Model-Based Clustering of High-Dimensional Data: Variable Selection versus Facet Determination[☆]

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Abstract

Variable selection is an important problem for cluster analysis of high-dimensional data. It is also a difficult one. The difficulty originates not only from the lack of class information but also the fact that high-dimensional data are often multifaceted and can be meaningfully clustered in multiple ways. In such a case the effort to find one subset of attributes that presumably gives the "best" clustering may be misguided. It makes more sense to identify various facets of a data set (each being based on a subset of attributes), cluster the data along each one, and present the results to the domain experts for appraisal and selection. In this paper, we propose a generalization of the Gaussian mixture models and demonstrate its ability to automatically identify natural facets of data and cluster data along each of those facets simultaneously. We present empirical results to show that facet determination usually leads to better clustering results than variable selection.

Keywords: Model-based clustering, Facet determination, Variable selection, Latent tree models, Gaussian mixture models

1. Introduction

Variable selection is an important issue for cluster analysis of high-dimensional data. The cluster structure of interest to domain experts can often be best described using a subset of attributes. The inclusion of other attributes can degrade clustering performance and complicate cluster interpretation. Recently there is a growing interest in the issue [2, 3, 4]. This paper is concerned with variable selection for model-based clustering.

In classification, variable selection is a clearly defined problem, i.e., to find the subset of attributes that gives the best classification performance. The problem is less clear for cluster analysis due to the lack of class information. Several methods have been proposed for model-based clustering. Most of them introduce flexibility into the generative mixture model to allow clusters to be related to subsets of (instead of all) attributes and determine the subsets alongside parameter estimation or during a separate model selection phase. Raftery and Dean [5] consider a variation of the Gaussian mixture model (GMM) where the latent variable is related to a subset of attributes and is independent of other attributes given the subset. A greedy algorithm is proposed to search among those models for one with high BIC score. At each search step, two nested models are compared using the Bayes factor and the better one is chosen to seed the next search step. Law et al. [6] start with the Naïve Bayes model (i.e., GMM with diagonal covariance matrices) and add a saliency parameter for each attribute. The parameter ranges between 0 and 1. When it is 1, the attribute depends on the latent variable. When it is 0, the attribute is independent of the latent variable and its distribution is assumed to be unimodal. The saliency parameters are estimated together with other

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model parameters using the EM algorithm. The work is extended by Li et al. [7] so that the saliency of an attribute can vary across clusters. The third line of work is based on GMMs where all clusters share a common diagonal covariance matrix, while their means may vary. If the mean of a cluster along an attribute turns out to coincide with the overall mean, then that attribute is irrelevant to cluster. Both Bayesian methods [8, 9] and regularization methods [10] have been developed based on this idea.

Our work is based on two observations. First, while clustering algorithms identify clusters in data based on the characteristics of data, domain experts are ultimately the ones to judge the interestingness of the clusters found. Second, high-dimensional data are often multifaceted in the sense that there may be multiple meaningful ways to partition them. The first observation is the reason why variable selection for clustering is such a difficult problem, whereas the second one suggests that the problem may be ill-conceived from the start.

Instead of the variable selection approach, we advocate a *facet determination* approach. The idea is to systematically identify all the different facets of a data set, cluster the data along each one, and present the results to the domain experts for appraisal and selection. The analysis would be useful if one of the clusterings is found interesting.

The difference between the two approaches can be elucidated by comparing their objectives. In variable selection, the aim is to find one subset of attributes that gives the "best" or "good" clustering result. In facet determination, the aim is to find multiple subsets of attributes such that each subset gives meaningful partition of data. In other words, performing facet determination can be considered as performing multiple variable selection simultaneously, without assuming that there is only a single "best" solution.

To realize the idea of facet determination, we generalize the GMMs to allow multiple latent variables. For computational tractability, we restrict that each attribute can be connected to only one latent variable and the relationships among the latent variables can be represented as a tree. The result is what we call pouch latent tree models (PLTMs). Analyzing data using a PLTM may result in multiple latent variables. Each latent variable represents a partition (clustering) of the data and is usually related primarily to only a subset of attributes. Consequently, facets of data can be identified by those subsets of attributes.

Facet determination has been investigated under different designations in previous work. Two other classes of models have been considered. Galimberti and Soffritti [11] use a collection of GMMs, each on a disjoint subset of attributes, for facet determination. We refer to their method as the GS method. The method starts by obtaining a partition of attributes using variable clustering. A GMM is built on each subset of attributes and the collection of GMMs is evaluated using the BIC score. To look for the optimal partition of attributes, the method repeatedly merges the two subsets of attributes that lead to the largest improvement in the BIC score. It stops when no subsets can be merged to improve the score. Similar to the GS models, PLTMs can be considered as containing a collection of GMMs on disjoint subsets of attributes. However, the GMMs in PLTMs are connected by a tree structure, whereas those in the GS models are disconnected.

Latent tree models (LTMs) [12, 13] are another class of models that have been used for facet determination [14]. LTMs and PLTMs are much alike. However, LTMs include only discrete variables and can handle only discrete data. PLTMs generalize LTMs by allowing continuous variables. As a result, PLTMs can also handle continuous data.

Two contributions are made in this paper. The first one is a new class of models in PLTMs, resulting from the marriage between GMMs and LTMs. PLTMs not only generalize GMMs to allow multiple latent variables, but they also generalize LTMs for handling continuous data. The generalization of LTMs is interesting for two reasons. First, it is desirable to have a tool for facet determination on continuous data. Second, the generalization is technically non-trivial. It requires new algorithms for inference and structural learning.

The second contribution is that we compare the variable selection approach and the facet determination approach for model-based clustering. The two approaches have not been compared in the previous studies on facet determination [11, 14]. In this paper, we show that facet determination usually leads to better clustering results than variable selection.

The rest of the paper is organized as follows. Section 2 reviews traditional model-based clustering using GMMs. PLTMs is then introduced in Section 3. In Sections 4–6, we discuss inference, estimation,

and structural learning for PLTMs. The empirical evaluation is divided into two parts. In Section 7, we analyze real-world basketball data using PLTMs. We aim to demonstrate the effectiveness of using PLTMs for facet determination. In Section 8, we compare PLTMs with other methods on benchmark data. We aim to compare the facet determination approach and the variable selection approach. After the empirical evaluation, we discuss related work in Section 9 and conclude in Section 10.

2. Background and Notations

In this paper, we use capital letters such as X and Y to denote random variables, lower case letters such as x and y to denote their values, and bold face letters such as X, Y, x, and y to denote sets of variables or values.

Finite mixture models are commonly used in model-based clustering [15, 16]. In finite mixture modeling, the population is assumed to be made up from a finite number of clusters. Suppose a variable Y is used to indicate this cluster, and variables X represent the attributes in the data. The variable Y is referred to as a *latent* (or unobserved) variable, and the variables X as manifest (or observed) variables. The manifest variables X is assumed to follow a mixture distribution

$$P(\boldsymbol{x}) = \sum_{y} P(y)P(\boldsymbol{x}|y).$$

The probability values of the distribution P(y) are known as mixing proportions and the conditional distributions P(x|y) are known as component distributions. To generate a sample, the model first picks a cluster y according to the distribution P(y) and then uses the corresponding component distribution P(x|y) to generate values for the observed variables.

Gaussian distributions are often used as the component distributions due to computational convenience. A Gaussian mixture model (GMM) has a distribution given by

$$P(\boldsymbol{x}) = \sum_{y} P(y) \mathcal{N}(\boldsymbol{x}|\boldsymbol{\mu}_{y}, \boldsymbol{\Sigma}_{y}),$$

where $\mathcal{N}(x|\mu_y, \Sigma_y)$ is a multivariate Gaussian distribution, with mean vector μ_y and covariance matrix Σ_y conditional on the value of Y.

The Expectation-Maximization (EM) algorithm [17] can be used to fit the model. Once the model is fit, the probability that a data point d belongs to cluster y can be computed by

$$P(y|\mathbf{d}) \propto P(y) \mathcal{N}(\mathbf{d}|\boldsymbol{\mu}_u, \boldsymbol{\Sigma}_u),$$

where the symbol \propto implies that the exact values of the distribution P(y|d) can be obtained by using the sum $\sum_{y} P(y) \mathcal{N}(d|\mu_{y}, \Sigma_{y})$ as a normalization constant.

The number G of components (or clusters) can be given manually or determined by model selection automatically. In the latter case, a score is used to evaluate a model with G clusters. The G that leads to the highest score is then chosen as the estimated number of components. The BIC score has been empirically shown to perform well for this purpose [18].

3. Pouch Latent Tree Models

A pouch latent tree model (PLTM) is a rooted tree, where each internal node represents a latent variable, and each leaf node represents a set of manifest variables. All the latent variables are discrete, while all the manifest variables are continuous. A leaf node may contain a single manifest variable or several of them. Because of the second possibility, leaf nodes are called *pouch nodes*. Figure 1a shows an example of PLTM.

¹In fact, PLTMs allow both discrete and continuous manifest variables. The leaf nodes may contain either a single discrete variable, a single continuous variable, or multiple continuous variables. For brevity, we focus on continuous manifest variables in this paper.

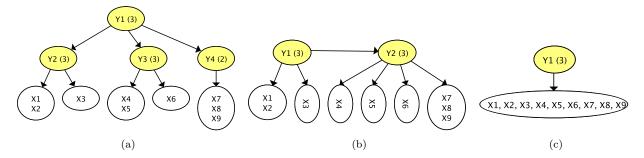


Figure 1: (a) An example of PLTM. The latent variables are shown in shaded nodes. The numbers in parentheses show the cardinalities of the discrete variables. (b) Generative model for synthetic data. (c) GMM as a special case of PLTM.

y_1	$P(y_1)$
s_1	0.33
s_2	0.33
s_3	0.34

	$P(y_2 y_1)$				
y_2	$y_1 = s_1$	$y_1 = s_2$	$y_1 = s_3$		
s_1	0.74	0.13	0.13		
s_2	0.13	0.74	0.13		
s_3	0.13	0.13	0.74		

Table 1: Discrete distributions in Example 1.

In this example, Y_1-Y_4 are discrete latent variables, where Y_1-Y_3 have three possible values and Y_4 has two. X_1-X_9 are continuous manifest variables. They are grouped into five pouch nodes, $\{X_1, X_2\}$, $\{X_3\}$, $\{X_4, X_5\}$, $\{X_6\}$, and $\{X_7, X_8, X_9\}$.

