Multi-Label Classification via Adaptive Resonance Theory-Based Clustering

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Abstract—This article proposes a multi-label classification algorithm capable of continual learning by applying an Adaptive Resonance Theory (ART)-based clustering algorithm and the Bayesian approach for label probability computation. The ART-based clustering algorithm adaptively and continually generates prototype nodes corresponding to given data, and the generated nodes are used as classifiers. The label probability computation independently counts the number of label appearances for each class and calculates the Bayesian probabilities. Thus, the label probability computation can cope with an increase in the number of labels. Experimental results with synthetic and real-world multi-label datasets show that the proposed algorithm has competitive classification performance to other well-known algorithms while realizing continual learning.

Index Terms—Multi-label classification, continual learning, clustering, adaptive resonance theory, correntropy

1 INTRODUCTION

Thanks to the recent advances of IoT technology, we can easily obtain a wide variety of data and utilize them to machine learning algorithms. Thus, the importance of continual learning is increasing for machine learning algorithms in order to efficiently utilize the data [1]. The requirement for continual learning is to handle both sequential learning and class-incremental learning without destroying the learned knowledge. In general, sequential learning is defined as a method that learns the data instance by instance, not in batches. Class-incremental learning is defined as a method that can deal with the situation where the number of classes (labels) increases during the learning process.

Since real-world phenomena and objects are complex and may have multiple semantics in nature, multi-label classification attracts a great deal of attention from machine learning and related fields such as web mining [2], rule mining [3], and information retrieval [4], [5]. In regard to multi-label classification algorithms, the sequential learning has realized by stream multi-label classification algorithms [6]. For those algorithms, however, data pre-processing such as normalization and standardization is often required. In addition, it is necessary for learning process to define the number of classes in advance. The class-incremental learning is theoretically feasible with the Bayesian approach for label probability computation in Multi-Label k-Nearest Neighbor (ML-kNN) [7]. The label learning process independently counts the number of label appearances for each class and calculates the Bayesian probabilities. Thus, the label probability computation can cope with an increase in the number of labels. However, ML-kNN cannot cope with the sequential learning because k-NN requires the entire data before the learning process. Multi-label Learning with Emerging New Labels (MuENL) [8] has realized continual learning by constructing two classifiers for classifying instances and for detecting new labels. However, MuENL cannot perform continual learning in the non-stationary environment, i.e., the situation where new data distributions are sequentially provided.

In the case of single-label classification algorithms, several types of clustering-based classifiers capable of continual learning have been proposed [9], [10], [11], [12], [13], [14]. In particular, classifiers designed by an Adaptive Resonance Theory (ART)-based clustering algorithm have shown comparable classification performance to typical classification algorithms such as SVM and k-NN. The main feature of the
above ART-based clustering algorithms is the use of the Correntropy-Induced Metric (CIM) [15] to a similarity threshold, which makes the self-organizing process fast and stable [11], [12], [13], [14].

This paper is an extended version of our conference paper [16]. In the conference paper, we have introduced the basic algorithm of Multi-Label CIM-based ART (MLCA). In this paper, we provide a comprehensive analysis of MLCA in terms of the classification performance for numerical and categorical datasets, the continual learning ability, and the computational complexity by comparing with recently proposed multi-label classification algorithms. Furthermore, we proposed two variants of MLCA by modifying the calculation method of the CIM to improve the classification performance of MLCA.

The contributions of this paper are summarized as follows:

i) A multi-label classification algorithm, called MLCA, is proposed by integrating a CIM-based ART and the Bayesian approach for label probability computation. MLCA computes the prior probability and likelihood by using nodes (representative points of training data) which have accumulated label counts while ML-$k$NN computes them by directly referencing to training instances.

ii) A new CIM-based ART is proposed by introducing an efficient node generation process and a bandwidth adaptation method for a kernel function in CIM. As a result, MLCA does not require any data pre-processing method such as normalization or scaling.

iii) Empirical studies show that MLCA and its variants have competitive classification performance to recent multi-label classification algorithms.

iv) The continual learning ability of MLCA is analyzed from multiple perspectives, and its usefulness and superiority are clarified.

The paper is organized as follows. Section 2 presents literature review for clustering algorithms and multi-label classification algorithms. Section 3 describes details of the proposed algorithm and modifications of the calculation method of the CIM. Section 4 presents extensive simulation experiments to evaluate the continual learning ability and classification performance of MLCA. Section 5 concludes this paper.

2 Literature Review

2.1 Clustering Algorithm

Cluster analysis is one of the widely applied approaches to extract hidden relation from data. Typical types of clustering algorithms are the Gaussian mixture model [17], $k$-means [18], and Self-Organizing Map (SOM) [19]. Although the above algorithms are quite simple and highly adaptable, the number of classes and network architectures are specified in advance. Growing Neural Gas (GNG) [20] and Self-Organizing Incremental Neural Network (SOINN) [9] are well-known growing self-organizing clustering algorithms that can overcome the drawbacks of the typical types of clustering algorithms. GNG and SOINN can adaptively generate topological networks corresponding to given data. However, since these algorithms permanently insert new nodes into their networks for memorizing new knowledge, they have a potential to forget learned knowledge (i.e., catastrophic forgetting). This trade-off is called the plasticity-stability dilemma [21]. A variant of GNG, called Grow When Required (GWR) [22] can avoid the plasticity-stability dilemma by adding nodes whenever the state of the current network does not sufficiently match the instance. One problem of GWR is that as the number of nodes in the network increases, the cost of calculating a threshold for each node increases, and thus the learning efficiency decreases.

A successful approach to avoid the plasticity-stability dilemma is the ART-based algorithms [23]. Because the ART-based algorithms realize sequential and class-incremental learning without the catastrophic forgetting, a number of the ART-based algorithms and their improvements are proposed in both supervised learning [24], [25], [26] and unsupervised learning [27], [28], [29], [30]. In the ART-based algorithms, a criterion of a new category (node) generation, i.e., a similarity measurement between a node and an instance, has a great impact on the classification/clustering performance. Previous studies have shown that algorithms with the CIM [15] as a similarity measurement are capable of faster and more stable learning than other self-organizing clustering algorithms [11], [12], [13], [14].

2.2 Multi-Label Classification

The multi-label classification algorithms are categorized into two approaches, namely, a problem transformation approach and an algorithm adaptation approach [31]. The problem transformation approach transforms a multi-label classification problem into multiple single-label classification problems. The problem transformation approach is further divided into two methods, namely, the Binary Relevance (BR) [32] and the Label Powerset (LP) [33]. The BR transforms a multi-label classification problem into multiple binary classification problems by decomposing multi-labels into multiple single labels. The LP transforms a multi-label classification problem into a multi-class classification problem by merging multi-labels into a single label. Various single-label classification algorithms have been used in the problem transformation approach thanks to its simplicity and applicability.

The algorithm adaptation approach extends existing single-label classification algorithms for handling multi-label classification problems. Various types of algorithm adaptation approach have been introduced based on $k$-NN [7], [34], [35], decision tree [36], regression [37], Support Vector Machine (SVM) [38], feed-forward neural networks [39], [40], and an ensemble approach [41]. In order to achieve high classification performance, recent studies consider label distributions and their correlations in a label learning process [42], [43], [44]. These algorithms, however, cannot cope with the situation where new label information is sequentially provided. ML-$k$NN [7] is a well-known algorithm adaptation method that integrates $k$-NN and the Bayesian approach for label probability computation. ML-$k$NN counts the number of relevant labels of neighbors for each instance in training data. Based on the counts of relevant labels, the likelihood and posterior probability of each
label are computed by the Bayesian approach. Here, the computation of the Bayesian probability is individually performed in each label. Therefore, in theory, the number of labels to be learned can be increased/decreased during the label probability computation. One disadvantage of ML-kNN is that it cannot efficiently cope with the situation where new training instances are sequentially provided. Multi-Label Self-Adjusting k Nearest Neighbors (MLSA-kNN) [45] is capable of handling a stream multi-label classification by employing a self-adjusting window to detect a concept drift and to adaptively control a parameter $k$ in kNN. MuENL [8] generates two classifiers for classifying instances and for detecting new labels. By using the two classifiers, MuENL realizes sequential learning and class-incremental learning, simultaneously. Although MuENL is capable of continual learning, MuENL requires a batch learning process in the initial stage of learning for constructing two classifiers for instances and labels. Moreover, MuENL cannot deal with the situation where new data distributions are provided in the non-stationary environment.

