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A Confident Information First Principle for Parameter Reduction and Model Selection of Boltzmann Machines

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Abstract—Typical dimensionality reduction (DR) methods are data-oriented, focusing on directly reducing the number of random variables (or features) while retaining the maximal variations in the high-dimensional data. Targeting unsupervised situations, this paper aims to address the problem from a novel perspective and considers model-oriented DR in parameter spaces of binary multivariate distributions. Specifically, we propose a general parameter reduction criterion, called confident-information-first (CIF) principle, to maximally preserve confident parameters and rule out less confident ones. Formally, the confidence of each parameter can be assessed by its contribution to the expected Fisher information distance within a geometric manifold over the neighborhood of the underlying real distribution. Then, we demonstrate two implementations of CIF in different scenarios. First, when there are no observed samples, we revisit the Boltzmann machines (BMs) from a model selection perspective and theoretically show that both the fully visible BM and the BM with hidden units can be derived from the general binary multivariate distribution using the CIF principle. This finding would help us uncover and formalize the essential parts of the target density that BM aims to capture and the nonessential parts that BM should discard. Second, when there exist observed samples, we apply CIF to the model selection for BM, which is in turn made adaptive to the observed samples. The sample-specific CIF is a heuristic method to decide the priority order of parameters, which can improve the search efficiency without degrading the quality of model selection results as shown in a series of density estimation experiments.

Index Terms—Boltzmann machine (BM), Fisher information, information geometry (IG), parametric reduction.

I. INTRODUCTION

RECENTLY, deep learning models (e.g., deep belief networks (DBNs) [1], stacked denoising autoencoder [2], and deep Boltzmann machine (DBM) [3]) have drawn increasing attention due to their impressive empirical performance in various application areas, such as computer vision [4]–[6], natural language processing [7], information retrieval [8], [9], and other classification problems [10]–[12]. Despite of these practical successes, there have been debates on the fundamental principle that governs the design and training of those deep architectures. In most situations, searching the parameter space for deep learning models is difficult. To tackle this difficulty, unsupervised pretraining has been introduced as an important process. In [13], it has been empirically shown that the unsupervised pretraining could fit the network parameters in a region of the parameter space that could well capture the data distribution, thus alleviating generalization error of the trained deep architectures.

From the density estimation point of view, the unsupervised learning can be interpreted as an attempt to discover a set of parameters for a generative model that describes the underlying distribution of the observed data. In real-world applications, the data sets are often high-dimensional and we would need a model with high-dimensional parameter space in order to effectively depict the underlying distribution. However, when the model becomes excessively complex, the model would easily overfit the limited training data. This leads to the issue of model selection, i.e., selecting a subset of parameters in an attempt to create a model of optimal complexity for the given data (a comprehensive review of model selection can be found in [14] and [15]).

Targeting unsupervised situations, this paper aims to address this issue by considering model-oriented dimensionality reduction (DR) in parameter spaces of binary multivariate distributions. Since BMs are fundamental building blocks for a number of widely used deep architectures (e.g., DBN and DBM), we will focus on a formal analysis of the essential parts of the target density (a multivariate binary distribution) that the BM aims to capture in terms of model selection.

Here, finding the essential parts for the underlying distribution is the main objective of model selection, which has two implications: 1) the choice of the number of free parameters and 2) the choice of parameters given that number. The former has been studied by classical model selection approaches
(such as Akaike information criterion (AIC) [16] and Bayesian information criterion (BIC) [17]), where the number of free parameters is adopted as a model complexity measure. However, the model complexity is a complicated quantity which should take into account the effectiveness of the selected free parameters (e.g., as illustrated in Section 3.5.3 of Bishop’s book [18] and the curvature measure [19]). Therefore, the latter problem (i.e., choice of parameters given a model complexity) is also of great significance for the model selection purpose and should be thoroughly studied.

Various model selection procedures have been proposed in the literature. In general, the model selection problem is tackled by the combination of a model selection criterion that allows comparison of alternative models and a search strategy that allows us to find the optimal model according to that criterion. Typical model selection criteria include $P$-value [20] that is based on the hypothesis test of the likelihood ratio [21] between two nested models; the information-theoretic approaches that estimate the expected information distance based on the empirical log-likelihood function when a certain model is used [14]. For the latter, the expected information distance can be estimated by different techniques, such as bootstrap methods using bootstrap samples [22], cross-validation methods that average the log likelihoods on validation samples [23], and asymptotic methods aiming to find an (asymptotically) unbiased estimator for the expected information distance that takes into account the log likelihood of the data under a candidate model and a penalty related to the effective number of parameters in the model (the well-known examples are the AIC that penalizes the number of parameters at the rate of $O(1)$ and BIC that penalizes at the rate of $O(\log N)$, $N$ is the number of samples). In addition to the above model selection criteria, the $L_1$ regularization methods can also be used for the model selection purpose by jointly minimizing the empirical error (or negative log likelihood) and penalty [24], [25], where the parameters with significant absolute values after training are selected to create the optimal model.

As described earlier, the model selection problem in high-dimensional parameter space can be divided into the choice of the number of free parameters and the choice of parameters given that number. The classical model selection criteria can be seen as the integration of the two, formalized by a single optimization objective. However, the existing model selection approaches are insufficient when the model’s parameters do not have a priority order, which is the case for BM.