Before we move on, we need to explain some other notations. We use capital letter $\Pi(V)$ to indicate the parent variable of a variable V and lower case letter $\pi(V)$ to denote its value. Also, we reserve the use of bold capital letter W for denoting the variables of a pouch node. When the meaning is clear from context, we use the terms 'variable' and 'node' interchangeably.

In a PLTM, the dependency of a discrete latent variable Y on its parent $\Pi(Y)$ is characterized by a conditional discrete distribution $P(y|\pi(Y))$. Let \boldsymbol{W} be the variables of a pouch node with a parent node $Y = \Pi(\boldsymbol{W})$. We assume that, given a value y of Y, \boldsymbol{W} follows the conditional Gaussian distribution $P(\boldsymbol{w}|y) = \mathcal{N}(\boldsymbol{w}|\boldsymbol{\mu}_y, \boldsymbol{\Sigma}_y)$ with mean vector $\boldsymbol{\mu}_y$ and covariance matrix $\boldsymbol{\Sigma}_y$. A PLTM can be written as a pair $M = (m, \boldsymbol{\theta})$, where m denotes the model structure and $\boldsymbol{\theta}$ denotes the parameters.

Example 1. Figure 1b gives another example of PLTM. In this model, there are two discrete latent variables Y_1 and Y_2 , each having three possible values $\{s_1, s_2, s_3\}$. There are six pouch nodes, namely $\{X_1, X_2\}$, $\{X_3\}$, $\{X_4\}$, $\{X_5\}$, $\{X_6\}$, and $\{X_7, X_8, X_9\}$. The variables in the pouch nodes are continuous.

Each node in the model is associated with a distribution. The discrete distributions $P(y_1)$ and $P(y_2|y_1)$, associated with the two discrete nodes, are given in Table 1.

The pouch nodes \mathbf{W} are associated with conditional Gaussian distributions. These distributions have parameters for specifying the conditional means $\boldsymbol{\mu}_{\pi(\mathbf{W})}$ and conditional covariances $\boldsymbol{\Sigma}_{\pi(\mathbf{W})}$. For the four pouch nodes with single variables, $\{X_3\}$, $\{X_4\}$, $\{X_5\}$, and $\{X_6\}$, these parameters have scalar values. The conditional means μ_y for each of these variables are either -2.5, 0, and 2.5, depending on whether $y=s_1$, s_2 , or s_3 , where y is the value of the corresponding parent variable $Y \in \{Y_1, Y_2\}$. The conditional covariances Σ_y can also have different values for different values of their parents. However, for simplicity in this example, we set $\Sigma_y = 1, \forall y \in \{s_1, s_2, s_3\}$.

Let p be the number of variables in a pouch node. The conditional means are specified by p-vectors and the conditional covariances by $p \times p$ matrices. For example, the conditional means and covariances of the

² The root node is regarded as the child of a dummy node with only one value, and hence is treated in the same way as other latent nodes.

pouch node $\{X_1, X_2\}$ are given by:

$$\boldsymbol{\mu}_{y_1} = \begin{cases} (-2.5, -2.5) & : y_1 = s_1 \\ (0, 0) & : y_1 = s_2 \\ (2.5, 2.5) & : y_1 = s_3 \end{cases} \quad and \quad \boldsymbol{\Sigma}_{y_1} = \begin{pmatrix} 1 & 0.5 \\ 0.5 & 1 \end{pmatrix}, \forall y_1 \in \{s_1, s_2, s_3\}.$$

The conditional means and covariances are specified similarly for pouch node $\{X_7, X_8, X_9\}$. The conditional means for $X_i, i \in \{7, 8, 9\}$, can be -2.5, 0, or 2.5. The variance of any of these variables is 1, and the covariance between any pair of these variables is 0.5.

PLTMs have a noteworthy two-way relationship with GMMs. On the one hand, PLTMs generalize the structure of GMMs to allow more than one latent variable in a model. Thus, a GMM can be considered as a PLTM with only one latent variable and one pouch node containing all manifest variables. As an example, a GMM is depicted as a PLTM in Figure 1c, in which Y_1 is a discrete latent variable and X_1 – X_9 are continuous manifest variables.

On the other hand, the distribution of a PLTM over the manifest variables can be represented by a GMM. Consider a PLTM M. Suppose W_1, \ldots, W_b are the b pouch nodes and Y_1, \ldots, Y_l are the l latent nodes in M. Denote as $X = \bigcup_{i=1}^b W_i$ and $Y = \{Y_j : j = 1, \ldots, l\}$ the sets of all manifest variables and all latent variables in M, respectively. The probability distribution defined by M over the manifest variables X is given by

$$P(\boldsymbol{x}) = \sum_{\boldsymbol{y}} P(\boldsymbol{x}, \boldsymbol{y})$$

$$= \sum_{\boldsymbol{y}} \prod_{j=1}^{l} P(y_j | \pi(Y_j)) \prod_{i=1}^{b} \mathcal{N}(\boldsymbol{w}_i | \boldsymbol{\mu}_{\pi(\boldsymbol{W}_i)}, \boldsymbol{\Sigma}_{\pi(\boldsymbol{W}_i)})$$

$$= \sum_{\boldsymbol{y}} P(\boldsymbol{y}) \mathcal{N}(\boldsymbol{x} | \boldsymbol{\mu}_{\boldsymbol{y}}, \boldsymbol{\Sigma}_{\boldsymbol{y}}).$$
(1)

Equation 1 follows from the model definition. Equation 2 follows from the fact that $\Pi(\mathbf{W}_i), \Pi(Y_j) \in \mathbf{Y}$ and the product of Gaussian distributions is also a Gaussian distribution. Equation 2 shows that $P(\mathbf{x})$ is a mixture of Gaussian distributions. Although it means that PLTMs are not more expressive than GMMs on the distributions of observed data, PLTMs have two advantages over GMMs. First, numbers of parameters can be reduced in PLTMs by exploiting the conditional independence between variables, as expressed by the factorization in Equation 1. Second, and more important, the multiple latent variables in PLTMs allow multiple clusterings on data.

Example 2. In this example, we compare the numbers of parameters in a PLTM and in a GMM. Consider a discrete node and its parent node with c and c' possible values, respectively. It requires $(c-1) \times c'$ parameters to specify the conditional discrete distribution for this node. Consider a pouch node with p variables, and its parent variable with c' possible values. This node has $p \times c'$ parameters for the conditional mean vectors and $\frac{p(p+1)}{2} \times c'$ parameters for the conditional covariance matrices. Now consider the PLTM in Figure 1a and the GMM in Figure 1c. Both of them define a distribution on 9 manifest variables. Based on the above expressions, the PLTM has 77 parameters and the GMM has 164 parameters.

Given the same number of manifest variables, a PLTM may appear to be more complex than a GMM, due to a larger number of latent variables. However, this example shows that a PLTM can still require fewer parameters than a GMM. \Box

The graphical structure of PLTMs looks similar to that of the Bayesian networks (BNs) [19]. In fact, a PLTM is different from a BN only because of the possibility of multiple variables in a single pouch node. It has been shown that any nonsingular multivariate Gaussian distribution can be converted to a complete Gaussian Bayesian network (GBN) with an equivalent distribution [20]. Therefore, a pouch node can be considered as a shorthand notation of a complete GBN. If we convert each pouch node into a complete

GBN, a PLTM can be considered as a conditional Gaussian Bayesian network (i.e., a BN with discrete distributions and conditional Gaussian distributions), or a BN in general.

Cluster analysis based on PLTMs requires learning PLTMs from data. It involves parameter estimation and structure learning. We discuss the former problem in Section 5, and the two problems as a whole in Section 6. Since parameter estimation involves inference on the model, we discuss this problem in the next section before the other two problems.

4. Inference

A PLTM defines a probability distribution P(X, Y) over manifest variables X and latent variables Y. Consider observing values e for the evidence variables $E \subseteq X$. For a subset of variables $Q \subseteq X \cup Y$, we are often required to compute the posterior probability P(q|e). For example, classifying a data point d to one of the clusters represented by a latent variable Y requires us to compute P(y|X = d).

Inference refers to the computation of the posterior probability P(q|e). It can be done on PLTMs similarly as the clique tree propagation on conditional GBNs [21]. However, due to the existence of pouch nodes in PLTMs, this propagation algorithm requires some modifications. The inference algorithm is discussed in details in Appendix A.

The structure of PLTMs allows an efficient inference. Let n be the number of nodes in a PLTM, c be the maximum cardinality of a discrete variable, and p be the maximum number of variables in a pouch node. The time complexity of the inference is dominated by the steps related to message passing and incorporation of evidence on continuous variables. The message passing step requires $O(nc^2)$ time, since each clique has at most two discrete variables due to the tree structure. Incorporation of evidence requires $O(ncp^3)$ time.

Suppose we have the same number of manifest variables. Since PLTMs generally has smaller pouch nodes than GMMs, and hence a smaller p, the term $O(ncp^3)$ shows that inference on PTLMs can be faster than that on GMMs. This happens even though PLTMs can have more nodes and thus a larger n.

5. Parameter Estimation

Suppose there is a data set \mathcal{D} with N samples d_1, \ldots, d_N . Each sample consists of values for the manifest variables. Consider computing the maximum likelihood estimate (MLE) θ^* of the parameters for a given PLTM structure m. We do this using the EM algorithm. The algorithm starts with an initial estimate $\theta^{(0)}$ and improves the estimate iteratively.