Several studies have realized multi-label classification by using a clustering algorithm. A typical type of clustering-based multi-label classification algorithm utilizes SOM [46], [47]. Although the learning process of SOM is performed in an unsupervised learning manner, the convergence of the SOM network is significantly slow and unstable. The online semi-supervised GNG for multi-label classification utilizes GNG as a base classifier [48]. Unlike SOM-based algorithms, this method adaptively generates a topological network corresponding to the given instances. However, as mentioned in the literature review for clustering in the previous subsection, GNG-based algorithms do not satisfy the requirements of continual learning. Multi-label Classification via Incremental Clustering (MCIC) [49] has realized continual learning by applying a continual clustering to a learning process. MCIC extracts and accumulates information from data by nodes through the continual clustering process that takes into account the arrival time of data. In addition, the continual clustering process also constructs and updates a distribution of labels in each node for a label estimation process. Some studies have employed Fuzzy ARTMAP [50] to a base classifier of the multi-label classification [51], [52], [53], [54]. ARTMAP is composed of two ART architecture to realize an explicit supervised learning process. Although Fuzzy ARTMAP has various advantages, there is a well-known problem, i.e., high sensitivity to statistical overlapping between the generated categories [55]. This sensitivity problem results in category proliferation (i.e., disordered generation of categories), which leads to a high computational cost and deterioration in the classification performance. The recent ART-based algorithms in [51], [52], [53], [54] potentially have this problem.

### 3 Proposed Algorithm

In this section, first the theoretical background of the CIM is briefly described. Next, the proposed algorithm, namely, MLCA, is explained in detail. Then, two variants of MLCA are introduced by modifying the calculation method of the CIM. Table 1 summarizes the main notations used in this paper.

| Notation | Description |
|----------|-------------|
| $X = (x_1, x_2, \ldots, x_n, \ldots)$ | Training instances |
| $x_n = (x_{n,1}, x_{n,2}, \ldots, x_{n,d})$ | $d$-dimensional training instance (the $n$th instance) |
| $L = (l_1, l_2, \ldots, l_n, \ldots)$ | Relevant labels for $X$ |
| $l_i = (l_{i,1}, l_{i,2}, \ldots, l_{i,N})$ | Relevant label for $x_i$ |
| $N$ | Dimension of the relevant label set for $x_n$ |
| $Y = (y_1, y_2, \ldots, y_K)$ | Prototype nodes |
| $y_i = (y_{i,1}, y_{i,2}, \ldots, y_{i,d})$ | $d$-dimensional prototype node (the $k$th node) |
| $S = (\sigma_1, \sigma_2, \ldots, \sigma_K)$ | Bandwidths for a kernel function |
| $\kappa$ | Kernel function with a bandwidth $\sigma$ |
| $k^+$ | Relevant label for $x_n$ |
| $k^-$ | Prototype node of $x_n$ |
| $V_{x_i}$, $V_{y_i}$ | Predefined similarity threshold |
| $V$ | The number of instances that have accumulated by the node $y_{k_1}$ |
| $\alpha_{y_i}$ | Neighbor node of $y_{k_1}$ |
| $\beta_{y_i}$ | The number of labels that have accumulated by the node $y_{k_1}$ |
| $V_{y_i}$ | The predefined number of neighbor nodes for $y_{k_1}$ |
| $N_y$ | The number of instances in each node for a label estimation process |
| $\lambda$ | Interval for adapting $\sigma$ |
| $H_i$ | Event that an instance has the $i$th label |
| $H_0$ | Event that an instance does not have the $i$th label |
| $P(E|H)$ | Likelihood for a label probability computation |
| $P(H|E)$ | Prior probability for a label probability computation |
| $P(H|E)$ | Posterior probability for a label probability computation |
| $c$ | Label counter |
| $x^* = (x^*_1, x^*_2, \ldots, x^*_d)$ | $d$-dimensional testing instance |
| $l^*$ | Predicted label vector of the test instance $x^*$ |
3.1 Correntropy and Correntropy-Induced Metric

Correntropy [15] provides a generalized similarity measure between two arbitrary instances $x = (x_1, x_2, \ldots, x_d)$ and $y = (y_1, y_2, \ldots, y_d)$ as follows:

$$C(x, y) = E[\kappa_\sigma(x - y)],$$  \hspace{1cm} (1)

where $E[\cdot]$ is the expectation operation, and $\kappa_\sigma(\cdot)$ denotes a positive definite kernel with a kernel bandwidth $\sigma$. The correntropy can be estimated as follows:

$$\hat{C}(x, y) = \frac{1}{d} \sum_{i=1}^{d} \kappa_\sigma(x_i - y_i).$$  \hspace{1cm} (2)

In this paper, we use the following Gaussian kernel in the correntropy:

$$\kappa_\sigma(x_i - y_i) = \exp\left(-\frac{(x_i - y_i)^2}{2\sigma^2}\right).$$  \hspace{1cm} (3)

A nonlinear metric called CIM is derived from the correntropy [15]. The CIM quantifies the similarity between two instances as follows:

$$CIM(x, y, \sigma) = \left[\kappa_\sigma(0) - \hat{C}(x, y)\right]^2,$$  \hspace{1cm} (4)

where $\kappa_\sigma(0) = 1$ from (3). Here, thanks to the Gaussian kernel without a coefficient $\frac{1}{\sqrt{2\pi}\sigma}$ as defined in (3), a range of the CIM is limited to $[0, 1]$.

3.2 Learning Procedure

We use the following notations: Training instances are $X = (x_1, x_2, \ldots, x_n)$, where $x_n = (x_{n,1}, x_{n,2}, \ldots, x_{n,d})$ is a $d$-dimensional feature vector. Relevant label sets for $X$ are $L = (l_1, l_2, \ldots, l_n)$, where $l_i = (l_{i,1}, l_{i,2}, \ldots, l_{i,n_K})$ is a binary vector used to show a set of relevant labels (classes) in the learned training instances. Note that MLCA is capable of continual learning, the algorithm can accept any number of training instances and labels. A set of $\lambda$ prototype nodes in MLCA is defined by

$$(x_{\lambda}, y_{\lambda}, k_{\lambda}) \in \mathbb{R}^d \times \mathbb{R}^{d+n_K} \times \mathbb{N},$$

where $x_{\lambda} = (x_{\lambda,1}, \ldots, x_{\lambda,d})$ has the same dimension as $x_n$. Furthermore, each node $y_{\lambda}$ has an individual bandwidth $\sigma$ for the CIM, i.e., $S = (\sigma_1, \sigma_2, \ldots, \sigma_K)$.

The learning procedure of MLCA is divided into five parts: 1) initialization process for a bandwidth of a kernel function in the CIM, 2) winner node selection, 3) vigilance test, 4) node learning, and 5) label probability computation. Each of them is explained in the following subsections.

3.2.1 Initialization Process for a Bandwidth of a Kernel Function in the CIM

Similarity measurement between an instance and a node has a large impact on the performance of clustering algorithms. MLCA uses the CIM as a similarity measure. As defined in (4), the state of the CIM is controlled by a bandwidth $\sigma$ of a kernel function which is a data-dependent parameter.

In general, the bandwidth of a kernel function can be estimated from $\lambda$ instances belonging to a certain distribution [56], which is defined as follows:

$$\Sigma = U(F_0)\Gamma \lambda^{-\frac{1}{d+1}},$$  \hspace{1cm} (5)

$$U(F_0) = \left(\frac{\pi^{d/2}2^{d+1}(d!)^2 R(F)^d}{\int_{\mathbb{R}^d}(2\pi)^d (d-1)!(d-1)!}ight) \frac{1}{\Gamma^{d+1}},$$  \hspace{1cm} (6)

where $\Gamma$ denotes a rescale operator ($d$-dimensional vector) which is defined by a standard deviation of the $d$ attributes among $\lambda$ instances. $\nu$ is the order of a kernel. $R(F)$ is a roughness function. $\kappa_\sigma(F)$ is the moment of a kernel. In this paper, we utilize the Gaussian kernel for the CIM. Therefore, $\nu = 2, R(F) = (2\sqrt{\pi})^{-1}$, and $\kappa_\sigma(F) = 1$ are derived. The details of the derivation of (5) and (6) can be found in [56].

In MLCA, the initial state of $\sigma$ in the CIM is defined by training instances. When a new node $y_{\lambda+1}$ is generated from $x_n$, a bandwidth $\sigma_{\lambda+1}$ is estimated from the past $\lambda$ instances, i.e., $(x_{\lambda-1}, x_{\lambda-2}, \ldots, x_{\lambda-\lambda})$, by using (5) and (6) with $\nu = 2, R(F) = (2\sqrt{\pi})^{-1}$, and $\kappa_\sigma(F) = 1$, as follows:

$$\Sigma = \left(4 \frac{1}{2+d}\right)^{\frac{1}{d+1}} \Gamma \lambda^{-\frac{1}{d+1}},$$  \hspace{1cm} (7)

where $\Gamma$ denotes a rescale operator ($d$-dimensional vector) which is defined by a standard deviation of the $d$ attributes among the past $\lambda$ training instances of MLCA. Here, $\Sigma$ contains the bandwidth of each attribute.