First, the lack of an explicit priority order of parameters would lead to serious efficiency issues. Assuming there exists a universal parametric probabilistic model $S$ (with $K$ free parameters) that is general enough to represent all system phenomena. Consequently, there exist $2^K$ candidate submodels, corresponding to different choices of parameters. When $K$ is large, it is computationally intractable to exhaustively test all submodels in classical model selection methods. By using a stepwise greedy search strategy [26], the number of tested submodels can be reduced to $O(K^2)$, which is often time-consuming in practice. Although $L_1$ regularization could avoid the heavy combinatorial search, the regularization coefficient needs to be determined through cross validation, which may also be time-consuming in practice.

Second, for BM, the lack of an explicit priority order of parameters would lead to a comparison among nonnested models. For $P$-value, since the likelihood ratio tests exist only for nested models, the tests of hypotheses within a data set are not independent (called multiple testing problem). As a result, it is difficult to make inferences [26]. For information criteria, there exist considerable controversies about their applicability for nonnested models. Burnham and Anderson [26] argued that the information criteria make no assumptions about nested candidate models and the order in which the information criterion is computed over the set of models is not relevant, while Ripley [27] stated that the differences in AIC for pairs of nested models can be much more precisely estimated than those for some nonnested pairs, and the sampling error can make comparisons of AIC meaningless unless the differences are large.

In summary, although the classical methods have demonstrated a sound theoretical basis for choosing suitable model complexity, they suffer from limitations in parameter choice when the model’s parameters do not have a priority order, especially for models involving a large number of parameters.

In this paper, we focus on addressing the parameter choice problem, for density estimation on the parameter space of multivariate distributions under a given number of free parameters. Note that the parameter reduction is related but different from the traditional DR. Traditional DR techniques are often data-oriented and focus on directly reducing the number of random variables (features) while retaining the maximal variations in the high-dimensional data, e.g., feature selection [28], [29] and feature extraction [30], [31]. Generally speaking, they cannot be applied to the parametric reduction problem that this paper aims to tackle.

We propose a general parameter reduction criterion, namely the confident-information-first (CIF) principle, to maximally preserve confident parameters and rule out less confident ones. Formally, the confidence of each parameter can be assessed by its contribution to the expected Fisher information distance (FID) within a geometric manifold over the neighborhood of the underlying real distribution, based on the theoretical framework of information geometry (IG) [32].

A. Motivation of CIF Under IG

In IG, the general model $S$ can be seen as a $K$-dimensional manifold and the goal of the parametric reduction is to derive a lower dimensional submodel $M$ by reducing the number of free parameters in $S$. $M$ is a smoothed submanifold of $S$. The number of free parameters in $M$ is restricted to be a constant $k$ ($k \ll K$). A more detailed introduction to IG is given in Section II.

The major difficulty in the parametric reduction procedure is the choice of parameters to keep or to remove. In this paper,
with parameters $\zeta_1$ and $\zeta_2$, the FID between $p_1$ and $p_2$ is

$$D(p_1, p_2) = \sqrt{(\zeta_1 - \zeta_2)^T G_{\zeta}(\zeta_1 - \zeta_2)}$$  \hspace{1cm} (2)

where $G_\zeta$ represents the Fisher information matrix [32].

The rationality of maximally preserving the FID can also be interpreted from the maximum-likelihood (ML) estimation’s point of view. Let $\hat{\zeta}$ be the ML estimators for $\zeta$. The asymptotic normality of ML estimation implies that the density of $\hat{\zeta}$ is the normal distribution with mean $\zeta$ and covariance $\Sigma$, that is

$$f(\hat{\zeta}) \sim \mathcal{N}(\zeta, \Sigma) = \frac{1}{Z} \exp \left\{ - \frac{1}{2} (\zeta - \hat{\zeta})^T \Sigma^{-1} (\zeta - \hat{\zeta}) \right\}$$  \hspace{1cm} (3)

where the inverse of $\Sigma$ can be asymptotically estimated using the Fisher information matrix $G_\zeta$, as suggested by the Cramér–Rao bound [36]. From the FID given in 2, the exponent part of 3 is just the opposite of the half squared FID between two distributions $p$ and $\hat{p}$ determined by the close parameters $\zeta$ and $\hat{\zeta}$, respectively. Hence, a larger FID means a lower likelihood. It turns out that, in density estimation, maximally preserving the expected FID after the projection $\Gamma_M$ (1) is equivalent to maximally preserving the likelihood structure among close distributions. In supervised learning tasks (e.g., classification), maximally preserving FID can also effectively preserve the likelihood structure among different class densities (the generative distributions of classes), which helps prevent the interference of sample noises. Recall that sample noises always reduce the FID among class densities in a statistical sense, leading to a reduced discrimination marginality between two class densities. Hence, for noisy data, the model that maximally preserving FID will have a better discriminative capability.

To solve the optimization problem in 1, we propose a parameter reduction criterion called the CIF principle, described as follows. For multivariate binary distributions, the squared FID $D^2(p_1, p_2)$ can be decomposed into the distances of two orthogonal parts by using the dually orthogonal coordinates in IG [32]

$$D^2(p_1, p_2) = D^2(\Gamma_M(p_1), \Gamma_M(p_2)) + D^2(\Gamma_M(p_1), \Gamma_M(p_2))$$

Here, the submanifold $\overline{M}$ is complementary to $M$, i.e., the free parameters in $\overline{M}$ are nonfree in $M$, and vice versa. Thus, it is possible to categorize the system parameters of $S$ into two sets, i.e., the set of parameters in $M$ with “major” variations and the set of parameters in $\overline{M}$ with “minor” variations, based on their contributions to the whole information distance. The former refers to parameters that are important for reliably distinguishing the true distribution from the sampling distribution, thus considered as “confident.” Meanwhile, the parameters in $\overline{M}$ with minor contributions can be regarded as less reliable. Hence, the CIF principle can be stated as parametric reduction that preserves the confident parameters and rules out less confident parameters.