Suppose the parameter estimate $\boldsymbol{\theta}^{(t-1)}$ is obtained after t-1 iterations. The t-th iteration consists of two steps, an E-step and an M-step. In the E-step, we compute, for each latent node Y and its parent $\Pi(Y)$, the distributions $P(y, \pi(Y)|\boldsymbol{d}_k, \boldsymbol{\theta}^{(t-1)})$ and $P(y|\boldsymbol{d}_k, \boldsymbol{\theta}^{(t-1)})$ for each sample \boldsymbol{d}_k . This is done by the inference algorithm discussed in the previous section. For each sample k, let \boldsymbol{w}_k be the values of variables \boldsymbol{W} of a pouch node for the sample \boldsymbol{d}_k . In the M-step, the new estimate $\boldsymbol{\theta}^{(t)}$ is obtained as follows:

$$\begin{split} P(y|\pi(Y), \pmb{\theta}^{(t)}) &\propto \sum_{k=1}^{N} P(y, \pi(Y)| \pmb{d}_k, \pmb{\theta}^{(t-1)}) \\ \pmb{\mu}_y^{(t)} &= \frac{\sum_{k=1}^{N} P(y| \pmb{d}_k, \pmb{\theta}^{(t-1)}) \pmb{w}_k}{\sum_{k=1}^{N} P(y| \pmb{d}_k, \pmb{\theta}^{(t-1)})} \\ \pmb{\Sigma}_y^{(t)} &= \frac{\sum_{k=1}^{N} P(y| \pmb{d}_k, \pmb{\theta}^{(t-1)}) (\pmb{w}_k - \pmb{\mu}_y^{(t)}) (\pmb{w}_k - \pmb{\mu}_y^{(t)})'}{\sum_{k=1}^{N} P(y| \pmb{d}_k, \pmb{\theta}^{(t-1)})}, \end{split}$$

where $\boldsymbol{\mu}_y^{(t)}$ and $\boldsymbol{\Sigma}_y^{(t)}$ here correspond to the distribution $P(\boldsymbol{w}|y,\boldsymbol{\theta}^{(t)})$ for node \boldsymbol{W} conditional on its parent $Y = \Pi(\boldsymbol{W})$. The EM algorithm proceeds to the (t+1)-th iteration unless the improvement of log-likelihood $\log P(\mathcal{D}|\boldsymbol{\theta}^{(t)}) - \log P(\mathcal{D}|\boldsymbol{\theta}^{(t-1)})$ falls below a certain threshold.

The starting values of the parameters $\boldsymbol{\theta}^{(0)}$ are chosen as follows. For $P(y|\pi(Y),\boldsymbol{\theta}^{(0)})$, the probabilities are randomly generated from a uniform distribution over the interval (0,1] and are then normalized. The initial values of $\boldsymbol{\mu}_y^{(0)}$ are set to equal to a random sample from data, while those of $\boldsymbol{\Sigma}_y^{(0)}$ are set to equal to the sample covariance.

Like in the case of GMMs, the likelihood is unbounded in the case of PLTMs. This might lead to spurious local maxima [15]. For example, consider a mixture component that consists of only one data point. If we set the mean of the component to be equal to that data point and set the covariance to zero, then the model will have an infinite likelihood on the data. However, even though the likelihood of this model is higher than some other models, it does not mean that the corresponding clustering is better. The infinite likelihood can always be achieved by trivially grouping one of the data points as a cluster. This is why we refer to this kind of local maxima as spurious.

To mitigate this problem, we use a variant of the method by Ingrassia [22]. In the M-step of EM, we need to compute the covariance matrix $\Sigma_y^{(t)}$ for each pouch node W. We impose the following constraints on the eigenvalues $\lambda^{(t)}$ of $\Sigma_y^{(t)}$: $\sigma_{min}^2/\gamma \leq \lambda^{(t)} \leq \sigma_{max}^2 \times \gamma$, where σ_{min}^2 and σ_{max}^2 are the minimum and maximum of the sample variances of the variables W and γ is a parameter for our method.

6. Structure Learning

Given a data set \mathcal{D} , we aim at finding the model m^* that maximizes the BIC score [23]:

$$BIC(m|\mathcal{D}) = \log P(\mathcal{D}|m, \theta^*) - \frac{d(m)}{2} \log N,$$

where θ^* is the MLE of the parameters and d(m) is the number of independent parameters in m. The first term is known as the likelihood term. It favors models that fit data well. The second term is known as the penalty term. It discourages complex models. Hence, the BIC score provides a trade-off between model fit and model parsimoniousness.

We have developed a hill-climbing algorithm to search for m^* . It starts with a model $m^{(0)}$ that contains one latent node as root and a separate pouch node for each manifest variable as a leaf node. The latent variable at the root node has two possible values. Suppose a model $m^{(j-1)}$ is obtained after j-1 iterations. In the j-th iteration, the algorithm uses some search operators to generate candidate models by modifying the base model $m^{(j-1)}$. The BIC score is then computed for each candidate model. The candidate model m' with the highest BIC score is compared with the base model $m^{(j-1)}$. If m' has a higher BIC score than $m^{(j-1)}$, m' is used as the new base model $m^{(j)}$ and the algorithm proceeds to the (j+1)-th iteration. Otherwise, the algorithm terminates and returns $m^* = m^{(j-1)}$ (together with the MLE of the parameters).

We now describe the search operators used. There are four aspects of the structure m, namely, the number of latent variables, the cardinalities of these latent variables, the connections between variables, and the composition of pouches. The search operators used in our hill-climbing algorithm modify all these aspects to effectively explore the search space. There are totally seven search operators. Five of them are borrowed from Zhang and Kočka [24], while two of them are new for PLTMs.

We first described the five operators borrowed from others. A node introduction (NI) operator involves one latent node Y and two of its neighbors. It creates a new model by introducing a new latent node Y_{new} to mediate between Y and the two neighbors. The cardinality of Y_{new} is set to be that of Y. A node deletion (ND) operator is the opposite of NI. It involves two neighboring latent nodes Y and Y_{delete} . It creates a new model by deleting Y_{delete} and making all neighbors of Y_{delete} (other than Y) neighbors of Y. Given a latent variable in PLTM, a state introduction (SI) operator creates a new model by adding a state to the domain of the variable. A state deletion (SD) operator does the opposite. A node relocation (NR) involves a node V, one of its latent node neighbors Y_{origin} and another latent node Y_{dest} . The node V can be a latent node or a pouch node. The NR operator creates a new model by relocating V to Y_{dest} , i.e., removing the link between V and Y_{origin} and adding a link between V and Y_{dest} . Figure 2 gives some examples of the use of NI, ND, and NR operators.

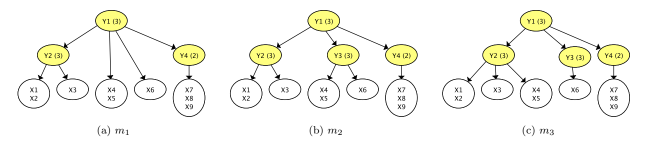


Figure 2: Examples of applying the node introduction, node deletion, and node relocation operators. Introducing Y_3 to mediate between Y_1 , $\{X_4, X_5\}$ and $\{X_6\}$ in m_1 gives m_2 . Relocating $\{X_4, X_5\}$ from Y_3 to Y_2 in m_2 gives m_3 . In reverse, relocating $\{X_4, X_5\}$ from Y_2 to Y_3 in m_3 gives m_2 . Deleting Y_3 in m_2 gives m_1 .

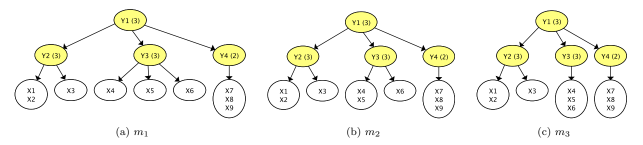


Figure 3: Examples of applying the pouching and unpouching operators. Pouching $\{X_4\}$ and $\{X_5\}$ in m_1 gives m_2 , and pouching $\{X_4, X_5\}$ and $\{X_6\}$ in m_2 gives m_3 . In reverse, unpouching X_6 from $\{X_4, X_5, X_6\}$ in m_3 gives m_2 , and unpouching X_5 from $\{X_4, X_5\}$ gives m_1 .

The two new operators are pouching (PO) and unpouching (UP) operators. The PO operator creates a new model by combining a pair of sibling pouch nodes \boldsymbol{W}_1 and \boldsymbol{W}_2 into a new pouch node $\boldsymbol{W}_{po} = \boldsymbol{W}_1 \cup \boldsymbol{W}_2$. The UP operator creates a new model by separating one manifest variable X from a pouch node \boldsymbol{W}_{up} , resulting in two sibling pouch nodes $\boldsymbol{W}_1 = \boldsymbol{W}_{up} \setminus \{X\}$ and $\boldsymbol{W}_2 = \{X\}$. Figure 3 shows some examples of the use of these two operators.

The purpose of the PO and UP operators is to modify the conditional independencies entailed by the model on the variables of the pouch nodes. For example, consider the two models m_1 and m_2 in Figure 3. In m_1 , X_4 and X_5 are conditionally independent given Y_3 , i.e., $P(X_4, X_5|Y_3) = P(X_4|Y_3)P(X_5|Y_3)$. In other words, covariance between X_4 and X_5 is zero given Y_3 . On the other hand, X_4 and X_5 need not be conditionally independent given Y_3 in m_2 . The covariances between them are allowed to be non-zero in the 2×2 conditional covariance matrices for the pouch node $\{X_4, X_5\}$.

The PO operator in effect postulates that two sibling pouch nodes are correlated given their parent node. It may improve the BIC score of the candidate model by increasing the likelihood term, when there is some degree of local dependence between those variables on the empirical data. On the other hand, the UP operator postulates that one variable in a pouch node is conditionally independent from other variables in the pouch node. It reduces the number of parameters in the candidate model and hence may improve the BIC score by decreasing the penalty term. These postulates are tested by comparing the BIC scores of the corresponding models in each search step. The postulate that leads to the model with the highest BIC score is considered as most appropriate.

For the sake of computational efficiency, we do not consider introducing a new node to mediate between Y and more than two of its neighbors. This restriction can be compensated by considering a restricted version of node relocation after a successful node introduction. Suppose Y_{new} is introduced to mediate between Y and its two neighbors. The restricted version of NR relocates one of the neighbors of Y (other than Y_{new}) to Y_{new} . The PO operator also has similar issue. Hence, after a successful pouching, we consider

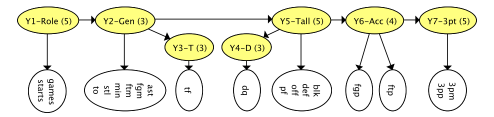


Figure 4: PLTM obtained on NBA data. The latent variables represent different clusterings on the players. They have been renamed based on our interpretation of their meanings. The abbreviations in these names stand for: role (Role), general ability (Gen), technical fouls (T), disqualification (D), tall-player ability (Tall), shooting accuracy (Acc), and three-pointer ability (3pt).

a restricted version of PO. The restricted version combines the new pouch node resulting from PO with one of its sibling pouch nodes.