In this paper, the median of $\Sigma$ is selected as a representative bandwidth for the new node $y_{\lambda+1}$, i.e.,

$$\sigma_{\lambda+1} = \text{median}(\Sigma).$$  \hspace{1cm} (8)

3.2.2 Winner Node Selection

Once an instance $x_n$ is presented to MLCA, two nodes which have a similar state to the instance $x_n$ are selected, namely, winner nodes $y_{\lambda_1}$ and $y_{\lambda_2}$. The winner nodes are determined based on the state of the CIM as follows:

$$k_1 = \arg\min_{k \notin K} \text{CIM}(x_n, y_k, \text{mean}(S)),  \hspace{1cm} (9)

$$k_2 = \arg\min_{k \notin K} \text{CIM}(x_n, y_k, \text{mean}(S)),  \hspace{1cm} (10)

$$\text{where } k_1 \text{ and } k_2 \text{ denote indexes of the 1st and 2nd winner nodes i.e., } y_{\lambda_1} \text{ and } y_{\lambda_2}, \text{ respectively. } S \text{ is a bandwidth for a kernel function of the CIM in each node.}$

Note that when there is no node in MLCA, the $(\lambda+1)$th instance becomes the initial node (i.e., $y_{1} = x_{\lambda+1}$). In the case, the bandwidth of $y_{\lambda}$ is estimated from the 1st to $\lambda$th instances in a set of training instances $X$ by (5)-(8), and the next instance is given without vigilance test until the 1st and 2nd winner nodes can be defined.

3.2.3 Vigilance Test

Similarities between an instance $x_n$ and the 1st and 2nd winner nodes are defined as follows:

$$V_{k_1} = \text{CIM}(x_n, y_{k_1}, \text{mean}(S)).$$  \hspace{1cm} (11)

$$V_{k_2} = \text{CIM}(x_n, y_{k_2}, \text{mean}(S)).$$  \hspace{1cm} (12)
The vigilance test classifies the relationship between an instance and a node into three cases by using a predefined similarity threshold \( V \).

- **Case I**
  
  A similarity between an instance \( x_n \) and the 1st winner node \( y_{k_1} \) is larger (i.e., less similar) than the similarity threshold \( V \), namely:
  \[
  V_{k_1} > V.
  \]

  If (13) is satisfied, \( V_{k_2} > V \) is also satisfied since \( V_{k_2} > V_{k_1} > V \). Thus, a new node is defined as \( y_{K+1} = x_n \), and the bandwidth \( \sigma_{K+1} \) is defined by (8).

  Moreover, two counters \( \alpha \) and \( \beta \) are initialized. One counter \( \alpha \) is the number of instances that have been accumulated by a node, which is initialized as \( \alpha_{K+1} = 1 \). The other counter \( \beta \) is the number of labels that have been accumulated by a node, which is initialized as \( \beta_{K+1} = 1 \).

- **Case II**
  
  A similarity between an instance \( x_n \) and the 1st winner node \( y_{k_1} \) is smaller (i.e., more similar) than the similarity threshold \( V \), and a similarity between the instance \( x_n \) and the 2nd winner node \( y_{k_2} \) is larger (i.e., less similar) than the similarity threshold \( V \), namely:
  \[
  V_{k_1} \leq V, \text{ and } V_{k_2} > V.
  \]

  If (14) is satisfied, node learning is performed. In addition, two counters \( \alpha \) and \( \beta \) for the winner node \( y_{k_1} \) are updated as follows:
  \[
  \alpha_{k_1} \leftarrow \alpha_{k_1} + 1,
  \]
  \[
  \beta_{k_1} \leftarrow \beta_{k_1} + \beta_{k_1}.
  \]

- **Case III**
  
  Similarities between an instance \( x_n \) and the 1st and 2nd winner nodes are both smaller (i.e., more similar) than the similarity threshold \( V \), namely:
  \[
  V_{k_1} \leq V, \text{ and } V_{k_2} \leq V.
  \]

  If (17) is satisfied, node learning is performed. Moreover, two counters \( \alpha \) and \( \beta \) for the node \( y_{k_1} \) are updated by (15) and (16), respectively.

### 3.2.4 Node Learning

Different node learning is performed based on the results of the vigilance test.

If Case II, the state of the 1st winner node \( y_{k_1} \) is updated as follows:

\[
y_{k_1} \leftarrow y_{k_1} + \frac{1}{\alpha_{k_1}} (x_n - y_{k_1}).
\]

When updating the node, the amount of change is divided by \( \alpha_{k_1} \), so the larger \( \alpha_{k_1} \) becomes, the smaller the node position changes. This is based on the idea that the information around a node, where instances are frequently given, is important and should be held by the node.

If Case III, the state of the 1st winner node \( y_{k_1} \) is updated by (18). In addition, all neighbor nodes of \( y_{k_1} \) (i.e., \( y_{k_1}^{\text{neighbor}} \)) are also updated as follows:

\[
y_{k_1}^{\text{neighbor}} \leftarrow y_{k_1}^{\text{neighbor}} + \frac{1}{N_{y_{k_1}}^{\text{neighbor}}} (y_{k_1} - y_{k_1}^{\text{neighbor}}),
\]

where \( N_{y_{k_1}}^{\text{neighbor}} \) is the predefined number of neighbor nodes for \( y_{k_1} \), \( \alpha_{k_1}^{\text{neighbor}} \) denotes the number of instances that have been accumulated by the node \( y_{k_1}^{\text{neighbor}} \).

Equation (19) has the same concept as (18), but it should be less affected by the instance than \( y_{k_1} \) because it is the neighbor node of \( y_{k_1} \). Thus, \( N_{y_{k_1}} \) is added as a coefficient.

### 3.2.5 Label Probability Computation

Similar to ML-kNN, MLCA employs the Bayesian approach for label probability computation. The prior probability and likelihood are updated if the condition for Case I or Case II of the vigilance test is satisfied. Note that ML-kNN computes the prior probability and likelihood by using training instances repeatedly, while MLCA computes the prior probability and likelihood by using nodes which have accumulated label counts. As a result, MLCA realizes continual learning by computing the prior probability and likelihood whenever an instance is given.

To update the prior probability and likelihood, an instance \( x_n \), with a set of labels \( l_i \), the 1st winner node \( y_{k_1} \) and its \( N_{y_{k_1}} \) neighbor nodes are considered. Note that \( N_{y_{k_1}} \) denotes the number of neighbor nodes and is a predefined parameter in MLCA. Here, we consider the situation where \((n-1)\) instances have been given to MLCA. The likelihood \( P(E|H) \) is computed as follows:

\[
P(E_{n,i}|H^\phi_{n,i}) = \frac{(s + c_i^\phi)}{s \times (N_{y_{k_1}} + 1) + \sum_{j=0}^{N_{y_{k_1}}} c_j^\phi}, \quad (i \in N_i, \phi \in \{+,-\}),
\]

where \( H_{n,i}^+ \) is the event that an instance has the \( i \)-th label (i.e., \( l_i = 1 \)), and \( H_{n,i}^- \) is the event that an instance does not have the \( i \)-th label (i.e., \( l_i = 0 \)). Here, \( H = H_{n,i}^+ \cup H_{n,i}^- \) is satisfied. \( E_i \) is the event that the frequency of the label \( l_i \) among the \( N_{y_{k_1}} \) neighbor nodes of \( y_{k_1} \). \( N_i \) is a size of a label set. \( s \) is a smoothing parameter. In this paper, \( s \) is set to be 1 which yields the Laplace smoothing. A label counter \( e_i \) is defined as follows:

\[
\begin{align*}
  c_{i,j=0}^{\phi} & \leftarrow c_{i,j=0}^{\phi} + 1, \quad \text{if } l_{n,i} = 1, \\
  c_{i,j=1}^{\phi} & \leftarrow c_{i,j=1}^{\phi} + 1, \quad \text{otherwise}.
\end{align*}
\]

Here, \( g_i \) is the \( i \)-th attribute of an \( N_i \)-dimensional counting vector \( g = (g_1, \ldots, g_{N_i}) \), which is defined as follows:

\[
g_i = \sum_{j \in N_{y_{k_1}}} \beta_{ij}, \quad (i \in N_i).
\]

In order to make the maximum of \( g_i \) the same as the number of neighbor nodes \( N_{y_{k_1}} \), the following operation is performed:

\[
g_i \leftarrow \text{round} \left[ N_{y_{k_1}} \cdot \frac{g_i}{\max(g)} \right].
\]
Algorithm 1. Learning Procedure of MLCA

Input:
- a training instance: \( \mathbf{z}_i = (x_{i1}, x_{i2}, \ldots, x_{id}) \),
- a label set: \( l_i = (l_{i1}, l_{i2}, \ldots) \),
- prototype nodes: \( Y = \{y_1, y_2, \ldots, y_K\} (K \in \mathbb{Z}^+) \),
- bandwidths of \( Y \): \( S = \{\sigma_1, \sigma_2, \ldots, \sigma_K\} \),
- the number of instances that have been accumulated by nodes \( Y \): \( \alpha = \{\alpha_1, \alpha_2, \ldots, \alpha_K\} \),
- the number of labels that have been accumulated by nodes \( Y \): \( \beta = \{\beta_1, \beta_2, \ldots, \beta_K\} \),
- a label counter: \( c \),
- the number of neighbor nodes: \( N_y \),
- an interval for adapting \( \sigma \): \( \lambda \),
- and a similarity threshold: \( V \).