B. Outline and Contributions

The CIF principle is a general parametric choice framework that can be used to determine the optimal choice of submodel’s free parameters. There exist different implementations of CIF
with respect to different underlying distributions and/or their coordinate representations. The implementation of CIF can be analogized to some general inference objectives, e.g., the ML, which can achieve solutions in different senses (e.g., globally optimal or locally optimal) with respect to different application contexts. In this paper, we demonstrate two kinds of implementation of the CIF principle in two different application scenarios.

First, when \( p_k \) is unknown, i.e., there is no observed information, the CIF principle could be used to derive an optimal probabilistic model in an expectation sense. If the l-mixed \( \zeta \)-coordinates \([\zeta]\) (8) is adopted, we theoretically show that CIF could lead to an optimal submanifold \( M \) for \( k = \sum_{l=1}^{L} C^l_n \) in terms of the optimization problem in 1 (see Section III). For \( l = 2 \), we further show that the optimal submanifold \( M \) derived by CIF is exactly the BM (see Section IV). Note that CIF may be used to derive other useful models if different coordinate systems are employed.

Second, when \( p_k \) is known, the CIF principle is applied in the model selection for BM as a heuristic method to decide the priority order of parameters, which can improve the search efficiency without degrading the quality of model selection results. For a given number of free parameters \( k \), we can directly deduce a set of preferred parameters from all \( k \)-dimensional submodels by selecting the top-\( k \) confident parameters, where the confidences can be precalculated for efficiency. Then, by integrating the information criteria (such as AIC) with CIF, we can alleviate the possible bias of the model complexity when AIC is used to compare nonnested models and improve the computational efficiency of the model selection procedure. In this case, we only need to calculate the AIC for \( K \) submodels to obtain a solution. Moreover, we have further developed separate hypothesis tests on the confidences of parameters to decide whether certain parameters should be preserved or not (see Section V-A), where the significance level \( \alpha \) used in a hypothesis test can be considered as a heuristic threshold for the confidences of parameters.

II. THEORETICAL FOUNDATIONS OF IG

In this section, we introduce and develop the theoretical foundations of IG [32] for the manifold \( S \) of binary multivariate distributions with a given number of variables \( n \), i.e., the open simplex of all probability distributions over binary vector \( x \in \{0, 1\}^n \). This will lay the foundation for our theoretical deviation of CIF.

A. Notations for Manifold \( S \)

In IG, a parametric family of probability distributions is regarded as a differentiable manifold equipped with certain coordinate systems. In the case of binary multivariate distributions, there exist four commonly used coordinate systems [32, 37]: \( p \)-coordinates, \( \eta \)-coordinates, \( \theta \)-coordinates, and the mixed \( \zeta \)-coordinates. The \( \zeta \)-coordinates is very important for the analysis in this paper.

For the \( p \)-coordinates \([p]\), the probability distribution over \( n \) binary variables can be completely specified by a vector of \( 2^n - 1 \) positive numbers, corresponding to the probability of any \( 2^n - 1 \) exclusive states of \( x \). For example, when \( n = 2 \), the \( p \)-coordinates could be \([p] = (p_{01}, p_{00}, p_{11})\). Note that all probability terms need to be strictly positive as required by IG [32]. For simplicity, the \( p \)-coordinates are indexed by capital letters \( I, J, \ldots \), where an index \( I \) denotes a subset of \( \{1, 2, \ldots, n\} \). \( p_I \) is defined as the probability that all variables included in \( I \) are one and the complemented variables equal to zero. For example, if \( I = \{1, 3\} \) and \( n = 3 \), we have

\[
p_{I} = p_{101} = \text{Prob}(x_1 = 1, x_2 = 0, x_3 = 1).
\]

Note that the null set \( I = \{} \) is also valid for the \([p]\) and \( p_{00..0} \) denotes the probability that all variables are zero.

The \( \eta \)-coordinates \([\eta]\) are defined by

\[
\eta_I = E[X_I] = \text{Prob} \left( \prod_{i \in I} x_i = 1 \right) \tag{4}
\]

where \( X_I = \prod_{i \in I} x_i \) and \( E[\cdot] \) denotes the expectation with respect to the probability distribution over \( x \). The \( \eta \)-coordinates can be grouped by coordinate orders and denoted as \([\eta] = (\eta_1^1, \eta_2^2, \ldots, \eta_n^n)\), where the superscript represents the order number of the corresponding parameter. For example, \( \eta_1^1 \) denotes the set of all \( \eta \) parameters with the order number two.