The above explains the basic principles for the hill-climbing algorithm. However, the algorithm as outlined above is inefficient and can handle only small data sets. We have developed acceleration techniques that make the algorithm efficient enough for some real-world applications. We leave these details in Appendix B.

7. Facet Determination on NBA Data

In the first part of our empirical study, we aim to demonstrate the effectiveness of facet determination by PLTMs. Real-world basketball data were used in this study. The objective is to see whether the facets identified and the clusterings given by PLTM on this data set are meaningful or not. The analysis can be considered as effective if the facets and clusterings are meaningful in general. We interpret the clusterings based on our basic basketball knowledge.

The data contain seasonal statistics of National Basketball Association (NBA) players. They were collected from 441 players who played in at least one game in the 2009/10 season. Each sample corresponds to one player. It includes the numbers of games played (games) and started (starts) by the player in that season. It also contains 16 other per game averages or seasonal percentages, including minutes (min), field goals made (fgm), field goal percentage (fgp), three pointers made (3pm), three pointer percentage (3pp), free throws made (ftm), free throw percentage (ftp), offensive rebounds (off), defensive rebounds (def), assists (ast), steals (stl), blocks (blk), turnovers (to), personal fouls (pf), technical fouls (tf), and disqualifications (dq). In summary, the data set has 18 attributes and 441 samples.

7.1. Facets Identified

We built a PLTM on the data and the structure of the resulting model is shown in Figure 4. The model contains 7 latent variables. Each of the them identifies a different facet of data. The first facet consists of attributes games and starts, which are related to the role of a player. The second one consists of attributes min, fgm, ftm, ast, stl, and to, which are related to some general performance of a player. The third and fourth facets each contain only one attribute. They are related to tf and dq, respectively. The fifth facet contains attributes blk, off, def, and pf. A player usually needs to jump higher than other players to get a block (blk) or a rebound (off or def). Therefore, the facet is mainly related to an aspect of performance in which taller players have an advantage. The sixth facet consists of two attributes ftp and fgp, which are related to the shooting accuracy. The last facet contains 3pm and 3pp, which are related to three pointers.

The facets identified consist mostly of related attributes. Hence, they appear to be reasonable in general.

³ The data were obtained from: http://www.dougstats.com/09-10RD.txt

Role	$\mathbf{P}(\mathtt{Role})$	games	starts
occasional players	0.32	29.1	2.0
irregular starters	0.11	46.1	31.7
regular substitutes	0.19	68.2	5.4
regular players	0.13	75.8	32.8
regular starters	0.25	76.0	73.7
overall	1.00	56.3	27.9

(a) Role of player (Role)

3pt	P(3pt)	3pm	Зрр
never	0.29	0.00	0.00
seldom	0.12	0.08	0.26
fair	0.17	0.33	0.28
good	0.40	1.19	0.36
extreme	0.02	0.73	0.64
overall	1.00	0.55	0.23

Acc	$\mathbf{P}(\mathtt{Acc})$	fgp	ftp
low ftp	0.10	0.44	0.37
low fgp	0.16	0.39	0.72
high ftp	0.47	0.44	0.79
high fgp	0.28	0.52	0.67
overall	1.00	0.45	0.71

(b) Three-pointer ability (3pt)

(c) Shooting accuracy (Acc)

Table 2: Attribute means on three different facets on NBA data. The means are conditional on the specified latent variables. The second columns show the marginal distributions of those latent variables. The last rows show the unconditional means of the attributes. The clusters are named based on our interpretation.

7.2. Meanings of Clusters

To understand the meanings of the clusters, we may examine the mean values of attributes of each cluster. We use three clusterings, Role, 3pt, and Acc, for illustration.

Table 2a shows the means of games and starts conditional on the clustering variable Role. Note that there are 82 games in a NBA season. We see that the mean value of games for the players in the first cluster is far below the overall mean. Hence, the first cluster contains players who played occasionally. The second group of players also played less often than average, but they usually started in a game when they played. This may look counter-intuitive. However, the cluster probably refers to those players who had the calibre of starters but had missed part of the season due to injuries. The third group of players played often, but usually as substitutes (not as starters). The fourth group of players played regularly and sometimes started in a game. The last group contains players who played and started regularly.

Table 2b shows the means of 3pm (three-pointer made) and 3pp (three-pointer percentage) conditional on 3pt (three-pointer ability). The variable 3pt partitions players into five clusters. The first two clusters contain players that never and seldom made a three-pointer, respectively. The next two clusters contains players that have fair and good three-pointer accuracies, respectively. The last group is an extreme case. It contains players shooting with unexpectedly high accuracy. As indicated by the marginal distribution, it consists of only a very small proportion of players. This is possibly a group of players who had made some three pointers during the sporadic games that they had played. The accuracy remained very high since they did not play often.

Table 2c shows the means of fgp (field goal percentage) and ftp (free throw percentage) conditional on Acc (shooting accuracy). The first three groups contain players of low ftp, low fgp, and high ftp, respectively. The fourth group of players had high fgp. However, those players also had below-average ftp. Both ftp and fgp are related to the shooting accuracies. Hence, one may expect that the two attributes should be positively correlated and the fourth group may look unreasonable. However, the group is indeed sensible due to the following observation. Taller players usually stay closer to basket in games. Therefore, they take high-percentage shots more often and have higher field goal percentage. On the other hand, taller players are usually poorer in making free throws and have lower free throw percentage. A typical example of such players is Dwight Howard, who is listed as 6'11" tall. He had a relatively high fgp (0.61) but a

	Gen				
Role	poor	fair	good		
occasional players	0.81	0.19	0.00		
irregular starters	0.00	0.69	0.31		
regular substitutes	0.22	0.78	0.00		
regular players	0.00	0.81	0.19		
regular starters	0.00	0.06	0.94		

	Acc					
Tall	low ftp	low fgp	high ftp	high fgp		
poor	0.28	0.53	0.00	0.18		
fair	0.00	0.02	0.95	0.03		
good	0.00	0.00	1.00	0.00		
good_big	0.11	0.04	0.00	0.86		
very good	0.00	0.00	0.14	0.86		

⁽a) P(Gen|Role): relationship between general ability (b) P(Acc|Tall): relationship between shooting accuracy (Acc) (Gen) and role (Role)

and tall-player ability (Tall)

Table 3: Conditional distributions of Gen and Acc on NBA data.

Subset of attributes	#clusters
starts, min, fgm, 3pm, ftm, off, def, ast, stl, to, blk, pf	11
games, tf	3
fgp, 3pp, ftp, dq	2

Table 4: Partition of attributes on NBA data by the GS method. The last column lists the number of clusters in the GMM built for each subset of attributes.

relatively low ftp (0.59). He was classified appropriately by the PLTM as "high fgp".

7.3. Relationships between Clusterings

Latent variables in PLTMs are connected. The edges between them represent probabilistic relationships between the clusterings. Therefore, PLTMs can also show some possibly interesting relationships between clusterings. We use two examples to demonstrate this.

Table 3a shows the conditional distribution P(Gen|Role). The clustering Gen consists of three clusters of players with poor, fair, and good general performances, respectively. We observe that those players playing occasionally had mostly poor performance, and almost all starters played well. The other three groups of players usually played fairly. However, more of the irregular starters played well than the regular players. Also none of the regular substitutes played well. The above relationship looks reasonable because the role of a player should be related to his general performance.

Table 3b shows the conditional distribution P(Acc|Tall). It is consistent with our observation that taller players usually shoot free throws more poorly. Most players who played very well ("very good") or particularly well ("good_big") as tall players belong to the group that has high field goal percentage but low free throw percentage ("high_fgp"). On the other hand, those who do not play well specifically as tall players ("fair" and "good") usually have average field goal percentage and higher free throw percentage ("high_ftp"). For those who played poorly as tall players, we cannot tell much about them.

7.4. Comparison with the GS Model

The GS models, mentioned in Section 1, contain a collection of GMMs, each built on a subset of attributes. These models can also be used for facet determination. We now compare the results obtained from the GS model and those from the PLTM.

Table 4 shows the partition of attributes given by the GS method. Each row corresponds to a subset of attributes and identifies a facet. A clustering is given by the GMM built on the subset of attributes. The number of clusters for each clustering is shown in the last column.

Compared to the results obtained from PLTM analysis, those from the GS method have three weaknesses. First, the facets found by the GS method appear to be less natural than those by PLTM analysis. In particular, attribute games can be related to many aspects of the game statistics. However, it is grouped together by the GS method with a less interesting attribute tf, which indicates the number of technical fouls, in the second subset. In the third subset, attributes fgp, 3pp, ftp are all related to shooting percentages. However, they are also grouped together with an apparently unrelated attribute dq (disqualifications). The first subset lumps together a large number of attributes. This means it has missed some more specific and meaningful facets that have been identified by PLTM analysis.

The second weakness is related to the numbers of clusters given by the GS method. One the one hand, there are a large number of clusters on the first subset. This makes it difficult to comprehend the clustering, especially with many attributes in this subset. On the other hand, there are only few clusters on the second and third subsets. This means some subtle clusters found in PLTM analysis were not found by the GS method. The third weakness is inherent in the structure of the GS models. Since disconnected GMMs are used, the latent variables are assumed to be independent. Consequently, the GS models cannot show those possibly meaningful relationships between the clusterings as PLTMs do.

7.5. Discussions

The results presented above in general show that PLTM analysis identified reasonable facets and suggest that it gave meaningful clusterings on those facets. We also see that better results were obtained from PLTM analysis than the GS method. Therefore, we conclude that PLTMs can perform facet determination effectively on NBA data.

If we think about how basketball games are played, we can expect that the heterogeneity of players can originate from various aspects, such as positions of the players, their competence on their corresponding positions, or their general competence. As our results show, PLTM analysis identified these different facets from the NBA data and allowed users to partition data based on them separately. If traditional clustering methods are used, only one clustering can be obtained, regardless of whether variable selection is used or not. Therefore, some of the facets cannot be identified by the traditional methods.

The number of attributes in NBA data may be small relatively to those data available nowadays. Nevertheless, we can still identify multiple facets and obtain multiple meaningful clusterings from the data. We thus can expect real-world data with higher dimensions are also multifaceted. Consequently, it is in general more appropriate to use the facet determination approach with PLTMs than the variable selection selection approach for clustering high-dimensional data.