Output:
- updated prototype nodes: \( Y \),
- updated bandwidths of \( Y \): \( S \),
- the updated number of instances that have been accumulated by the nodes \( Y \): \( \alpha \),
- the updated number of labels that have been accumulated by the nodes \( Y \): \( \beta \),
- the updated label counter: \( c \),
- a prior probability: \( P(H) \),
- and a likelihood: \( P(E | H) \).

1 function (LearningMLCA(\( \mathbf{z}_i, l_i, Y, S, \alpha, \beta, c, N_y, \lambda, V \))
2 Input an instance \( \mathbf{z}_i \).
3 Input a label set \( l_i \).
4 if The index \( i \) is multiple of \( \lambda \) then
5 Compute a bandwidth for the CIM by (8).
6 if \( K < 1 \) do
7 Generate a new node as \( y_{K+1} = \mathbf{z}_i \).
8 Assign a bandwidth \( \sigma_{K+1} \) which is defined by (8).
9 Update \( \alpha_{K+1} \) and \( \beta_{K+1} \) by (15) and (16), respectively.
10 else
11 Compute the CIM by (4).
12 Search indexes of winner nodes \( k_1 \) and \( k_2 \) by (9) and (10), respectively.
13 if CIM(\( \mathbf{z}_i, y_{k_1}, \text{mean}(S) \)) > \( \lambda \) then
14 Generate a new node as \( y_{K+1} = \mathbf{z}_i \).
15 Compute a bandwidth \( \sigma_{K+1} \) which is defined by (8).
16 Update \( \alpha_{K+1} \) and \( \beta_{K+1} \) by (15) and (16), respectively.
17 Update similarities between an instance \( \mathbf{z}_i \) and nodes \( Y \) by the CIM.
18 Update a likelihood \( P(E | H) \) by (20).
19 else
20 Update a state of \( y_{k_1} \) by (18).
21 Update \( \alpha_{k_1} \) and \( \beta_{k_1} \) by (15) and (16), respectively.
22 if CIM(\( \mathbf{z}_i, y_{k_2}, \text{mean}(S) \)) \leq \( \lambda \) then
23 Update the state of \( N_y \) neighbor nodes of \( y_{k_2} \) by (19).
24 Update a likelihood \( P(E | H) \) by (20).
25 Compute a prior probability by (24) and (25).
26 return \( Y, S, \alpha, \beta, c, P(E | H) \), and \( P(H) \).

The prior probability \( P(H) \) is computed as follows:

\[
P(H^+_i) = \frac{\sum_{k=1}^{N} \beta_k}{(s \times 2 + n)}, \quad (i \in N_i), \quad (24)
\]

\[
P(H^-_i) = 1 - P(H^+_i), \quad (i \in N_i), \quad (25)
\]

where \( n \) denotes the number of instances that have been given. Here, \( P(H) = 1 \) is satisfied.

The learning procedure of MLCA is summarized in Algorithm 1.

### 3.3 Label Prediction Procedure

We use the following notations: A testing instance is \( \mathbf{z}^* = (x_{1}^*, x_{2}^*, \ldots, x_{d}^*) \). A set of prototype nodes in MLCA after the learning procedure is \( Y = \{y_1, y_2, \ldots, y_K\} \). In addition, each node \( y_k \) has an individual bandwidth \( \sigma_k \) for the CIM, i.e., \( S = \{\sigma_1, \sigma_2, \ldots, \sigma_K\} \). Moreover, the likelihood \( P(E | H) \), the prior probability \( P(H) \), and the label counter \( \beta \) are utilized for the Bayesian approach.

First of all, the winner node \( y_{k_1} \) for a test instance \( \mathbf{z}^* \) and its \( N_y \) neighbor nodes are determined. In the same manner as the learning procedure, a membership counting vector \( g \) for the test instance \( \mathbf{z}^* \) is computed by (23). The posterior probability \( P(H^+_i | E | g) \) for the test instance \( \mathbf{z}^* \) is defined by the Bayes rule as follows:

\[
P(H^+_i | E | g) = \frac{P(E | H^+_i) P(H^+_i)}{\sum_{\phi \in \{+,-\}} P(E | H^+_i) P(H^+_i)}, \quad (i \in N_i). \quad (26)
\]

A predicted label vector \( \Gamma^* \) of the test instance \( \mathbf{z}^* \) is determined by a simple thresholding method as follows:

\[
\Gamma^*_i = \begin{cases} 
1 & \text{if } P(H^+_i | E | g) > 0.5, \quad (i \in N_i), \\
0 & \text{otherwise}
\end{cases} \quad (27)
\]

Since the label prediction process is a binary classification, it is reasonable to specify the threshold to 0.5. ML-kNN also uses the same value.

The prediction procedure of MLCA is summarized in Algorithm 2.

### 3.4 Attribute Processing for the CIM

As shown in (2) and (3), the CIM in MLCA uses a common bandwidth \( \sigma \) even if the range of attribute values is different, and the average value as the similarity between an instance \( \mathbf{z}_i \) and a node \( y_k \). Therefore, if the range of attribute values is significantly different, a specific attribute may have a large impact on the value of the CIM when the common bandwidth \( \sigma \) is not appropriate for that attribute.

In this section, we propose two approaches in order to mitigate the above-mentioned effects: 1) one approach calculates the CIM by using an each individual attribute separately, and the average value as the similarity measurement, and 2) the other approach applies a clustering algorithm to attribute values, then attributes with similar value ranges are grouped. The CIM is calculated by using an each attribute group, and the average CIM value is used for similarity measurement.

#### 3.4.1 Individual-Based Approach

In this approach, the CIM is calculated by using an each individual attribute separately, and the average CIM value is used for similarity measurement. The similarity between an instance \( \mathbf{z}_i \) and a node \( y_k \) is defined by the CIM as follows:

\[
\text{CIM}^i(\mathbf{z}_i, y_k, \sigma_k) = \frac{1}{d} \sum_{i=1}^{d} \left[ \text{CIM}_i(x_{ni}, y_{ki}) \right]^2, \quad (28)
\]

\[
\text{CIM}_i(x_{ni}, y_{ki}) = \kappa_{\sigma_k}(x_{ni} - y_{ki}), \quad (29)
\]

where \( \sigma_k \) is the bandwidth for attribute \( x_k \).
where $\sigma_i = (\sigma_1, \sigma_2, \ldots, \sigma_d)$ is a bandwidth of a node $y_k$. A bandwidth for the $i$th attribute is defined as follows:

$$\sigma_i = \left( \frac{4}{2 + \Delta} \right)^{\frac{1}{\beta}} \Gamma_i \lambda^{-\frac{1}{\beta}},$$

(30)

where $\Gamma_i$ denotes a rescale operator which is defined by a standard deviation of $i$th attribute values among the $\lambda$ instances.

In this paper, MLCA with the individual-based approach is called MLCA-Individual (MLCA-I).

**Algorithm 2. Prediction Procedure of MLCA**

**Input:**
- a testing instance: $x^* = (x^*_1, x^*_2, \ldots, x^*_d)$,
- prototype nodes: $Y = (y_1, y_2, \ldots, y_K)$ ($K \in \mathbb{Z}^+$),
- the bandwidths of $Y$: $\mathcal{S} = (\sigma_1, \sigma_2, \ldots, \sigma_K)$,
- the number of labels that have been accumulated by nodes $Y$: $\beta = (\beta_1, \beta_2, \ldots, \beta_K)$,
- the number of neighbor nodes: $N_y$,
- and a similarity threshold: $V$.