The \( \theta \)-coordinates (natural coordinates) \([\theta]\) are defined by

\[
\log p(x) = \sum_{I \subseteq \{1, 2, \ldots, n\}, I \not= \text{NullSet}} \theta^I X_I - \psi(\theta) \tag{5}
\]

where \( \psi(\theta) = \log(\sum_x \exp(\sum_i \theta_i X_i(x))) \) is the cumulant generating function and its value equals to \(-\log \text{Prob}(x_i = 0, \forall i \in \{1, 2, \ldots, n\})\). By solving the linear system 5, we have \( \theta^I = \frac{1}{K-I} \sum_{K \subseteq I} (-1)^{I-K} \log(p_K) \). The \( \theta \)-coordinate is denoted as \([\theta] = (\theta_1^1, \theta_2^2, \ldots, \theta_n^n)\), where the subscript represents the order number of the corresponding parameter. Note that the order indices locate differently in \([\eta]\) and \([\theta]\) by convention in [33].

There exists bijective map between coordinate systems \([\eta]\) and \([\theta]\), formally given by the Legendre transformation

\[
\theta^I = \frac{\partial \phi(\eta)}{\partial \eta_I}, \quad \eta_I = \frac{\partial \psi(\theta)}{\partial \theta^I} \tag{6}
\]

where \( \psi(\theta) \) is specified in 5 and \( \phi(\eta) = \sum_x p(x; \eta) \log p(x; \eta) \) denotes the negative entropy. It can be shown that \( \psi(\theta) \) and \( \phi(\eta) \) satisfy the following equation [32]:

\[
\psi(\theta) + \phi(\eta) - \sum \theta^I \eta_I = 0. \tag{7}
\]

The l-mixed \( \zeta \)-coordinates \([\zeta]\) are defined by

\[
[\zeta] = [\eta^l, \theta_{l+}] = (\eta_1^1, \eta_2^2, \ldots, \eta_{l,j \ldots}^l, \theta_{l+}^j, \ldots, \theta_{l+}^{1,\ldots,n}) \tag{8}
\]

where \( l \in \{1, \ldots, n - 1\} \). The former part \( \eta^l \) consists of \( \eta \)-coordinates with order less or equal to \( l \) and the latter part \( \theta_{l+} \) consists of \( \theta \)-coordinates with order greater than \( l \).
B. Fisher Information Matrix for Parametric Coordinates

For a general coordinate system \( [\zeta] \), the \( i \)th row and \( j \)th column element of the Fisher information matrix \( G_{\zeta} \) for \([\zeta]\) is defined as the covariance of the scores of \( \zeta_i \) and \( \zeta_j \) [36]

\[
g_{ij} = E \left[ \frac{\partial \log p(x; \zeta)}{\partial \zeta_i} \frac{\partial \log p(x; \zeta)}{\partial \zeta_j} \right] \]

where \( E[\cdot] \) denotes the expectation with respect to the probability distribution over \( x \). The Fisher information measures the amount of information in the data that a statistic carries about the unknown parameters [38]. The Fisher information matrix is of vital importance to our analysis, since the inverse of the matrix gives an asymptotically tight lower bound to the covariance matrix of any unbiased estimators for the parameters under consideration [36]. Another important concept related to our analysis is the orthogonality defined by Fisher information. Two coordinate parameters \( \zeta_i \) and \( \zeta_j \) are called orthogonal if and only if their Fisher information vanishes, i.e., \( g_{ij} = 0 \), meaning that their influences on the log-likelihood function are uncorrelated.

According to [32], the Fisher information for \([\theta]\) can be rewritten as \( g_{ij} = (\partial^2 \psi(\theta))/((\partial \theta_i \partial \theta_j)) \), and for \([\eta]\), it is \( g_{ij} = (\partial^2 \phi(\eta))/((\partial \eta_i \partial \eta_j)) \). Let \( G_{\theta} = (g_{ij}) \) be the Fisher information matrix for \([\theta]\) and \( G_{\eta} = (g_{ij}) \) for \([\eta]\). As shown in [32], \( G_{\theta} \) and \( G_{\eta} \) are mutually inverse matrices, i.e., \( \sum_J g_{IJ} g_{JK} = \delta_{IK} \), where \( \delta_{IK} = 1 \) if \( I = K \) and zero if \( I \neq K \). Next, we will develop the two propositions to compute \( G_{\theta} \) and \( G_{\eta} \) generally. Note that Proposition 1 is a generalization of Theorem 2 in [33].

**Proposition 1:** The Fisher information between two parameters \( \theta^I \) and \( \theta^J \) in \([\theta]\) is given by

\[
g_{IJ}(\theta) = \eta_{IJ} - \eta_{IJ}. \quad (9)\]

**Proof:** See Appendix A. \( \blacksquare \)

**Proposition 2:** The Fisher information between two parameters \( \eta^I \) and \( \eta^J \) in \([\eta]\) is given by

\[
g_{IJ}(\eta) = \sum_{K \subseteq I \cap J} (-1)^{|I-K|+|J-K|} \cdot \frac{1}{p_K} \quad (10)
\]

where \( |\cdot| \) denotes the cardinality operator.

**Proof:** See Appendix B. \( \blacksquare \)

We take the probability distribution with three variables for example. Based on 10, the Fisher information between \( \eta^I \) and \( \eta^J \) can be calculated, e.g., \( g_{IJ} = 1/(p_{00}) + 1/(p_{010}) \) if \( I = [1, 2] \) and \( J = [2, 3] \), \( g_{IJ} = -1/(1/p_{00}) + 1/(p_{010}) + 1/(p_{100}) \) if \( I = [1, 2] \) and \( J = [1, 2, 3] \), and so on.

The next proposition shows that the Fisher information matrix \( G_{\zeta} \) for the \([\zeta]\) can be calculated based on \( G_{\eta} \) and \( G_{\theta} \).

