasets. The evaluation settings are as follows:

- Sample Size: n=30 features, missing rate 30%;
- **Feature Dimension:** n=1000 features, missing rate 30%;
- Missing Rate: d=30 samples, n=1000 features.

Sample Size. HI-PMK demonstrates efficient scaling with increasing sample size and remains faster than deep models like GAIN for  $d \leq 2000$ , making it well-suited for small to medium-scale datasets.

**Feature Dimension.** As dimensionality increases, HI-PMK's runtime grows moderately due to kernel computations. It consistently outperforms kernel methods such as KPCA and PPCA, which exhibit steeper growth under high dimensions.

**Missing Rate.** HI-PMK maintains stable runtime across a wide range of missing rates. In contrast, deep generative models incur additional cost due to prolonged training. This stability highlights HI-PMK's robustness under high sparsity.

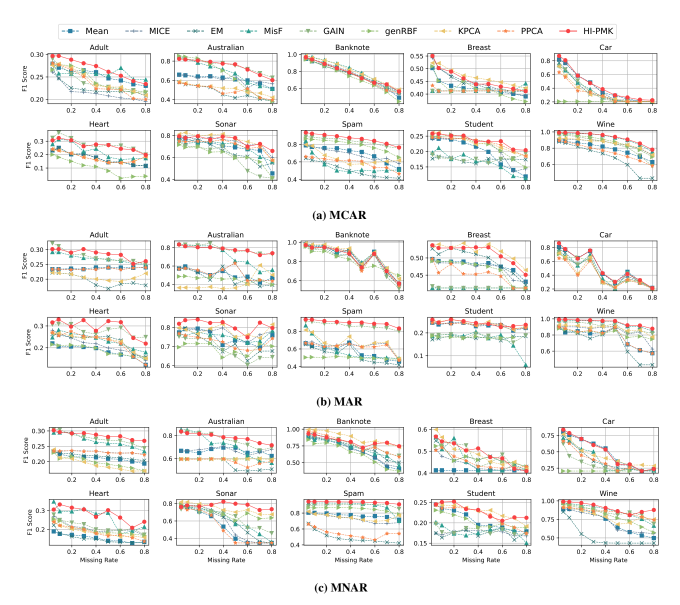


Figure 5. F1 scores for classification tasks under different missingness mechanisms (MCAR, MAR, MNAR), evaluated across varying missing rates. Each plot illustrates how classifier performance responds to increased data degradation, highlighting sensitivity to the underlying missing data pattern.

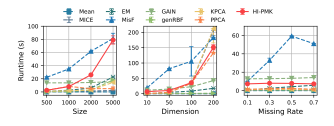


Figure 6. Scalability across sample size, dimension, and missing rate.

# 5.4 Ablation Study

We conduct two ablation studies to assess the contribution of HI-PMK components: (1) a module-level decomposition and (2) a comparison of uncertainty modeling strategies. Table 4 reports average F1 scores over complete datasets under MCAR, MAR, and MNAR

conditions (left), and average classification and clustering performance on real incomplete datasets (right).

Component-Level Analysis. We evaluate: PMK (baseline using complete data only), H-PMK (PMK with MICE Imputer), I-PMK-w/o MaxU (removes uncertainty-aware term), I-PMK-w/o  $\mathcal{B}_k$  (excludes frequency-based missing bucket adjustment), I-PMK (includes both MaxU and  $\mathcal{B}_k$ ), and HI-PMK (our model).

Uncertainty Strategy Comparison. We further compare MaxU with AvgU and MinU, which estimate average-case and optimistic dissimilarities, respectively. MaxU assumes worst-case uncertainty by assigning the largest plausible dissimilarity when values are missing. Results show that MaxU consistently yields superior performance, especially under MNAR, validating the benefit of conservative uncertainty modeling [18].