**Output:**
- a predicted label vector: $l^*$.

1. function (PredictMLCA($x^*$, $Y$, $\mathcal{S}$, $\beta$, $N_y$, $V$))
2. Input an instance $x^*$.
3. Compute similarities between an instance $x^*$ and nodes $Y$ by the CIM.
4. Compute a membership counting vector $g$ by (23).
5. Compute a posterior probability $P(V|E)$ by (26).
6. Determine a predicted label vector $l^*$ by (27).
7. return $l^*$.

### 3.4.2 Clustering-Based Approach

In this approach, for every $\lambda$ instances, the clustering algorithm presented in Section 3.2 is applied to the attribute values. Each attribute value of $\lambda$ instances is regarded as a one-dimensional vector and used as the input to the clustering algorithm. As a result, attributes with similar value ranges are grouped together, i.e., an instance $x_n = (x_1, x_2, \ldots, x_d)$ is transformed into $x^C_n = (u_{n,1}, u_{n,2}, \ldots, u_{n,j})$ ($J \leq d$) by the clustering algorithm, where $u_{j}$ represents the $j$th attribute group. The dimensionality of each attribute group is represented by $d = (d_1, d_2, \ldots, d_J)$ where $d_j$ is the dimensionality of the $j$th attribute group.

In this approach, the similarity between an instance $x^C_n$ and a node $y_k$ is defined by the CIM as follows:

$$\text{CIM}(x^C_n, y_k, \sigma^C_n) = \frac{1}{J} \sum_{j=1}^{J} \left[ \sigma_j (0) - \hat{C}_C(u_j, v_j) \right]^2,$$

(31)

$$\hat{C}_C(u_j, v_j) = \frac{1}{d_j} \sum_{i=1}^{d_j} \kappa_{\sigma_j} (u_i - v_i),$$

(32)

where $y^C_k = (v_1, v_2, \ldots, v_j)$ is a node $y_k$, but its attributes are grouped by referencing to the attribute indexes of $x^C_n$. A bandwidth $\sigma_j$ is defined as follows:

$$\sigma_j = \left( \frac{4}{2 + \Delta} \right)^{\frac{1}{\beta}} \Gamma_j \lambda^{-\frac{1}{\beta}},$$

(33)

where $\Gamma_j$ denotes a rescale operator which is defined by the standard deviation of the $i$th attribute value in the $j$th attribute group among the $\lambda$ instances.

In this paper, MLCA with the clustering-based approach is called MLCA-Clustering (MLCA-C).

The differences in attribute processing among the general approach, the individual-based approach (MLCA-I), and the clustering-based approach (MLCA-C) are depicted in Fig. 1.

The codes of MLCA, MLCA-I, and MLCA-C are available on GitHub.¹

### 4 Simulation Experiments

In this section, the ability of MLCA is evaluated from various perspectives. First, the continual learning ability of MLCA is analyzed with a two-dimensional synthetic multi-label dataset under the stationary and non-stationary environments. Next, the classification performance of MLCA is compared with other algorithms by using real-world multi-label datasets. Third, we evaluate the effect of a multi-epoch learning process to MLCA. Finally, we analyze the computational complexity of each algorithm.

#### 4.1 Evaluation Metrics

We use the six metrics to evaluate the classification performance of multi-label classification algorithms [31].

- **Exact Match**
  This is possibly the most strict performance metric. This metric measures whether a predicted set of labels for an instance is exactly equal to the true labels. The higher, the better.

- **$F_1$-score**
  This is a harmonic mean between the Precision and the Recall, namely, a weighted measure of how many

1. https://github.com/Masuyama-lab/MLCA
true labels are predicted and how many predicted labels are truly relevant. The higher, the better.

- **Macro-averaged AUC**
  This is the area under the receiver operating characteristic curve. The Macro-averaged AUC is the arithmetic mean of the AUC for each label. This metric gives a better sense of the performance across all labels. The higher, the better.

- **Hamming Loss**
  This metric computes the symmetric difference between the predicted and true labels, and divided by the total number of labels in a dataset. The lower, the better.

- **Ranking Loss**
  This metric computes how many times a relevant label (a member of the true labels) appears ranked lower than a non-relevant label, namely, the average proportion of label pairs that are incorrectly ordered for an instance. The lower, the better.

- **Coverage**
  This metric is defined as the distance to cover all possible labels assigned to an instance, namely, how many top-scored predicted labels are included without missing any true labels. The lower, the better. In this paper, we scaled the Coverage by the number of labels \(N_l - 1\) thus the range of the Coverage is \([0, 1]\).

### 4.2 Continual Learning Ability

In theory, ART-based clustering is capable of learning new knowledge and preserving the learned knowledge without catastrophic forgetting by setting a fixed similarity threshold (i.e., a vigilance parameter in ART). Thus, MLCA can continually learn and preserve knowledge by adaptively generating nodes in response to changes of data distributions. Moreover, since MLCA has a fixed similarity threshold \(V\) and there is no node deletion process, MLCA does not inherently cause catastrophic forgetting.

In this section, we verify the continual learning ability of MLCA by using a two-dimensional synthetic multi-label dataset in the stationary and non-stationary environments.

For comparison, we apply MCIC [49] which is capable of continual learning through a clustering process. Similar to MLCA, MCIC extracts and accumulates information from data by nodes. Therefore, MCIC is a competitive algorithm for MLCA. Note that, as mentioned in Section 2.2, MuENL [8] can perform sequential learning and class-incremental learning, i.e., continual learning. However, MuENL cannot deal with the situation where new data distributions are provided in the non-stationary environment, and thus MuENL cannot cope with the experimental conditions in this section.

Fig. 2 shows the two-dimensional synthetic multi-label dataset. The dataset consists of three distributions where each has 10,000 instances that follow a uniform distribution. In addition, as shown in Fig. 2b, seven types of label sets are defined. In this experiment, the instances are given to each algorithm in three different conditions: (1) all the instances are given at the same time, (2) the three distributions are given in sequential order, and (3) the seven distributions are given in sequential order. The condition (1) is the stationary environment while the conditions (2) and (3) are the non-stationary environment. Here, the label information of each distribution is not changed. It should be noted that the given instances are presented as a sequence of three distributions in Fig. 3 whereas they are divided into seven distributions in Fig. 4. As a result, Fig. 4 is an easier problem than Fig. 3 because there are no overlap regions.

In the case that the three distributions are given in sequential order (Fig. 3), the instances in the overlapping regions are designed to contain label information of the already given distribution (e.g., \(I_i = (1, 1)\)) for generating a situation where the number of labels increases in a pseudo manner. Thus, as shown in Table 2, seven types of label sets are defined after the three distributions are given. As the number of labels increases, the length of the label set also changes incrementally, e.g., \(I_A = (1) \rightarrow I_A = (1, 0) \rightarrow I_A = (1, 0, 0)\). The similar label transition is occurred in the case of Fig. 4.

In order to analyze the continual learning capability, we evaluate the classification performance after each distribution is given. After learning the training instances of each distribution, the instances belonging to the learned distribution is used as test data. Namely, 1) after learning the distribution
#1, the classification performance is evaluated by using the test data of distribution #1. Next, II) after learning the distribution #2, the classification performance is evaluated by using the test data of the distributions #1 and #2. Then, III) after learning the distribution #3, the classification performance is evaluated by using the test data of the distributions #1 and #3. We continue this procedure until all the distributions are given. We repeat the experiment 20 times with a different random seed to obtain consistent results. The parameters of MLCA are specified as follows: \( N_f = 10 \), \( \lambda = 50 \), and \( V = 0.10 \). The parameters of MCIC are specified as follows: \( \delta = 0.1 \), \( K = 3 \), \( \lambda = 0.25 \), \( \beta = 2 \), \( \epsilon = \{0.011, 0.007, 0.005\} \), and a processing speed is 10,000. Under the above parameter settings, MLCA and MCIC generate the similar number of nodes.

Figs. 5, 6, 7, 8, and 9 show the visualization of generated nodes and their label information in each algorithm. The results in Figs. 5, 6, 7, 8, and 9 are a trial which showed the highest Exact Match in each algorithm among 20 trials. The color of a node indicates a label set that the node predicts. Once a testing instance is given, the nearest node predicts a label set for the testing instance corresponding to the color shown in the legend of Figs. 5, 6, 7, 8, and 9. The quantitative results corresponding to Figs. 5, 6, 7, 8, and 9 are provided in Table 1 in the supplementary file, which can be found on the Computer Society Digital Library at http://doi.ieeecomputersociety.org/10.1109/TPAMI.2022.3230414.

Comparing Figs. 5a and 5b, MLCA can represent the seven distributions very well while MCIC fails to represent overlapped regions. Let us focus on Figs. 6 and 7, MLCA and MCIC can properly preserve the information of the distribution #1. On the other hand, after learning the distributions #2 and #3, MLCA properly represents overlapped regions, but MCIC fails to do so. A similar tendency can be seen in Figs. 8 and 9. These results suggest that MLCA is capable of continuous learning in various environments while MCIC cannot accumulate and preserve information when distributions are adjacent or overlapping.