**Proposition 3:** The Fisher information matrix \( G_{\zeta} \) of \([\zeta]\) is given by

\[
G_{\zeta} = \begin{pmatrix} A & 0 \\ 0 & B \end{pmatrix} \quad (11)
\]

where \( A = ((G_{\eta}^{-1})_{\eta})^{-1} \), \( B = ((G_{\theta}^{-1})_{\theta})^{-1} \), \( G_{\eta} \) and \( G_{\theta} \) are the Fisher information matrices of \([\eta]\) and \([\theta]\), respectively, \( I_{\eta} \) is the index set of the parameters shared by \([\eta]\) and \([\zeta]\), i.e., \( \{\eta_i, \ldots, \eta_{i,j}, \ldots\} \), and \( J_{\theta} \) is the index set of the parameters shared by \([\theta]\) and \([\zeta]\), i.e., \( \{\theta_{i,j}, \ldots, \theta_{n}^{1, \ldots, n}\} \).

**Proof:** See Appendix C. \( \blacksquare \)

### III. General CIF Principle

The general manifold \( S \) of all probability distributions over binary vector \( x \in \{0, 1\}^n \) could be exactly represented using \( 2^n - 1 \) parametric coordinates. Given a target distribution \( q(x) \in S \), we would like to realize it by a distribution in some lower dimensional submanifold \( M \). This is defined as the parametric reduction problem for multivariate binary distributions.

In this section, we will formally illuminate the general CIF for parametric reduction. Assume we can construct a coordinate system whose parameters entail a natural hierarchy according to their confidences, that high confident parameters are significantly distinguished from and orthogonal to lowly confident ones. Then, we could implement the CIF conveniently by assigning the lowly confident parameters to neutral values and keeping the high confident parameters unchanged. As described in Section I, we should assess the confidence of parameters according to their contributions to the expected information distance. Therefore, the choice of coordinates is crucial for the CIF principle. This strategy is infeasible in terms of \( p \)-coordinates, \( \eta \)-coordinates, or \( \theta \)-coordinates for two main reasons: first, the orthogonality does not hold in those coordinates, and hence, we cannot safely prune some parameters without affecting the values of others; second, in those coordinate systems, highly confident parameters cannot be significantly distinguished from lowly ones. In this section, we will focus on the \( l \)-mixed coordinates \([\zeta]\) and show how \([\zeta]\) meets all requirements of CIF.

We will first show that \([\zeta]\) meets the requirements of CIF in typical distributions that generate real-world data sets, so as to grasp an intuitive picture for the general CIF strategy. Then, we will prove that CIF could lead to an optimal submanifold with respect to the parametric reduction problem in 1, in general cases.

#### A. CIF in Typical Distributions

In this section, we consider the typical situation in real-world data collections to facilitate our analysis, where there are only a small fraction of all system states are frequent and meaningful patterns [39]. More formally, the underlying distributions \( q(x) \) is assumed to have at least \((2^n - 2^{n/2})\) \( p \)-coordinates of the scale \( \epsilon \), where \( \epsilon \) is a sufficiently small value. Therefore, the residual at most \( 2^{n/2} \) \( p \)-coordinates is of scale \( \Theta(1/2^{(n/2)}) \), and their sum approximates 1.

Next, for the true distribution \( q(x) \), we introduce a small perturbation \( \Delta p \) to the \( p \)-coordinates \([p]\), and the perturbed distribution is denoted by \( q'(x) \). For those \( p \)-coordinates with a sufficiently small value, we assume that the scale of the fluctuation \( \Delta p_I \) for \( p_I \) is proportional to \( a \cdot p_I \), where \( a \) is a small constant coefficient. For \( p \)-coordinates that are significantly larger than zero, the scale of the fluctuation \( \Delta p_I \) for \( p_I \) is assumed to be proportional to the standard
deviation of the estimate of the corresponding $p_1$. According to the Cramér–Rao bound theory, the standard deviation can be approximated by the inverse of the square root of the Fisher information. After some algebra, we can hence assume the perturbation $\Delta p_1$ to be $a(p_1)^{1/2}$.

In our previous work [40], we have analyzed the $l$-mixed coordinates $[\zeta]$ of $\Delta \eta_q = (\Delta \eta^2; \Delta \theta_{\zeta})$, where $l = 2$. The incremental coordinates of mixed coordinates caused by perturbation $\Delta p$ is denoted as $\Delta \eta_q = (\Delta \eta^2; \Delta \theta_{\zeta})$. By decomposing the squared FID $D^2(q, q') = \Delta \eta_q^T G \Delta \eta_q$ into the direction of each coordinate in $[\zeta]$, we have shown that the scale of the FID in each coordinate of $\eta^l$ is significantly larger than that of $\theta_{\zeta}$ (refer to our previous paper for more details [40]).

As described earlier, the coordinates of confidence parameters (measured by the decomposed FID) in $[\zeta]$ entail a natural hierarchy. Moreover, the parameters in $[\eta^l]$ are orthogonal to the ones in $[\theta_{\zeta}]$ [37]. Additionally, those low confident parameters $[\theta_{\zeta}]$ have the neutral value of zero. Hence, we can perform parametric reduction in $[\zeta]$ using the CIF principle by setting low confident parameters to be 0 and reconstructing the distribution according to the new coordinates. The resulting submanifold $M$ tailored by CIF becomes $[\zeta]_{M_i} = (\eta^l_{i_1, \ldots, i_k}, 0, \ldots, 0)$, which is called $[\zeta]_{M_i}$ the $l$-tailed-mixed-coordinates.