8. Facet Determination versus Variable Selection

In the previous section, we have demonstrated that multiple meaningful clusters can be found on a data set. In such cases, variable selection methods are inadequate in the sense they can produce only single clustering solutions and thus cannot identify all the meaningful clusterings. However, it is still possible that the single clustering produced by a variable selection method is more interesting to users than all the clusterings produced by a facet determination method.

In the second part of the empirical study, we aim to address this issue and compare the two approaches for clustering. We want to see which method can produce the most interesting clustering. Due to the lack of domain experts, we use data with class labels. We assume that the partition indicated by the class labels is the partition most interesting to users on a data set. The method that can produce a partition closest to the class partition is deemed to be the best method in this study.

8.1. Data Sets and Methods

We used both synthetic and real-world data sets in this study. The synthetic data were generated from the model described in Example 1, of which the variable Y_1 is designated as the class variable. The real-world data sets were borrowed from the UCI machine learning repository.⁴ We chose 9 labeled data sets that have often been used in the literature and that contain only continuous attributes. Table 5 shows the basic information of these data sets.

⁴ http://www.ics.uci.edu/~mlearn/MLRepository.html

Data Set	#Attributes	#Classes	#Samples	#Latents
glass	9	6	214	3.0
image	18^{5}	7	2310	4.4
ionosphere	33^{5}	2	351	9.9
iris	4	3	150	1.0
vehicle	18	4	846	3.0
wdbc	30	2	569	9.4
wine	13	3	178	2.0
yeast	8	10	1484	5.0
zernike	47	10	2000	6.9

Table 5: Descriptions of UCI data sets used in our experiments. The last column shows the average numbers of latent variables obtained by PTLM analysis over 10 repetitions.

We compare PLTM analysis with five methods based on GMMs. The first method is plain GMM (PGMM) analysis. The second one is MCLUST [25],⁶ which reduces the number of parameters by imposing constraints on the eigenvalue decomposition of the covariance matrices. The third one is CLUST-VARSEL [5],⁷ which is denoted as CVS for short. The last one is the method of Law et al. [6], which we call LFJ, using the the first letters of the three author names. Among these four methods, the last two perform variable selection while the first two do not. The fifth method is the GS method [11]. Similar to PLTM analysis, it follows the facet determination approach. Note that this study aims to compare methods following the variable selection approach or the facet determination approach. Those methods without variable selection are included for reference.

In our experiments, the parameters of PGMMs and PLTMs were estimated using the EM algorithm. The same settings were used for both cases. EM was terminated when it failed to improve the log-likelihood by 0.01 in one iteration or when the number of iterations reached 500. We used a variant of multiple-restart approach [26] with 64 starting points to avoid local maxima in parameter estimation. For the scheme to avoid spurious local maxima in parameter estimation as described in Section 5, we set the constant γ at 20. For PGMM and CVS, the true numbers of classes were given as input. For MCLUST, LFJ, and GS, the maximum number of mixture components was set at 20.

8.2. Method of Comparison

Our experiments started with labeled data. In the training phase, models were learned from data with the class labels removed. In the testing phase, the clusterings contained in models were evaluated and compared based on the class labels. The objective is to see which method recovers the class variable the best.

A model produced by PGMM, MCLUST, CVS, or LFJ contains a single latent variable Y. It represents one way to partition the data. We follow Strehl and Ghosh [27] and evaluate the partition using normalized mutual information NMI(C;Y) between Y and the class variable C. The NMI is given by

$$NMI(C;Y) = \frac{MI(C;Y)}{\sqrt{H(C)H(Y)}},$$

where MI(C;Y) is the mutual information between C and Y and H(V) is the entropy of a variable V [28]. These quantities can be computed from P(c,y), which in turn is estimated by $P(c,y) = \frac{1}{N} \sum_{k=1}^{N} I_{c_k}(c) P(y|\boldsymbol{d}_k)$, where $\boldsymbol{d}_1, \ldots, \boldsymbol{d}_N$ are the samples in testing data, $I_{c_k}(c)$ is an indicator function having value of 1 when $c = c_k$ and 0 otherwise, and c_k is the class label for the k-th sample.

⁵Attributes having single values had been removed.

⁶ http://cran.r-project.org/web/packages/mclust

⁷ http://cran.r-project.org/web/packages/clustvarsel

	Facet Determination		Variable Selection		No Variable Selection	
Data Set	PLTM	GS	CVS	LFJ	PGMM	MCLUST
synthetic	.85 (.00)	.69 (.00)	.34 (.00)	.56 (.02)	.56 (.00)	.64 (.00)
glass	.43 (.03)	.38 (.00)	.29 (.00)	.35 (.03)	.28 (.03)	.33 (.00)
image	.71 (.03)	.65 (.00)	.41 (.00)	.51 (.03)	.52 (.04)	.66 (.00)
vehicle	.40 (.04)	.31 (.00)	.23 (.00)	.27 (.01)	.25 (.08)	.36 (.00)
wine	.97 (.00)	.83 (.00)	.71 (.00)	.70 (.19)	.50 (.06)	.69 (.00)
zernike	.50 (.02)	.39 (.00)	.33 (.00)	.45 (.01)	.44 (.03)	.41 (.00)
ionosphere	.36 (.01)	.26 (.00)	.41 (.00)	.13 (.07)	.57 (.04)	.32 (.00)
iris	.76 (.00)	.74 (.00)	.87 (.00)	.68 (.02)	.73 (.08)	.76 (.00)
wdbc	.45 (.03)	.36 (.00)	.34 (.00)	.41 (.02)	.44 (.08)	.68 (.00)
yeast	.18 (.00)	.22 (.00)	.04 (.00)	.11 (.04)	.16 (.01)	.11 (.00)

Table 6: Clustering performances as measured by NMI. The averages and standard deviations over 10 repetitions are reported. Best results are highlighted in bold. The first row categorizes the methods according to their approaches.

A model resulting from PLTM analysis or the GS method contains a set Y of latent variables. Each of the latent variables represents a partition of the data. In practice, the user may find several of the partitions interesting and use them all in his work. In this section, however, we are talking about comparing different clustering algorithms in terms of the ability to recover the original class partition. So, the user needs to choose one of the partitions as the final result. The question becomes whether this analysis provides the possibility for the user to recover the original class partition. Consequently, we assume that the user chooses, among all the partitions produced, the one closest to the class partition and we evaluate the performance of PLTM analysis and the GS method using this quantity:

$$\max_{Y \in \boldsymbol{Y}} NMI(C;Y).$$

Note that NMI was used in our experiments due to the absence of a domain expert to evaluate the clustering results. In practice, class labels are not available when we cluster data. Hence, NMI cannot be used to select the appropriate partitions in the facet determination approach. A user needs to interpret the clusterings and find those interesting to her. It might also be possible to use clustering validity indices [29] for selection. Investigation into this possibility is left for future research.

8.3. Results

The results of our experiments are given in Table 6. In terms of NMI, PLTM had clearly superior performances over the two variable selection methods, CVS and LFJ. Specifically, it outperformed CVS on all but one data sets and LFJ on all data sets. PLTM also performed better than GS, the other facet determination method. It outperformed GS on all but one data. Besides, PLTM has clear advantages over PGMM and MCLUST, the two methods that do not do variable selection. PLTM outperformed PGMM on all but one data sets and outperformed MCLUST except for two data sets. On those data sets in our experiments, PLTM usually outperformed the other methods by large margins.

Note that multiple clusterings are produced by PLTMs and maximum NMI scores are used for comparison. It is therefore possible that not all of the clusterings produced by PLTM analysis have higher NMI scores than clusterings obtained by other methods, even if PLTM analysis has a higher maximum NMI score. Moreover, the clusterings produced by other methods may appear more interesting if the interest of users lies not in the class partitions. Nevertheless, our comparison assumes that users are only interested in the class partitions. Indeed, this is an assumption taken implicitly when the class partitions are used for the evaluation of clusterings. Based on this assumption, the experimental results indicate that PLTM analysis usually provides the best possibility for recovering the partitions of interest.

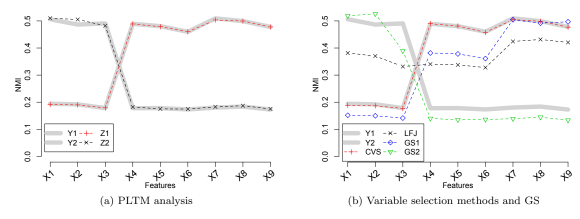


Figure 5: Feature curves of the partitions obtained by various methods and that of the original class partition on synthetic data.

8.4. Explaining the Results

We next examine models produced by the various methods to gain insights about the superior performance of PLTM analysis. This also allows us to judge the facets identified by PLTM analysis.

8.4.1. Synthetic Data

Before examining models obtained from synthetic data, we first take a look at the data set itself. The data were sampled from the model shown in Figure 1b, with information about the two latent variables Y_1 and Y_2 removed. Nonetheless, the latent variables represent two natural ways to partition the data. To see how the partitions are related to the attributes, we plot the NMI⁸ between the latent variables and the attributes in Figure 5a. We call the curve for a latent variable its feature curve. We see that Y_1 is strongly correlated with X_1 – X_3 , but not with the other attributes. Hence it represents a partition based on those three attributes. Similarly, Y_2 represents a partition of the data based on attributes X_4 – X_9 . So, we say that the data has two facets, one represented by X_1 – X_3 and another by X_4 – X_9 . The designated class partition Y_1 is a partition along the first facet.

The model produced by PLTM analysis has the same structure as the generative model. We name the two latent variables in the model Z_1 and Z_2 respectively. Their feature curves are also shown in Figure 5a. We see that the feature curves of Z_1 and Z_2 match those of Y_1 and Y_2 well. This indicates that PLTM analysis has successfully recovered the two facets of the data. It has also produced a partition of the data along each of the facets. If the user chooses the partition Z_1 along the facet X_1 – X_3 as the final result, then the original class partition is well recovered. This explains the good performance of PLTM (NMI=0.85).