From the results in this section, it can be seen that MLCA adaptively generates nodes and incrementally learns label information from the given instances while maintaining the extracted knowledge.

In this section, since the label information of each distribution is not changed, the learning ability has been verified only for the knowledge that is represented by node positions/distributions. Regarding the label information (the prior probabilities and likelihood) of nodes in MLCA, it is sequentially updated based on the frequency of a label appearance. Therefore, although the location or distribution of nodes do not change, the meaning of the knowledge may change in the case where the label distribution is changed. This is known as concept drift.

From another point of view, the label forgetting nature of MLCA may lead to superior classification performance for stream data with concept drift than other algorithms. This is an interesting future research topic.

### 4.3 Quantitative Analysis

This section presents a comparison on the classification performance of MLCA, MLCA-I, and MLCA-C with that of MCIC [49], MuENL [8], mIODM [43], GLOCAL [42], MLSA-kNN [45], and ML-kNN [7] by utilizing real-world multi-label datasets.

The source code of MCIC,2 MuENL,3 mIODM,4 GLOCAL,5 MLSA-kNN,6 and ML-kNN7 are provided by authors.
4.3.1 Datasets
We use 16 real-world multi-label datasets that six numerical and six categorical datasets from the Mulan repository [57] and two numerical and two categorical datasets from the Extreme Classification repository [58]. Table 3 shows the statistics of the datasets. During our experiments, the training instances in each dataset are presented to each algorithm only once. In regard to pre-processing for datasets, mlODM and GLOCAL need [0, 1] scaling to maintain high classification performance, while other algorithms do not need the scaling. Therefore, we prepare the [0, 1] scaled data for mlODM and GLOCAL. For other algorithms, we use the raw data with no pre-processing.

4.3.2 Parameter Specifications
MLCA, MLCA-I, MLCA-C, and all the compared algorithms have parameters which have an impact on the classification performance. This section presents parameter specifications of each algorithm in detail.

We use grid search to specify parameter values of each algorithm. Before grid search, we separate a dataset into training instances and test instances. The training instances are 90% of the dataset, and the testing instances are the remaining 10%. The testing instances are used only for the performance evaluation of the designed classifier (i.e., they are not used for parameter specifications). During grid search, we train an algorithm with the training instances, and test the algorithm by using the training instances again. Thus, we obtain a parameter setting which shows the highest classification performance to the training instances. Once the best parameter setting, which is optimal to the training instances, is specified, the classification performance of the algorithm with the best parameter setting is evaluated by using the testing instances. Since grid search for specifying the parameters does not use the testing instances, the generalization ability of each algorithm can be properly evaluated in Section 4.3.4.

For specifying the best parameter setting, we calculate the Exact Match in each parameter specification for the training instances in each dataset listed in Table 3 except for large-scale datasets due to a time-consuming training process. Regarding the parameters for the large-scale datasets, we apply the same specification with Langlog dataset because it has the largest number of labels among small-scale datasets.

We repeat the evaluation 20 times (i.e., 2×10-fold Cross Validation (CV)) with training instances selected by different random seeds. Although six evaluation metrics were introduced in Section 4.1, the Exact Match is used for specifying the parameters since it is the most strict evaluation metric.

### Table 3: Statistics of Multi-Label Datasets

| Dataset       | Number of Instances | Number of Attributes | Number of Labels |
|---------------|---------------------|----------------------|------------------|
| Small-scale   |                     |                      |                  |
| Flags         | 194                 | 10                   | 9                |
| Emotions      | 593                 | 72                   | 0                |
| Birds         | 645                 | 258                  | 2                |
| Image         | 2,000               | 294                  | 0                |
| Scene         | 2,407               | 294                  | 0                |
| Yeast         | 2,417               | 103                  | 0                |
| VirusGO       | 207                 | 0                    | 749              |
| GpositiveGO   | 519                 | 0                    | 912              |
| Genbase       | 662                 | 0                    | 1,186            |
| Medical       | 978                 | 0                    | 1,449            |
| PlantGo       | 978                 | 0                    | 3,091            |
| Langlog       | 1,460               | 0                    | 1,004            |
| Large-scale   |                     |                      |                  |
| EURLex-4K     | 19,348              | 5,000                | 0                |
| Mediamill     | 43,907              | 120                  | 3,993            |
| Bitex         | 7,395               | 0                    | 1,836            |
| Delicious     | 16,105              | 0                    | 500              |

Fig. 8. Visualization of nodes in MLCA in the case the seven distributions are given in sequential order.

Fig. 9. Visualization of nodes in MCIC in the case the seven distributions are given in sequential order.
Table 2 in the supplementary file, available online summarizes the parameters of all the algorithms. In the following, the settings and results of grid search are explained in detail for each of MLCA, its variants and the compared algorithms.

- **MLCA and its Variants**
  
  MLCA, MLCA-I, and MLCA-C have three parameters, i.e., the number of neighbor nodes $N_\nu$, an interval $\lambda$ for adapting $\sigma$ in the CIM, and a similarity threshold $V$. Among those parameters, the similarity threshold $V$ has a large impact on the classification performance. Therefore, we perform grid search for $V$ in increments of 0.05 over the range of $V = [0.05, 0.95]$ while fixing $N_\nu = 10$ and $\lambda = 50$. The parameters $N_\nu$ and $\lambda$ are the same specification as in [16]. In addition, the case of $V = 0.01$ is also considered. This is because that MLCA with a smaller $V$ will tend to generate more nodes which yields higher classification performance than that with a larger $V$.

The detailed results of grid search for the similarity threshold $V$ in MLCA and its variants are shown in Fig. 1 of the supplementary file, available online.

In this paper, the parameters of MLCA and its variants are not specified for each dataset, but the same settings are applied to all datasets. Note that the parameters of the compared algorithms are specified for each dataset in order to achieve the maximum classification performance. To specify the appropriate parameters for all datasets, we adopt the Friedman test and Nemenyi post-hoc analysis [59] to conduct statistical comparisons among the different parameter specifications ($V = [0.01, 0.45]$) in Fig. 1 of the supplementary file, available online) by using results on the Exact Match of all datasets. The Friedman test is used to test the null hypothesis that all algorithms perform equally. If the null hypothesis is rejected, the Nemenyi post-hoc analysis is then conducted. The Nemenyi post-hoc analysis is utilized for all pairwise comparisons based on the ranks of results on the Exact Match of all datasets. The difference in the performance of two algorithms is treated as statistically significant if the p-value defined by the Nemenyi post-hoc analysis is smaller than the significance level. Here, the null hypothesis is rejected at the significance level of 0.05 both in the Friedman test and the Nemenyi post-hoc analysis.

Fig. 10 shows critical difference diagrams for different parameter specifications of each algorithm. A better specification has lower average ranks, i.e., on the right side of a critical distance diagram. In theory, the parameter specifications within a critical distance (i.e., a red line) do not have a statistically significance difference [59]. From the results in Fig. 10, parameter values to be used in comparisons of classification performance by testing instances are specified as follows: $V = 0.01$ and 0.30 for MLCA, $V = 0.01$ and 0.15 for MLCA-I, and $V = 0.01$ and 0.25 for MLCA-C. As mentioned earlier, MLCA and its variants utilize the above $V$ values for all the datasets in the classification experiments.

- **Compared Algorithms**

  When grid search was used in the original paper where each algorithm was proposed, we use the same grid search with the same range of parameter values for each algorithm. Otherwise, we choose one or two parameters which have the largest effects on the classification performance. The details are as follows:

  - **MCIC:** The range of the decay controlled parameter $\lambda$ is the same range in the original paper. Although the boundary of a cluster $\theta$ is fixed in the original paper, it has a large effect to the clustering performance. The range of $\theta$ is theoretically defined as $0 < \theta < 0.5$.
  
  - **MuENL:** In the original paper, grid search is performed to $\lambda_1, \lambda_2 \in \{0.001, 0.01, 0.1, 1\}$. In this paper, we set a wider range that includes the above one.
  
  - **mIODM:** The parameters and their grid ranges are the same as in the original paper.
  
  - **GLOCAL:** The parameters and their grid ranges are the same as in the original paper.
  
  - **MLSA-kNN:** In the original paper, the authors mentioned that a window size $kAd_j$ and the maximum window size $m_{\text{max}}$ need to be set appropriately values in advance depending on datasets. In the original paper, these parameters are fixed as $kAd_j = 100$ and $m_{\text{max}} = 1,000$. In this paper, grid search was performed in the ranges around these values.
  