To verify our theoretical analysis, we conduct a simulation on the ratio of FID that is preserved by the $l$-tailed-mixed-coordinates $[\zeta]_{M_i}$ with respect to the original mixed-coordinates $[\zeta]$. We also show the corresponding ratio of preserved parameters: number of free parameters/total number of parameters. First, we randomly select real distribution $p_1$ with $n$ variables, where the distribution satisfies the basic assumption that we make in the beginning of this section. The $2n/2$ significant $p$-coordinates are generated based on Jeffery prior (the Dirichlet distribution with alpha parameters set to 0.5), and the left $p$-coordinates are set to a small constant. Then, we generate the sample distribution $p_2$ based on random samples drawn from the real distribution. Last, we calculate the FID between $p_1$ and $p_2$ in terms of the $[\zeta]$ and $[\zeta]_{M_i}$, respectively. The result is shown in Table I. We can see that $[\zeta]_{M_i}$ can indeed preserve most of the FID, which is consistent with our theoretical analysis.

### B. CIF Leads to an Optimal Submanifold $M$

Let $B_q$ be a $\varepsilon$-sphere surface centered at $q(x)$ on manifold $S$, i.e., $B_q = \{q' \in S||K L(q, q') = \varepsilon\}$, where $K L(\cdot, \cdot)$ denotes the KL divergence and $\varepsilon$ is small. Let $q(x)$ be a close neighbor of $q(x)$, which is uniformly sampled from the surface $B_q$, as shown in Fig. 2. Recall that, for a small $\varepsilon$, the KL divergence can be approximated by half of the squared FID. Thus, using the parameterization of $[\zeta]$, $B_q$ is indeed the surface of a hyperellipsoid determined by $G \zeta$ with the center at $q(x)$. The expected FID can indeed preserve most of the FID, which is consistent with our previous work [40].

### IV. Interpretation of Boltzmann Machine Using CIF

In Section III, an implementation of CIF is uncovered in the $[\zeta]$ coordinates for multivariate binary distributions. Next, when $l$ equals to 2, we show that the optimal submanifold $M$ derived by CIF is exactly the BM. This can help us uncover and formalize the essential parts of the target density that BM aims to capture and the nonessential parts that BM discards.

#### A. Introduction to the Boltzmann Machines

In general, a BM [41] is defined as a stochastic neural network consisting of visible units $x \in \{0, 1\}^{n_x}$ and hidden units $h \in \{0, 1\}^{n_h}$. The parameters of a BM are usually denoted as $\theta$, and the network $\theta$ is denoted as $\theta(x, h) = \log \frac{P(x, h)}{P(x)}$, where $P(x)$ and $P(x, h)$ are the marginal distributions of $x$ and $P(x, h)$, respectively. The goal of BM is to learn the parameters $\theta$ of $P(x, h)$ from the observed data.

#### B. Learning BM Using CIF

In our previous work [40], we have analyzed the $l$-mixed coordinates $[\zeta]$ of $\Delta \eta_q = (\Delta \eta^2; \Delta \theta_{\zeta})$, where $l = 2$. The incremental coordinates of mixed coordinates caused by perturbation $\Delta p$ is denoted as $\Delta \eta_q = (\Delta \eta^2; \Delta \theta_{\zeta})$. By decomposing the squared FID $D^2(q, q') = \Delta \eta_q^T G \Delta \eta_q$ into the direction of each coordinate in $[\zeta]$, we have shown that the scale of the FID in each coordinate of $\eta^l$ is significantly larger than that of $\theta_{\zeta}$ (refer to our previous paper for more details [40]).

As described earlier, the coordinates of confidence parameters (measured by the decomposed FID) in $[\zeta]$ entail a natural hierarchy. Moreover, the parameters in $[\eta^l]$ are orthogonal to the ones in $[\theta_{\zeta}]$ [37]. Additionally, those low confident parameters $[\theta_{\zeta}]$ have the neutral value of zero. Hence, we can perform parametric reduction in $[\zeta]$ using the CIF principle by setting low confident parameters to be 0 and reconstructing the distribution according to the new coordinates. The resulting submanifold $M$ tailored by CIF becomes $[\zeta]_{M_i} = (\eta^l_{i_1, \ldots, i_k}, 0, \ldots, 0)$, which is called $[\zeta]_{M_i}$ the $l$-tailed-mixed-coordinates.

To verify our theoretical analysis, we conduct a simulation on the ratio of FID that is preserved by the $l$-tailed-mixed-coordinates $[\zeta]_{M_i}$ with respect to the original mixed-coordinates $[\zeta]$. We also show the corresponding ratio of preserved parameters: number of free parameters/total number of parameters. First, we randomly select real distribution $p_1$ with $n$ variables, where the distribution satisfies the basic assumption that we make in the beginning of this section. The $2n/2$ significant $p$-coordinates are generated based on Jeffery prior (the Dirichlet distribution with alpha parameters set to 0.5), and the left $p$-coordinates are set to a small constant. Then, we generate the sample distribution $p_2$ based on random samples drawn from the real distribution. Last, we calculate the FID between $p_1$ and $p_2$ in terms of the $[\zeta]$ and $[\zeta]_{M_i}$, respectively. The result is shown in Table I. We can see that $[\zeta]_{M_i}$ can indeed preserve most of the FID, which is consistent with our theoretical analysis.