• MaxU leads to significant performance gains, particularly in in-

MCAR										
Model	Adult	Australian	Banknote	Breast	Car	Heart	Sonar	Spam	Student	Wine
Mean	0.2546	0.6127	0.7619	0.4278	0.4377	0.1774	0.6912	0.6978	0.1985	0.7962
MICE	0.2178	0.6123	0.7725	0.4508	0.4285	0.1774	0.7406	0.6933	0.2126	0.8563
EM	0.2269	0.4800	0.7709	0.4515	0.3736	0.1774	0.6628	0.5148	0.1742	0.6900
MisF	0.2584	0.6909	0.8109	0.4160	0.3741	0.2281	0.6977	0.5694	0.1660	0. <u>9234</u>
GAIN	0.2423	0.7447	0.7613	0.4135	0.3688	0.2608	0.6046	0.8351	0.1781	0.8463
genRBF	0.2443	0.6575	0.7345	0.4236	0.2059	0.1045	0.6202	0.7939	0.2195	0.8827
KPCA	0.2438	0.5161	0.8102	0.4307	0.3517	0.1817	<u>0.7519</u>	0.6500	0.2285	0.8610
PPCA	0.2384	0.4987	0.7702	0.4148	0.3339	0.1817	0.7142	0.5909	0.2281	0.7617
HI- PMK	0.2697	0.7500	<u>0.8107</u>	0.4605	0.4602	0.2694	0.7538	0.8673	0.2363	0.9329
MAR										
Mean	0.2371	0.5115	0.8116	0.4781	0.5181	0.1848	0.7507	0.5755	0.2329	0.7780
MICE	0.2351	0.5245	0.8161	0.4715	0.5096	0.2159	0.7739	0.6347	0.2374	0.8515
EM	0.1960	0.4840	0.8255	0.4943	0.4560	0.2310	0.7117	0.5374	0.1792	0.7179
MisF	0.2710	0.7191	0.8550	0.4127	0.4626	0.2355	0.7556	0.5948	0.1706	0.9470
GAIN	0.2791	0.7905	0.8106	0.4133	0.4458	0.2814	0.6892	0.8665	0.1840	0.9077
genRBF	0.2017	0.4543	0.8089	0.4667	0.4513	0.1889	0.6919	0.5002	0.1808	0.8235
KPCA	0.2084	0.4098	0.8107	0.5108	0.4264	0.2313	0.7819	0.6770	0.2402	0.8843
PPCA	0.2351	0.5236	<u>0.8360</u>	0.4526	0.4027	0.2270	0.7261	0.6229	0.2352	0.7857
HI- PMK	0.2852	<u>0.7841</u>	0.8390	0.5141	0.5250	0.2945	0.8129	0.8984	0.2429	0.9577
				M	INAR					
Model	Adult	Australian	Banknote	Breast	Car	Heart	Sonar	Spam	Student	Wine
Mean	0.2129	0.6663	0.6855	0.4127	0.5025	0.1555	0.5402	0.7712	0.2072	0.7237
MICE	0.2316	0.6704	0.7034	0.4413	0.4950	0.2098	0.5809	0.7310	0.1757	0.7268
EM	0.2184	0.5538	0.7506	0.4746	0.3871	0.1547	0.5987	0.4988	0.1743	0.5309
MisF	0.2694	0.7200	0.7222	0.4638	0.3901	0.2672	0.6216	0.8913	0.1793	0.8658
GAIN	0.2769	0.7678	0.7710	0.4849	0.3117	0.2178	0.6414	0.8979	0.1806	0.8543
genRBF	0.1990	0.5967	0.6390	0.4719	0.2059	0.2008	0.6973	0.8493	0.2050	0.8073
KPCA	0.1940	0.5967	0.8011	<u>0.4919</u>	0.4527	0.1882	0.7471	0.7516	0.2275	0.8034
PPCA	0.2311	0.5791	<u>0.7915</u>	0.4541	0.3885	0.1870	0.5289	0.5493	0.2181	0.8483
HI- PMK	0.2858	0.7851	0.8253	0.4925	0.5030	0.2825	0.7737	0.9347	0.2290	0.9111

**Table 3.** Classification results showing the F1 scores of HI-PMK and other baseline methods at a missing rate of 20% across multiple datasets and mechanisms (MCAR, MAR, and MNAR). The highest score for each dataset has been highlighted in bold, and the second-highest score is underlined.

complete data with structured missingness. Its conservative treatment of uncertainty helps avoid biased similarity estimates.

- B<sub>k</sub> enhances robustness for categorical features, especially under MAR and MNAR, where missingness correlates with specific values or categories.
- While these components offer marginal gains on complete data, they are critical for performance under real-world incomplete conditions
- HI-PMK achieves the best overall results, benefiting from the combined effects of hybrid modeling and uncertainty-aware similarity estimation.

These findings confirm the necessity of jointly modeling uncertainty and missing value frequency for achieving generalization across diverse missingness patterns. The effectiveness of HI-PMK stems from its principled design that integrates both data semantics and statistical uncertainty.

# 6 Conclusion

We introduced HI-PMK, a data-dependent kernel framework that directly models similarity under incomplete and heterogeneous settings

Variant	MCAR	MAR	MNAR	Cls.	Clus.
PMK	0.509	0.507	0.502	0.751	0.253
H-PMK	0.549	0.521	0.530	0.844	0.340
I-PMK-w/o MU	0.572	0.581	0.572	0.816	0.328
I-PMK-w/o $\mathcal{B}_k$	0.551	0.608	0.610	0.818	0.316
I-PMK	0.592	0.637	0.613	0.827	0.339
HI-PMK	0.630	0.651	0.647	0.869	0.362
HI-PMK-AvgU	0.622	0.616	0.603	0.860	0.359
HI-PMK-MinU	0.592	0.587	0.593	0.842	0.312
HI-PMK-MaxU	0.630	0.651	0.647	0.869	0.362

Table 4. Ablation study comparing components and uncertainty strategies.

without requiring imputation. By incorporating uncertainty-aware modeling and frequency-based correction, HI-PMK handles diverse missingness mechanisms and supports mixed-type features. Extensive experiments demonstrate consistent gains in classification and clustering tasks across 15+ benchmarks. Future work will focus on reducing quadratic complexity and extending the method to temporal or graph-structured domains, where missingness patterns exhibit richer dependencies.

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