  - **ML-kNN:** In the original paper, grid search was performed in the range of $k = \{8, 9, 10, 11, 12\}$. However, there was no significant difference in the classification performance. In this paper, we set wider range for finding an appropriate value.

  Table 4 shows the parameters that indicate the highest Exact Match for the training instances in each dataset as determined by grid search. N/A indicates that an algorithm can not build a predictive model within 5 days under the available computational resources. In this paper, we assign the worst evaluation value to each metric if an algorithm can not build a predictive model. The rest of parameters in each algorithm and the range of grid search are summarized in Table 2 of the supplementary file, available online.

Since MuENL has 10 parameters, we only consider three parameters because the rest of the parameters are related to class-incremental learning that is not performed in this section. Regarding mIODM and GLOCAL, there is a large difference in the classification performance depending on whether a dataset is pre-processed or not. In this paper, therefore, we use the [0, 1] scaled data and raw data without pre-processing in the learning process of mIODM and
and parameter specifications also. In Fig. 4, we can observe that MCIC has a clearly lower rank than the others. Moreover, MLCA and its variants show critical difference diagrams for the supplementary file, available online.  

Fig. 11 shows a critical difference diagram based on the classification performance for all the datasets. A better specification has a lower average rank, i.e., on the right side of the critical distance diagram. In theory, any parameter specifications within a critical distance (i.e., a red line) do not have a statistically significant difference [59]. Fig. 11 shows that MLCA (V = 0.01), MLCA-I (V = 0.01), and MLCA-C (V = 0.01) are superior to the compared algorithms with a lower average rank. Let us focus on MLCA and its variants with large V specifications (i.e., V = 0.15, 0.25, 0.30). In Fig. 11, there is no statistically significant difference among MLCA-C (V = 0.25), MLCA-I (V = 0.15), and MLCA (V = 0.30) whereas MLCA-C (V = 0.25) has a clearly lower rank than the others. Moreover, MLCA and its variants with large V specifications also do not show a statistically significant difference to the two batch learning algorithms mLODM and GLOCAL. In contrast, MLCA and its variants with large V specifications show a statistically significant difference to MCIC, MLSA-kNN, and MuENL which are capable of handling streaming data. The above observations suggest that MLCA and its variants have information compression ability while maintaining superior classification performance than other algorithms.

In order to discuss the features of each algorithm in detail, Figs. 12 and 13 show critical difference diagrams for each algorithm by using a critical difference diagram defined by the Nemenyi post-hoc analysis. The detailed results of six evaluation metrics for each dataset are summarized in Tables 3-6 on the supplementary file, available online.

### 4.3.4 Experimental Results

We compare the classification performance of each algorithm by using a critical difference diagram defined by the Nemenyi post-hoc analysis. The detailed results of six evaluation metrics for each dataset are summarized in Tables 3-6 on the supplementary file, available online.

| Dataset    | MCIC | MuENL | mLODM (Scaling) | mLODM (Raw) | GLOCAL (Scaling) | GLOCAL (Raw) | MLSA-kNN | ML-kNN |
|------------|------|-------|-----------------|-------------|-----------------|-------------|---------|-------|
|             | θ    | λ     | λ₃              | λ₄          | θ               | μ           | λ₃      | λ₄    |
| Flags      | 0.100 | 0.00  | 1.0             | 1.0         | 0.90            | 0.30        | 0.00    | 1.0   |
| Emotions   | 0.100 | 0.00  | 1.0E-6          | 1.0E-6      | 0.70            | 0.10        | 1.0E-4  | 1.0   |
| Birds      | 0.100 | 0.00  | 1.0             | 1.0         | 0.70            | 0.40        | 1.0E-1  | 1.0   |
| Image      | 0.100 | 0.00  | 1.0E-6          | 1.0E-6      | 0.70            | 0.40        | 1.0E-5  | 1.0E-4|
| Scene      | 0.100 | 0.00  | 1.0             | 1.0         | 0.90            | 0.10        | 0.0     | 0.0   |
| Yeast      | 0.100 | 0.00  | 1.0             | 1.0         | 0.90            | 0.30        | 1.0     | 1.0E-1|
| VirusGO    | 0.100 | 0.00  | 1.0E-6          | 1.0E-6      | 0.70            | 0.10        | 0.0     | 0.0   |
| GpositiveG | 0.495 | 0.45  | 1.0             | 1.0         | 0.90            | 0.10        | 0.0     | 0.0   |
| Genbase    | 0.200 | 0.00  | 1.0             | 1.0         | 0.90            | 0.10        | 0.0     | 0.0   |
| Medical    | 0.100 | 0.20  | 1.0             | 1.0         | 0.10            | 0.10        | 1.0E-4  | 1.0E-4|
| PlantGO    | 0.100 | 0.00  | 1.0E-6          | 1.0E-6      | 0.20            | 0.10        | 1.0E-3  | 1.0E-4|
| Langlog    | 0.100 | 0.10  | 1.0             | 1.0         | N/A             | N/A         | 1.0E-1  | 1.0E-5|
| EURlex-4K  | 0.100 | 0.10  | 1.0             | 1.0         | N/A             | N/A         | 1.0     | 1.0E-5|
| Mediamill  | 0.100 | 0.10  | 1.0             | 1.0         | N/A             | N/A         | 1.0E-1  | 1.0E-5|
| Bittex     | 0.100 | 0.10  | 1.0             | 1.0         | N/A             | N/A         | 1.0E-1  | 1.0E-5|
| Delicious  | 0.100 | 0.10  | 1.0             | 1.0         | N/A             | N/A         | 1.0E-1  | 1.0E-5|

N/A indicates that an algorithm could not build a predictive model within 5 days under the available computational resources.
numerical datasets and categorical datasets, respectively. In the case of numerical datasets (Fig. 12), the classification performance of MLCA and its variants ($V = 0.01$) are superior to compared algorithms with a statistically significant difference except for ML-$k$NN. Similarly, MLCA and its variants with large $V$ specifications (i.e., $V = 0.15$, $0.25$, $0.30$) show a statistically significant difference to MCIC, MLSA-$k$NN, and MuENL. For the numerical datasets, MLCA-I ($V = 0.01$) shows the lowest (i.e., best) rank. MLCA-I ($V = 0.15$) also shows a good classification performance, which has a statistically significant difference to the two batch learning algorithms GLOCAL (Scaling, Raw) and mLODM (Raw). Thus, we can see that MLCA-I is suitable for the numerical data sets. On the categorical databases (Fig. 13), ML-$k$NN and GLOCAL perform very well in contrast to the case of numerical datasets. However, these algorithms do not have a statistically significant difference to MLCA and its variants. Moreover, these algorithms have an obvious drawback compared to MLCA and its variants because they require all the training instances in advance. With respect to MLCA-C, there is no statistically significant difference among different $V$ specifications. On the other hand, the performance of MLCA and MLCA-I is greatly affected by the specification of a similarity threshold $V$, i.e., a statistically significant difference is exist among different specifications of $V$. Therefore, we can conclude that MLCA-C has stable classification performance on the categorical databases.

Table 5 shows the average number of generated nodes after learning the training instances. Let us focus on MLCA and MLCA-I, these algorithms generate only a very small number of nodes in the case of nominal datasets, especially when a similarity threshold $V$ is large. On the other hand, MLCA-C can maintain the sufficient number of nodes for classification even in the case of nominal datasets. Thus, MLCA-C is considered to be a strong algorithm for nominal datasets. This property can be seen in Figs. 1g-11 of the supplementary file, available online.

The experimental results in Table 5 show that MLCA and its variants have superior classification performance with several advantages over the compared algorithms. In mLODM, GLOCAL, MLSA-$k$NN, and ML-$k$NN, raw training instances are directly used for classification tasks. As a result, local distribution biases and outliers in training instances have a large effects on their classification performance. In MLCA and its variants, prototype nodes are used for classification tasks instead of directly using raw training instances. As a result, negative effects of local distribution biases and outliers are decreased. Moreover, their negative effects are decreased in prior probability learning and likelihood computation for label estimation in MLCA and its variants by using accumulated label counts at each node.

From the viewpoint of a clustering algorithm, an ART-based clustering algorithm shows fast and stable clustering performance. Furthermore, the CIM, which is used in MLCA and its variants, can calculate the similarity between instances more precisely than the euclidean Distance-based similarity measure, especially for high-dimensional data. As a result, MLCA and its variants achieve better classification performance than MuENL and MCIC, which utilize prototype nodes generated by clustering for classification tasks.