### B. CIF Leads to an Optimal Submanifold $M$

Let $B_q$ be a $\varepsilon$-sphere surface centered at $q(x)$ on manifold $S$, i.e., $B_q = \{q' \in S||K L(q, q') = \varepsilon\}$, where $K L(\cdot, \cdot)$ denotes the KL divergence and $\varepsilon$ is small. Let $q(x)$ be a close neighbor of $q(x)$, which is uniformly sampled from the surface $B_q$, as shown in Fig. 2. Recall that, for a small $\varepsilon$, the KL divergence can be approximated by half of the squared FID. Thus, using the parameterization of $[\zeta]_{M_i}$, $B_q$ is indeed the surface of a hyperellipsoid determined by $G \zeta$ with the center at $q(x)$. The expected FID can indeed preserve most of the FID, which is consistent with our previous work [40].

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In general, a BM [41] is defined as a stochastic neural network consisting of visible units $x \in \{0, 1\}^{n_x}$ and hidden units $h \in \{0, 1\}^{n_h}$. The parameters of a BM are usually denoted as $\theta$, and the network $\theta$ is denoted as $\theta(x, h) = \log \frac{P(x, h)}{P(x)}$, where $P(x)$ and $P(x, h)$ are the marginal distributions of $x$ and $P(x, h)$, respectively. The goal of BM is to learn the parameters $\theta$ of $P(x, h)$ from the observed data.
The energy function is defined as follows:

\[ E_{BM}(x, h; \xi) = -\frac{1}{2} x^T U x - \frac{1}{2} h^T V h - x^T W h - b^T x - d^T h \]

(12)

where \( \xi = \{U, V, W, b, d\} \) are the free parameters. \( U, V, \) and \( W \) represent the visible-visible, hidden-hidden, and visible-hidden interactions, respectively. \( b \) and \( d \) represent the visible and hidden self-connections, respectively. The diagonals of \( U \) and \( V \) are set to zero. The joint Boltzmann distribution of \( x \) and \( h \) can be expressed as follows:

\[ p(x, h; \xi) = \frac{1}{Z} \exp\{-E_{BM}(x, h; \xi)\} \]

(13)

where \( Z \) is a normalization factor.

1) Coordinates for Boltzmann Machines: Let \( B \) be the set of Boltzmann distributions realized by BM. Actually, \( B \) is a submanifold of the general manifold \( S_{ch} \) over \( \{x, h\} \). From (12) and (13), we can see that \( \xi = \{U, V, W, b, d\} \) plays the role of \( B \)'s coordinates in \( \theta \)-coordinates (5) as follows:

\[
\begin{align*}
\theta_1 & : \theta_1^{x i} = b_{x i}, \quad \theta_1^{h j} = d_{h j} \quad (\forall x_i \in x, \ h_j \in h) \\
\theta_2 & : \theta_2^{x_i x_j} = U_{x_i x_j}, \quad \theta_2^{h_i h_j} = W_{h_i h_j}, \\
\theta_2^{h_i h_j} & = V_{h_i h_j}, \quad (\forall x_i, x_j \in x; \ h_i, \ h_j \in h) \\
\theta_2^{h_i h_j} & : \theta_2^{m i h_{i+1} h_{i+2} \ldots h_{i+m}} = 0, \ m > 2, \\
& \quad (\forall x_i, \ldots, x_j \in x; \ h_0, \ldots, h_0 \in h).
\end{align*}
\]

(14)

So the \( \theta \)-coordinates for BM is given by

\[ [\theta]_{BM} = (\theta_1^{x i}, \theta_1^{h j}, \theta_2^{x_i x_j}, \theta_2^{h_i h_j}, 0, 0, \ldots, 0). \]

(15)

The fully visible BM (VBM) and restricted BM (RBM) are special cases of the general BM. Since VBM has no hidden units \( (n_h = 0) \) and all the visible units are connected to each other, the parameters of VBM are \( \xi_{VBM} = \{U, b\} \) and \( \{V, W, d\} \) are all set to zero. For RBM, it has connections only between hidden and visible units. Thus, the parameters of RBM are \( \xi_{RBM} = \{W, b, d\} \) and \( \{U, V\} \) are set to zero.

2) Gradient-Based Learning of BM: Given the sample \( x \) that generated from the underlying distribution, the ML is a commonly used gradient ascent method for training BM in order to maximize the log likelihood \( \log p(x; \xi) \) of the parameters \( \xi \) [42]. Based on (13), the log likelihood is given as follows:

\[ \log p(x; \xi) = \log \sum_h e^{-E(x, h; \xi)} - \log \sum_{x', h'} e^{-E(x', h'; \xi)}. \]

Differentiating the log likelihood, the gradient with respect to \( \xi \) is as follows:

\[
\begin{align*}
\frac{\partial \log p(x; \xi)}{\partial \xi} &= \sum_h p(h|x) \frac{\partial}{\partial \xi} [-E(x, h; \xi)] \\
& \quad - \sum_{x', h'} p(h'|x') \frac{\partial}{\partial \xi} [-E(x', h'; \xi)].
\end{align*}
\]