The feature curves of the partitions obtained by LFJ and CVS are shown in Figure 5b. We see that the LFJ partition is not along any of the two natural facets of the data. Rather it is a partition based on a mixture of those two facets. Consequently, the performance of LFJ (NMI=0.56) is not as good as that of PLTM. CVS did identify the facet represented by X_4 – X_9 , but it is not the facet of the designated class partition. In other words, it picked the wrong facet. Consequently, the performance of CVS (NMI=0.34) is the worst among all the methods considered. GS succeeded to identify two facets. However, the their feature curves do not match those of Y_1 and Y_2 well. This probably explains why its performance (NMI=0.69) is worse than PLTM.

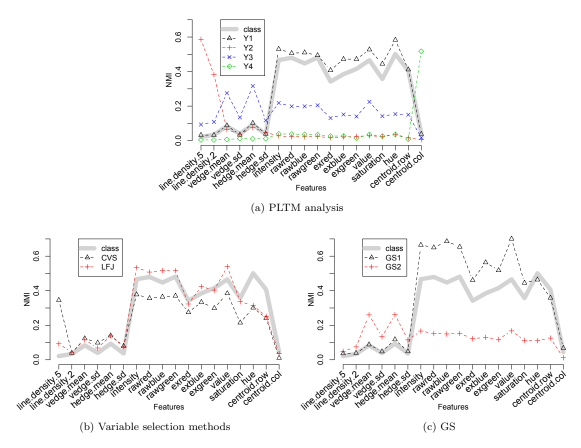


Figure 6: Feature curves of the partitions obtained by various methods and that of the original class partition on image data.

8.4.2. Image Data

In the image data, each instance represents a 3×3 region of an image. It is described by 18 attributes. The feature curve of the original class partition is given in Figure 6a. We see that it is a partition based on 10 color-related attributes from intensity to hue and the attribute centroid.row.

The structure of the model produced by PLTM analysis is shown in Figure 7. It contains 4 latent variables Y_1-Y_4 . Their feature curves are shown in Figure 6a. We see that the feature curve of Y_1 matches that of the class partition beautifully. If the user chooses the partition represented by Y_1 as the final result, then the original class partition is well recovered. This likely explains the good performance of PLTM (NMI=0.71).

The feature curves of the partitions obtained by LFJ and CVS are shown in Figure 6b. The LFJ curve matches that of the class partition quite well, but not as well as the feature curve of Y_1 , especially on the attributes line.density.5, hue and centroid.row. This is a possible reason why the performance of LFJ (NMI=0.51) is not as good as that of PLTM. Similar things can be said about the partition obtained by CVS. Its feature curve differs from the class feature curve even more than the LFJ curve on the attribute line.density.5, which is irrelevant to the class partition. Consequently, the performance of CVS (NMI=0.41) is even worse than that of LFJ.

GS produced four partitions on image data. Two of them comprise only one component and hence are discarded. The feature curves of the remaining two partitions are shown in Figure 6c. We see that one of them corresponds to the facet of the class partition, but does not match that well. This likely explains why

⁸To compute NMI(X;Y) between a continuous variable X and a latent variable Y, we discretized X into 10 equal-width bins, so that P(X,Y) could be estimated as a discrete distribution.

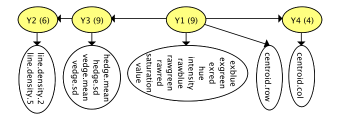


Figure 7: Structure of the PLTM learned from image data.

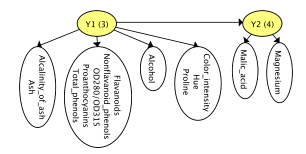


Figure 8: Structure of the PLTM learned from wine data.

the performance of GS (NMI=0.65) is better than the other methods but is not as good as that of PLTM. Two remarks are in order. First, the 10 color-related attributes semantically form a facet of the data. PLTM analysis has identified the facet in the pouch below Y_1 . Moreover, it obtained a partition based on not only the color attributes, but also the attribute centroid row, the vertical location of a region in an

PETM analysis has identified the facet in the pouch below Y_1 . Moreover, it obtained a partition based on not only the color attributes, but also the attribute centroid.row, the vertical location of a region in an image. This is interesting. It is because centroid.row is closely related to the color facet. Intuitively, the vertical location of a region should correlate with the color of the region. For example, the color of the sky occurs more frequently at the top of an image and that of grass more frequently at the bottom.

Second, the latent variable Y_2 is strongly correlated with the two line density attributes. This is another facet of the data that PLTM analysis has identified. PLTM analysis has also identified the edge-related facet in the pouch node below Y_3 . However, it did not obtain a partition along the facet. The partition represented by Y_3 depends on not only the edge attributes but others as well. The two coordinate attributes centroid.row and centroid.col semantically form one facet. The facet has not been identified probably because the two attributes are not correlated.

8.4.3. Wine Data

The PLTM learned from wine data is shown in Figure 8. The model structure also appears to be interesting. While we are not experts on wine, it seems natural to have Ash and Alcalinity_of_ash in one pouch as both are related to ash. Similarly, Flavanoids, Nonflavanoid_phenols, and Total_phenols are related to phenolic compounds. These compounds affect the color of wine, so it is reasonable to have them in one pouch along with the opacity attribute OD280/OD315. Moreover, both Magnesium and Malic_acid play a role in the production of ATP (adenosine triphosphate), the most essential chemical in energy production. So, it is not a surprise to find them connected to a second latent variable.

8.4.4. WDBC Data

After examining some positive cases for PLTM analysis, we now look at a data set that PLTM did not perform so well. Here we consider wdbc data. The data were obtained from 569 digitalized images of cell

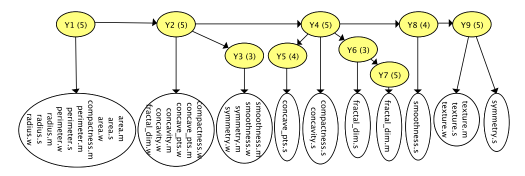


Figure 9: Structure of the PLTM learned from wdbc data.

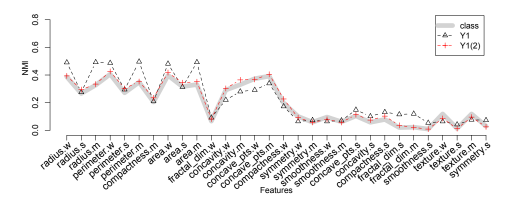


Figure 10: Features curves of the partition Y_1 , obtained by PLTM, and that of the original class partition on wdbc data. $Y_1(2)$ is obtained by setting the cardinality of Y_1 to 2.

nuclei aspirated from breast masses. Each image corresponds to a data instance. It is labeled as either benign or malignant. There are 10 computed features of the cell nuclei in the images. The attributes consist of the mean value (m), standard error (s), and worst value (w) for each of these features.

Figure 9 shows the structure of a PLTM learned from this data. We can see that this model identifies some meaningful facets. The pouch below Y_1 identifies a facet related to the size of nuclei. It mainly includes attributes related to area, perimeter, and radius. The second facet is identified by the pouch below Y_2 . It is related to concavity and consists primarily of the mean and worst values of the two features related to concavity. The third facet is identified by the pouch below Y_3 . It includes the mean and worst values of smoothness and symmetry. The facet appears to show whether the nuclei have regular shapes or not. The pouch below Y_9 identifies a facet primarily related to texture. This facet includes three texture-related attributes but also the attribute symmetry.s. The remaining attributes are mostly standard errors of some features and may be considered as the amount of variation of the features. They are connected to the rest of the model through Y_4 and Y_8 .

Although the model appears to have a reasonable structure, it did not achieve a high NMI on this data (NMI=0.45). To have better understanding, we compare the feature curve of the class partition with that of the closest partition (Y_1) given by PLTM in Figure 10. The two feature curves have roughly similar shapes. We also look at the confusion matrix for Y_1 (Table 7). We see that Y_1 gives a reasonable partition. The first four states of Y_1 group together the benign cases or malignant cases almost perfectly, while the remaining state groups together some uncertain cases.

	Y_1				
class	s1	s2	s3	s4	s5
malignant	43	116	6	0	47
benign	0	9	193	114	41

Table 7: Confusion matrix for PLTM on wdbc data.

One possible reason for the relatively low NMI is that Y_1 has 5 states but the class variable has only 2 states. The higher number of states of Y_1 may lead to a lower NMI due to an increase of the entropy term. It may also lead to the mismatch of the feature curves. To verify this explanation, we manually set the cardinality of Y_1 to 2. The feature curve of this adjusted latent variable (denoted by $Y_1(2)$) is shown in Figure 10. It now matches the feature curve of the class partition well. The adjustment also improved the NMI to 0.69, and made it become highest on this data set. This example shows that an incorrect estimated number of clusters could be a reason why PLTM performed worse than other methods on some data sets.

8.4.5. Discussions

We have also performed PLTM analysis on the other data sets. The last column of Table 5 lists the average numbers of latent variables obtained over 10 repetitions. We see that multiple latent variables have been identified except on iris data, which has only four attributes. Many of the latent variables represent partitions of data along natural facets of the data.

In general, PLTM analysis has the ability to identify natural facets of data and cluster data along those facets. In practice, a user may find several of the clusterings useful. In the setting where clustering algorithms are evaluated using labeled data, PLTM performs well if the original class partition is also along some of the natural facets, and poorly otherwise.

9. Related Work

Some recent work also produces multiple clusterings. One important difference between PLTMs and the recent work lies in the relationship between clusterings. PLTMs allow the multiple clusterings to be correlated. On the other hand, most recent work attempts to find multiple clusterings that are dissimilar from each other. Two approaches are commonly used. The first approach finds multiple clusterings sequentially. In this approach, an alternative clustering is found based on a given clustering. The alternative clustering is made dissimilar to the given one with conditional information bottleneck [30], by "cannot-links" constraints [31], by orthogonal projection [32], or with an optimization problem constrained by the similarity between clusterings [33]. The second approach finds multiple clusterings simultaneously. It includes a method based on k-means [34] and one based on spectral clustering [35]. Both methods find dissimilar clusterings by adding penalty terms that discourage the similarity between the clusterings to their corresponding objective functions. Besides, there is another method that finds multiple clusterings simultaneously using the suboptimal solutions of spectral clustering [36].

There are some other lines of work that produce multiple clusterings. Caruana et al. [37] generate numerous clusterings by applying random weights on the attributes. The clusterings are presented in an organized way so that a user can pick the one he deems the best. Biclustering [38] attempts to find groups of objects that exhibit the same pattern (e.g., synchronous rise and fall) over a subset set of attributes. Unlike our work, the above related work is distance-based rather than model-based.