From the results in Figs. 11, 12, and 13, and Table 5, the characteristics of MLCA and its variants can be analyzed as follows:

- **MLCA**
  This algorithm can be the first-choice algorithm because it shows stable and high classification performance for both numerical and nominal datasets. In other words, it has an advantage if the attribute type of the dataset is unknown. Furthermore, in the case of a numerical dataset, MLCA shows high classification performance regardless of a specification of the similarity threshold $V$. This means that MLCA can maintain high classification performance and high information compression performance simultaneously.

- **MLCA-I**
  This algorithm shows the outstanding classification performance on the numerical datasets. On the other hand, the classification performance on the nominal datasets is low in comparison with the results on to numerical datasets. Therefore, MLCA-I has an advantage if the dataset contains only numerical attributes.

- **MLCA-C**
  This algorithm shows stable and high classification performance for both numerical and nominal datasets although it is not as good as MLCA. It is notable that MLCA-C stably has small rank values for both nominal and numerical datasets even when a specification of a similarity threshold $V$ is large. Therefore, MLCA-C can achieve high classification performance and high information compression for both numerical and nominal datasets.

Table 6 summarizes the characteristics of MLCA and its variants based on the above analysis.
4.4 Effects of a Multi-Epoch Learning Process

MLCA, MLCA-I, and MLCA-C utilize generated nodes as classifiers. This means that the clustering performance on the training instances has a huge impact on the classification performance. The nodes of MLCA, MLCA-I, and MLCA-C are adaptively and continually generated/updated by the given instances. Therefore, it is possible to improve the clustering performance by learning the training instances in multiple epochs, and consequently to improve the classification performance of MLCA, MLCA-I, and MLCA-C. This feature is one of the advantages of MLCA, MLCA-I, and MLCA-C against the other compared algorithms.

We only apply the datasets listed in Table 3 except for the large-scale datasets due to a time-consuming training process. Fig. 14 shows results of the Exact Match of MLCA, MLCA-I, and MLCA-C with the learning of the training instances for 1 to 10 epochs. The conditions of this experiment are the same as in Section 4.3.

The following observation is obtained: As the number of epochs increases, the value of the Exact Match increases or remains roughly the same in most cases except for the Birds dataset. In particular, it is effective for categorical data. Therefore, we regard that the multi-epoch learning process is generally beneficial for MLCA, MLCA-I, and MLCA-C.

4.5 Computational Complexity

This section presents the computational complexity of MLCA, MLCA-I, and MLCA-C. Specifically, MLCA and its variants are analyzed in detail.

Regarding MLCA, the computational complexity of each process is as follows: For computing a bandwidth of a kernel function in the CIM is $O(n^2d)$ (line 5 in Alg. 1), for computing the CIM is $O(ndK)$ (line 11 in Alg. 1), for sorting the result of the CIM is $O(K\log K)$ (line 12 in Alg. 1), and for computing the label probability is $O(N_lN_y)$ (line 24 in Alg. 1). Here, $n$ is the number of training instances, $\lambda$ is an interval for adapting $\sigma$, $K$ is the number of nodes, $N_l$ is a size of a label set, and $N_y$ is the predefined number of neighbor nodes for $y_k$. Thus, the total computational complexity of MLCA is $O(n^2d + ndK + K\log K + N_lN_y)$. In general, $\lambda \ll n$, $K \ll n$, and $\lambda < K$. As a result, the total computational complexity of MLCA is $O(ndK + N_lN_y)$.

In terms of MLCA-I and MLCA-C, the difference of a training process is only in CIM and CIM$^C$ which is defined in (28) and (31), respectively. Since CIM$^C$ considers an individual attribute of a training instance separately, it takes $O(ndK)$. CIM$^C$ applies a clustering approach to attributes of a training instance every $\lambda$ instances. Thus, $O(\frac{n}{\lambda}(d + ndK + K\log K))$ is additionally required. In general, $\lambda \ll n$, $K \ll n$, and $\lambda < K$. As a result, the computational complexity of MLCA-I and MLCA-C are $O(nd^2K + N_lN_y)$ and $O(n(ndK + N_lN_y))$, respectively.

Table 7 summarizes the computational complexity of MLCA, MLCA-I, MLCA-C, and compared algorithms. Here, variables in the computational complexity of compared algorithms are as follows:

- **MCIC**: $T$ is a time period defined as $T = \frac{1}{\lambda}\log\left(\frac{\beta_L}{\beta_M-1}\right) + 1$. Here, $n$ is the number of instances, $\lambda$ and $\beta_L$ are the parameters of MCIC. $K_M$ and $K_L$ are the number of mature and immature clusters of MCIC, respectively.
- **MuENL**: $N_l$ is the size of a label set, $n$ is the number of instances, and $d$ is the dimension of instances. Note that the complexity of MuENL in Table 7 shows a pairwise label ranking classifier, not including a label incremental learning process.
- **mlODM**: $n$ is the number of instances, $N_l$ is the dimension of a current relevant label, and $I$ is the number of iterations of an optimization process.
- **GLOCAL**: $n$ is the number of instances, $n_{lm}$ is the number of instances of a partitioned training instances, $k$ is a rank of a label matrix which satisfies $k < N_l$. 

### TABLE 6

| Algorithm | Similarity Threshold $V$ | Classification Performance | Nominal |
|-----------|--------------------------|----------------------------|---------|
| MLCA      | Small                     | Very High                  | Very High |
|           | Large                     | Medium                     | Low     |
| MLCA-I    | Small                     | Very High                  | Medium  |
|           | Large                     | Medium                     | Low     |
| MLCA-C    | Small                     | High                       | Medium  |
|           | Large                     | Medium                     | Medium  |
MLSA-\(k\)-NN: \(d\) is the dimension of a training instance, \(N_i\) is a size of a label set, \(m_{\text{max}}\) is the maximum size of the window, and \(m_{\text{min}}\) is the minimum size of the window.

ML-\(k\)-NN: \(n\) is the number of training instances, \(d\) is the dimension of a training instance, \(k\) is the number of nearest neighbors of \(k\)-NN, and \(N_i\) is a size of a label set. Here, \(O(n^2d)\) is for \(k\)-NN computation, and \(O(nkN_i)\) is for the label probability computation.

With respect to the computational complexity of each algorithm, MLSA-\(k\)-NN and MCIC shows their superior computational efficiency than the other algorithms. MLCA, MLCA-I, MLCA-C, and ML-\(k\)-NN have moderate computational efficiency. These algorithms do not dramatically increase the computational complexity even when the number of instances and labels in a dataset is large. In contrast, MuENL, miODM, and GLOCAL have a polynomial complexity in terms of the number of instances or the size of a label set. Therefore, we consider that these algorithms are time-consuming and could not generate a predictive model in a valid time, especially for the large-scale datasets with a large number of training instances and labels.

The training and testing time on a CPU are summarized in Tables 7 and 8 of the supplementary file, available online.

### 5 CONCLUDING REMARKS

This paper proposed a multi-label classification algorithm capable of continual learning by extending our preliminary research [16], namely MLCA. In addition, two variants of MLCA were proposed by modifying the calculation method of the CIM, namely MLCA-I and MLCA-C. The proposed algorithms consist of two components: The CIM-based ART and the Bayesian approach for label probability computation. Because both components can deal with a situation where new training instances and corresponding labels are sequentially provided, the proposed algorithms can realize continual learning. The results of extensive experiments from qualitative and quantitative perspectives showed that MLCA has competitive classification performance to other well-known algorithms while maintaining the continual learning ability. Furthermore, the results also showed that the performance of MLCA can be enhanced by modifying the calculation method of the CIM.

The ability to adapt to concept drift [62] and to handle mixed numerical and categorical data [63] are significant factors for clustering-based algorithms capable of continual learning. A future research topic is to examine the performance of the proposed algorithms under concept drift and to improve them. It is also an important future research topic to modify the proposed algorithms for mixed datasets with both numerical and categorical attributes.

### TABLE 7

| Algorithm | Computational Complexity | Ref. |
|-----------|--------------------------|------|
| MLCA      | \(O(ndK + N_iN_p)\)     | —    |
| MLCA-I    | \(O(nd^2K + N_iN_p)\)   | —    |
| MLCA-C    | \(O(n(dK + N_iN_p))\)   | —    |
| MCIC      | \(O(n + \frac{K_p + K_s}{2})\) | —    |
| MuENL     | \(O(ndN_p)\)            | [8]  |
| miODM     | \(O(nN_p^2I)\)          | [43] |
| GLOCAL    | \(O(n^2 + n_m^2 + kn)\) | [60] |
| MLSA-\(k\)-NN | \(O\left(m_{\text{max}}\left(d + N_i + \log_2\frac{m_{\text{max}}}{m_{\text{min}}}\right)\right)\) | [45] |
| ML-\(k\)-NN | \(O(n(d + kN_i))\)      | [61] |

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