Our work is also related to projected clustering and subspace clustering [39]. Projected clustering can be considered as performing local variable selection, where each cluster may have a different subset of attributes. It produces only a single clustering as variable selection methods do. Subspace clustering considers dense regions as clusters and tries to identify all clusters in all subspaces. However, it does not partition data along those identified subspaces. In other words, some data points may not be assigned to any cluster in an

identified subspace. This is different from the facet determination approach, in which data are partitioned along every identified facet.

In terms of model definition, the AutoClass models [40] and the MULTIMIX models [41] are some mixture models that are related to our work. The manifest variables in those models have multivariate Gaussian distributions and are similar to the pouch nodes in PLTMs. However, those models do not allow multiple latent variables.

10. Concluding Remarks

In this paper, we have proposed PLTMs as a generalization of GMMs and empirically compared PLTM analysis with several GMM-based methods. Real-world high-dimensional data are usually multifaceted and interesting clusterings in such data often are relevant only to subsets of attributes. One way to identify such clusterings is to perform variable selection. Another way is to perform facet determination by PLTM analysis. Our work has shown that PLTM analysis is often more effective than the first approach.

One drawback of PLTM analysis is that the training is slow. For example, one run of PLTM analysis took around 5 hours on data sets of moderate size (e.g., image, ionosphere, and wdbc data) and around 2.5 days on the largest data set (zernike data) in our experiments. This prohibits the use of PLTM analysis on data with very high dimensions. For example, PLTM analysis is currently infeasible for data with hundreds or thousands of attributes, such as those for text mining and gene expression analysis. On the other hand, even though NBA data has only 18 attributes, our experiment demonstrated that PLTM analysis on that data could still identify multiple meaningful facets. This suggests that PLTM analysis can be useful for data sets that have tens of attributes. One possibly fruitful application of PLTMs is for analyzing survey data. Those data usually have only tens of attributes and are likely to be multifaceted.

A future direction of this work is to speed up the learning of PLTMs, so that PLTM analysis can be applied in more areas. This can possibly be done by parallelization. Recently, some structural learning methods for LTMs have been proposed to follow a variable clustering approach [13, 42, 43] or a constraint-based approach [44, 45]. The learning of PLTMs can also possibly be speed up by following those approaches.

We have made available online the data sets and the Java implementation of the algorithm used in the experiments. They can be found on http://www.cse.ust.hk/faculty/lzhang/ltm/index.htm.

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Appendix A. Inference Algorithm

Consider a PLTM M with manifest variables X and latent variables Y. Recall that inference on M refers to the computation of the posterior probability P(q|e) of some variables of interest, $Q \subseteq X \cup Y$, after observing values e of evidence variables $E \subseteq X$. To perform inference, M has to be converted into a clique tree T. A propagation scheme for message passing can then be carried out on T.

Construction of clique trees is simple due to the tree structure of PLTMs. To construct \mathcal{T} , a clique C is added to \mathcal{T} for each edge in M, such that $C = \mathbf{V} \cup \{\Pi(\mathbf{V})\}$ contains the variable(s) \mathbf{V} of the child node and variable $\Pi(\mathbf{V})$ of its parent node. Two cliques are then connected in \mathcal{T} if they share any common variable. The resulting clique tree contains two types of cliques. The first type are discrete cliques. Each one contains two discrete variables. The second type are mixed cliques. Each one contains the continuous variables of a pouch node and the discrete variable of its parent node. Observe that in a PLTM all the internal nodes are discrete and only leaf nodes are continuous. Consequently, the clique tree can be considered as a clique tree consisted of all discrete cliques, with some mixed cliques attaching to it on the boundary.

After a clique tree is constructed, propagation can be carried out on it. Algorithm 1 outlines a clique tree propagation, based on the Hugin architecture [46, 47], for PLTMs. It consists of four main steps:

Algorithm 1 Inference Algorithm

```
1: procedure Propagate(M, \mathcal{T}, \boldsymbol{E}, \boldsymbol{e})
                                                                   19: procedure DISTRIBUTEMESSAGE(C, C')
        Initialize \psi(C) for every clique C
                                                                   20:
                                                                           SendMessage(C, C')
                                                                           for all C'' \in Ne(C') \setminus \{C\} do
 3:
        Incorporate evidence to the potentials
                                                                   21:
                                                                               DistributeMessage(C', C'')
 4:
        Choose an arbitrary clique in \mathcal{T} as C_P
                                                                   22:
 5:
        for all C \in Ne(C_P) do
                                                                   23:
                                                                           end for
           CollectMessage(C_P, C)
                                                                   24: end procedure
 6:
 7:
        end for
                                                                   25: procedure SENDMESSAGE(C, C')
        for all C \in Ne(C_P) do
 8:
                                                                           \phi \leftarrow \text{RetrieveFactor}(C \cap C')
 9:
           DISTRIBUTEMESSAGE(C_P, C)
                                                                   26:
                                                                           \phi' \leftarrow \sum_{C \setminus C'} \psi(C)
10:
                                                                   27:
                                                                           SaveFactor(C \cap C', \phi')
        Normalize \psi(C) for every clique C
                                                                   28:
12: end procedure
                                                                           \psi(C') \leftarrow \psi(C') \times \phi'/\phi
                                                                   29:
                                                                   30: end procedure
13: procedure CollectMessage(C, C')
14:
        for all C'' \in Ne(C') \setminus \{C\} do
                                                                   31: procedure RetrieveFactor(S)
           CollectMessage(C', C'')
15:
                                                                           If SAVEFACTOR(S, \phi) has been called, return \phi;
16:
        end for
                                                                        otherwise, return 1.
17:
        SendMessage(C', C)
                                                                   33: end procedure
18: end procedure
                                                                        // Ne(C) denotes the neighbors of C
```

initialization of cliques, incorporation of evidence, message passing, and normalization. The propagation on the discrete part of the clique tree is done as usual. Here we focus on the part related to the mixed cliques.

Step 1: Initialization of cliques (line 2). Consider a mixed clique C_m containing continuous variables W and discrete variable $Y = \Pi(W)$. The potential $\psi(w, y)$ of C_m (also denoted as $\psi(C_m)$ in the algorithm) is initialized by the corresponding conditional distribution,

$$\psi(\boldsymbol{w},y) = P(\boldsymbol{w}|y) = \mathcal{N}(\boldsymbol{w}|\boldsymbol{\mu}_y, \boldsymbol{\Sigma}_y).$$

Step 2: Incorporation of evidence (line 3). The variables in a pouch node W can be divided into two groups, depending on whether the values of the variables have been observed. Let $E' = W \cap E$ denote those variables of which values have been observed, and let $U = W \setminus E$ denote those of which values have not been observed. Furthermore, let $[\mu]_S$ denote the part of mean vector μ containing elements corresponding to variables S, and let $[\Sigma]_{ST}$ denote the part of the covariance matrix Σ that has rows and columns corresponding to variables S and T, respectively. To incorporate the evidence e' for E', the potential of C_m changes from $\psi(w, y)$ to

$$\psi'(\boldsymbol{w}, y) = P(\boldsymbol{e}'|y) \times P(\boldsymbol{w}|y, \boldsymbol{e}') = \mathcal{N}\left(\boldsymbol{e}' | \begin{bmatrix} \boldsymbol{\mu}_y \end{bmatrix}_{\boldsymbol{E}'}, \begin{bmatrix} \boldsymbol{\Sigma}_y \end{bmatrix}_{\boldsymbol{E}'\boldsymbol{E}'}\right) \times \mathcal{N}\left(\boldsymbol{w}|\boldsymbol{\mu}_y', \boldsymbol{\Sigma}_y'\right),$$

where μ'_y and Σ'_y can be divided into two parts. The part related to the evidence variables E' is given by:

$$egin{bmatrix} m{\mu}_y' m{]}_{m{E}'} = m{e}', & m{\left[m{\Sigma}_y'
ight]}_{m{U}m{E}'} = m{0}, & m{\left[m{\Sigma}_y'
ight]}_{m{E}'m{E}'} = m{0}, & ext{and} & m{\left[m{\Sigma}_y'
ight]}_{m{E}'m{U}} = m{0}. \end{split}$$

The other part is given by:

$$\begin{split} \left[\boldsymbol{\mu}_y'\right]_{\boldsymbol{U}} &= \left[\boldsymbol{\mu}_y\right]_{\boldsymbol{U}} + \left[\boldsymbol{\Sigma}_y\right]_{\boldsymbol{U}\boldsymbol{E}'} \left[\boldsymbol{\Sigma}_y^{-1}\right]_{\boldsymbol{E}'\boldsymbol{E}'} \left(\boldsymbol{e}' - \left[\boldsymbol{\mu}_y\right]_{\boldsymbol{E}'}\right), \\ \left[\boldsymbol{\Sigma}_y'\right]_{\boldsymbol{U}\boldsymbol{U}} &= \left[\boldsymbol{\Sigma}_y\right]_{\boldsymbol{U}\boldsymbol{U}} - \left[\boldsymbol{\Sigma}_y\right]_{\boldsymbol{U}\boldsymbol{E}'} \left[\boldsymbol{\Sigma}_y^{-1}\right]_{\boldsymbol{E}'\boldsymbol{E}'} \left[\boldsymbol{\Sigma}_y\right]_{\boldsymbol{E}'\boldsymbol{U}}. \end{split}$$

Step 3: Message passing (lines 5–10). In this step, C_m involves two operations, marginalization and combination. Marginalization of $\psi'(\boldsymbol{w}, y)$ over \boldsymbol{W} is required for sending out a message from C_m (line 27). It results in a potential $\psi'(y)$, involving only the discrete variable y, as given by:

$$\psi'(y) = P(\boldsymbol{e}'|y) = \mathcal{N}\left(\boldsymbol{e}' \middle| \left[\boldsymbol{\mu}_y\right]_{\boldsymbol{E}'}, \left[\boldsymbol{\Sigma}_y\right]_{\boldsymbol{E}'\boldsymbol{E}'}\right).$$

Combination is required for sending a message to C_m (line 29). The combination of the potential $\psi'(\boldsymbol{w}, y)$ with a discrete potential $\phi(y)$ is given by $\psi''(\boldsymbol{w}, y) = \psi'(\boldsymbol{w}, y) \times \phi(y)$. When the message passing completes (line 10), $\psi''(\boldsymbol{w}, y)$ represents the distribution

$$\psi''(\boldsymbol{w}, y) = P(y, \boldsymbol{e}) \times P(\boldsymbol{w}|y, \boldsymbol{e}') = P(y, \boldsymbol{e}) \times \mathcal{N}\left(\boldsymbol{w}|\boldsymbol{\mu}_y', \boldsymbol{\Sigma}_y'\right).$$

Step 4: Normalization (line 11). In this step, a potential changes from $\psi''(w,y)$ to

$$\psi'''(\boldsymbol{w}, y) = P(y|\boldsymbol{e}) \times P(\boldsymbol{w}|y, \boldsymbol{e}') = P(y|\boldsymbol{e}) \times \mathcal{N}\left(\boldsymbol{w}|\boldsymbol{\mu}_y', \boldsymbol{\Sigma}_y'\right).$$

For implementation, the potential of a mixed clique is usually represented by two types of data structures: one for the discrete distribution and one for the conditional Gaussian distribution. More details for the general clique tree propagation can be found in [46, 21, 48].

Appendix B. Search Algorithm

Some issues for the search algorithm of latent tree models are studied in [49, 14]. These issues also affect PLTMs. In this section, we discuss how these issues are addressed for PTLMs. We summarize the entire search algorithm for PLTMs at the end of this section.

Appendix B.1. Three Search Phases

In every search step, a search operator generates all possible candidate models for consideration. Let l and b be the numbers of latent nodes and pouch nodes, respectively, and n=l+b be total number of nodes. Let p, q, and r be the maximum number of variables in a pouch node, the maximum number of sibling pouch nodes, and the maximum number of neighbors of a latent node, respectively. The numbers of possible candidate models that NI, ND, SI, SD, NR, PO, and UP can generate are O(lr(r-1)/2), O(lr), O(l), O(ll), O(ll), O(lq(q-1)/2), and O(bp), respectively. If we consider all seven operators in each search step, many candidate models are generated but only one of them is chosen for the next step. Those suboptimal models are discarded and are not used for the next step. Therefore, in some sense, much time are wasted for considering the suboptimal models.

A more efficient way is to consider fewer candidate models in each search step. This can be achieved by considering only a subset of search operators at a time. The search operators can be naturally partitioned into three categories. SI, NI, and PO belong to the expansion category, since all of them can create candidate models that are more complex than the current one. SD, ND, and UP, which simplify a model, belong to the simplification category. NR does not change the complexity considerably. It belongs to the adjustment category. We therefore perform search in three phases, each of which considers only operators in one category. The best model found in each phase is used to seed the next phase, and the search repeats these three phases until all three phases cannot find a better model. This search is called EAST, since it does Expansion, Adjustment, and Simplification until Termination.

Appendix B.2. Efficient Model Evaluation

Suppose m is the current model after a number of search steps. A candidate model m' can be obtained from m by applying a search operator. Very often, the search operator modifies only a small part of m, so that m and m' share a large number of parameters. For example, consider the model m_2 , that is obtained from model m_1 using the PO operator, as shown in Figure 3. m_1 and m_2 shares many parameters such as $P(x_1, x_2|y_2)$, $P(y_2|y_1)$, $P(y_3|y_1)$, $P(x_6|y_3)$, etc. On the other hand, some parameters are not shared by m

and m'. In this example, parameters $P(x_4|y_3)$ and $P(x_5|y_3)$ are specific to m_1 , while parameter $P(x_4, x_5|y_3)$ is specific to m_2 . We can divide the parameters of m' into two groups and write the parameters $\boldsymbol{\theta}'$ of m' as $(\boldsymbol{\theta}'_1, \boldsymbol{\theta}'_2)$. $\boldsymbol{\theta}'_1$ consists of parameters shared with m, while $\boldsymbol{\theta}'_2$ consists of parameters specific to m'. Similarly, we write the parameters of m as $\boldsymbol{\theta} = (\boldsymbol{\theta}_1, \boldsymbol{\theta}_2)$.

Suppose we have computed the MLE $\boldsymbol{\theta}^* = (\boldsymbol{\theta}_1^*, \boldsymbol{\theta}_2^*)$ of the parameters of m. $\boldsymbol{\theta}_1^*$ can be used as estimates for the shared parameters of m'. Consequently, we can obtain a likelihood function $P(\mathcal{D}|m', \boldsymbol{\theta}_1^*, \boldsymbol{\theta}_2')$ that depends on only the unshared parameters $\boldsymbol{\theta}_2'$. This function is referred to as restricted likelihood function of m'. As previously described, we use the BIC score to evaluate a candidate model. This requires the maximum log-likelihood (ML) of m'. Instead of computing the ML over all parameters of m', we may approximate it by maximizing the restricted log-likelihood over only the subset of parameters $\boldsymbol{\theta}_2'$. This results in an approximate score given by

$$BIC_{RL}(m'|\mathcal{D}) = \max_{\boldsymbol{\theta}_2'} \log P(\mathcal{D}|m', \boldsymbol{\theta}_1^*, \boldsymbol{\theta}_2') - \frac{d(m')}{2} \log N.$$

The advantage of using BIC_{RL} is that it allows a more efficient implementation of EM. This is due to two reasons. First, it involves a maximization of fewer parameters in the M-step. This also means fewer computations in the E-step, since only distributions relevant to θ'_2 need to be computed. Second, the E-step can exploit the fixed parameters θ^*_1 to allow sharing of computation among all iterations of EM. Specifically, the E-step requires inference on a PLTM, which in turn requires passing messages in the clique tree. As the parameters θ^*_1 do not change in these iterations, the messages that depend on only these parameters remain the same in all iterations. Therefore, some messages can be cached for each data case and can be shared for the inference used in subsequent E-steps.

In our implementation, both BIC and BIC_{RL} are used to evaluate models, but at different situations. The approximation BIC_{RL} is used for quickly evaluating candidate models generated by the same search operator, while the real BIC is used for accurately evaluating the candidate model with the highest BIC_{RL} for each of the search operators. This improves the speed for the evaluation of models within search operators, but maintains the accuracy for the evaluation of models across search operators.

Appendix B.3. Operation Granularity

Operation granularity refers to the phenomenon where some search operators might increase the complexity of the current model much more than other search operators. If we use BIC to evaluate candidate models given by the search operators, those having a much larger increase in complexity are usually preferred. This might lead to local maxima. Therefore, we adopt a simple and effective strategy called the cost-effectiveness principle for evaluating the candidate models. Let m be the current model and m' be a candidate model. Define the improvement ratio of m' over m given data \mathcal{D} to be

$$IR(m', m|\mathcal{D}) = \frac{BIC(m'|\mathcal{D}) - BIC(m|\mathcal{D})}{d(m') - d(m)}.$$

It is the increase in model score per unit increase in model complexity. The cost-effectiveness principle stipulates that one chooses, among a list of candidate models, the one with the highest improvement ratio.

The principle is applied only on candidate models generated by the SI, NI, and PO operators. In other words, it is used only during the expansion phase. The principle is not applied for other operators since they do not or do not necessarily increase model complexity.

Appendix B.4. PLTM-EAST

The entire search algorithm for PLTMs is named PLTM-EAST. It is outlined in Algorithm 2. The search process starts with the initial model $m^{(0)}$ described in Section 6. The procedure PLTM-EAST uses $m^{(0)}$ as the first current model. It then repeatedly tries to improve the current model in the three different phases.

In procedure EXPAND, the improvement ratio IR is used to pick the best models among the candidate models generated by SI, NI, and PO. It stops when the best candidate model fails to improve over the

Algorithm 2 Search Algorithm

```
1: procedure PLTM-EAST(m, \mathcal{D})
                                                                                23: procedure Adjust(m, \mathcal{D})
 2:
         repeat
                                                                                          return RepPickModel(m, NR, \mathcal{D})
3:
              m' \leftarrow m
                                                                                25: end procedure
 4:
              m \leftarrow \text{Expand}(m)
 5:
              m \leftarrow \text{Adjust}(m)
                                                                                26: procedure SIMPLIFY(m, \mathcal{D})
              m \leftarrow \text{Simplify}(m)
                                                                                          m \leftarrow \text{RepPickModel}(m, UP, \mathcal{D})
 6:
                                                                                27:
         until BIC(m|\mathcal{D}) \leq BIC(m'|\mathcal{D})
 7:
                                                                                28:
                                                                                          m \leftarrow \text{RepPickModel}(m, ND, \mathcal{D})
         return m'
                                                                                          m \leftarrow \text{RepPickModel}(m, SD, \mathcal{D})
8:
                                                                                29:
9: end procedure
                                                                                30: end procedure
10: procedure Expand(m, \mathcal{D})
                                                                                31: procedure RepPickModel(m, Op, \mathcal{D})
         loop
11:
                                                                                32:
                                                                                          repeat
12:
              m' \leftarrow m
                                                                                33:
13:
              \mathcal{M} \leftarrow SI(m') \cup NI(m') \cup PO(m')
                                                                                34:
                                                                                              m \leftarrow \text{PickModel}(Op(m'), \mathcal{D})
14:
              m \leftarrow \text{PickModel-IR}(\mathcal{M}, \mathcal{D})
                                                                                35:
                                                                                          until BIC(m|\mathcal{D}) \leq BIC(m'|\mathcal{D})
15:
              if BIC(m|\mathcal{D}) \leq BIC(m'|\mathcal{D}) then
                                                                                36:
                                                                                          return m'
16:
                  return m'
                                                                                37: end procedure
              end if
17:
18:
              if m \in NI(m') \cup PO(m') then
19:
                  m \leftarrow \text{Enhance}(m, \mathcal{D})
20:
              end if
21:
         end loop
22: end procedure
```

previous model. On the other hand, if the best candidate model is better, and if it comes from NI or PO, procedure Enhance is called. This procedure iteratively improves the model by using a restricted version of search operator. If the given model is generated by NI, restricted version of NR is used. If it is generated by PO, restricted version of PO is used.

In procedure Adjust, it calls the procedure RepPickModel to improve the model using the NR operator repeatedly. In procedure Simplify, it first tries to improve the model using UP. It then uses ND, and uses SD lastly.

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