(16)

where \( \frac{\partial E(x, h; \xi)}{\partial \xi} \) and \( \frac{\partial E(x', h'; \xi)}{\partial \xi} \) can be easily calculated from (12). Then, we can obtain the stochastic gradient using Gibbs sampling [43] in two phases: sample \( h \) given \( x \) for the first term, called the positive phase, and sample \( (x', h') \) from the stationary distribution \( p(x', h'; \xi) \) for the second term, called the negative phase. Now with the resulting stochastic gradient estimation, the learning rule is to adjust \( \xi \) by

\[ \Delta_{BM} = \epsilon \frac{\partial \log p(x; \xi)}{\partial \xi} \]

\[ \propto -\left\langle \frac{\partial E(x, h; \xi)}{\partial \xi} \right\rangle_0 + \left\langle \frac{\partial E(x', h'; \xi)}{\partial \xi} \right\rangle_{\infty} \]

(17)

where \( \epsilon \) is the learning rate, \( \langle \cdot \rangle_0 \) denotes the average using the sample data, and \( \langle \cdot \rangle_{\infty} \) denotes the average with respect to the stationary distribution \( p(x, h; \xi) \) after the corresponding Gibbs sampling phases.

To avoid the difficulty of computing the log-likelihood gradient, the contrastive divergence (CD) [42] optimizes a different objective function based on KL divergence, shown as follows:

\[ \Delta_{CD} = -\epsilon \frac{\partial (KL(p_0||p) - KL(p_m||p))}{\partial \xi} \]

\[ \propto -\left\langle \frac{\partial E(x, h; \xi)}{\partial \xi} \right\rangle_0 + \left\langle \frac{\partial E(x', h'; \xi)}{\partial \xi} \right\rangle_m \]

(18)

where \( KL(\cdot||\cdot) \) denotes the KL divergence, \( p_0 \) represents the sample distribution, and \( p_m \) denotes the distribution sampled from the Markov chain after running \( m \) steps (begins with the sample data). CD can be seen as an approximation to ML by replacing the last expectation \( \langle \cdot \rangle_{\infty} \) with \( \langle \cdot \rangle_m \).

B. Fully Visible Boltzmann Machine

Consider the parametric reduction on the manifold \( S \) over \( \{x\} \) and result with a \( k \)-dimensional submanifold \( M \) of \( S \), where \( k \ll 2^n - 1 \). \( M \) is set to be the same dimensionality as VBM, i.e., \( k = (n_x(n_x + 1))/2 \), then all candidate \( M \) are comparable to the submanifold \( M_{sub} \) endowed by VBM.

In the following corollary, we theoretically show that the CIF principle leads to the optimal submanifold that is exactly the submanifold specified by Boltzmann machines.

**Corollary 1:** Given the general manifold \( S \) in 2-mixed-coordinates \( \{x\}_2 \), VBM (with coordinates \( \{x\}_2 \)) defines an \( k \)-dimensional submanifold of \( S \) that can maximally preserve the expected squared FID induced by Fisher–Rao metric.

**Proof:** See Appendix E.

To learn such \( \{x\}_2 \), we need to learn the parameters \( \xi \) of VBM, such that its stationary distribution preserves the same coordinates \( \{\eta^2\} \) as target distribution \( q(x) \). Actually, this is exactly what traditional gradient-based learning algorithms intend to do. Proposition 5 shows that the ML learning of VBM is equivalent to learn the coordinates \( \{x\}_2 \): 1) high-order \( (\theta_2) \) components are set to zero and 2) low-order \( (\eta^2) \) components are unchanged comparing to the original distribution.

**Proposition 5:** Given the target distribution \( q(x) \) with 2-mixed coordinates

\[ [x]_2 = (\eta^1, \eta^2, \theta_2) \]

the coordinates of the stationary distribution of VBM trained by ML are uniquely given by [\eta_1^2]
\begin{align*}
C[\eta_2] = (\eta_i^1, \eta_j^2, \theta_{2+} = 0)
\end{align*}

\textbf{Proof:} See Appendix F.

\section*{C. Boltzmann Machine With Hidden Units}

In section IV.B, the CIF principle is applied to models without hidden units and leads to VBM by preserving the first-order and second-order \eta-coordinates. In this section, we will investigate the cases where hidden units are introduced.

Let \(S_{xh}\) be the manifold of distributions over the joint space of visible units \(x\) and hidden units \(h\). A general BM produces a stationary distribution \(p(x, h; \xi) \in S_{xh}\) over \(\{x, h\}\). Let \(B\) denote the submanifold of \(S_{xh}\) with probability distributions \(p(x, h; \xi)\) realizable by BM.

Given any target distribution \(q(x)\), only the marginal distribution of BM over the visible units is specified, leaving the distributions on hidden units vary freely. Let \(H_q\) be the submanifold of \(S_{xh}\), consisting of probability distributions \(q(x, h)\) that have the same marginal distribution as \(q(x)\) and the conditional distribution \(q(h|x)\) of hidden units is realized by the BM’s activation functions with some parameter \(\xi_{bm}\).

Then, the best BM is the one that minimizes the distance between \(B\) and \(H_q\). Due to the existence of hidden units, the solution may not be unique. In this section, the training process of BM is analyzed in terms of manifold projection (described in Section I), following the framework of the learning rule proposed in [33]. And we will show that the learning of BM with hidden units can be interpreted using CIF.

1) \textbf{Iterative Projection Learning for BM:} The learning algorithm using the iterative manifold projection is first proposed in [33] and theoretically compared with expectation and maximization algorithm in [44]. The learning of RBM can be implemented by the following iterative projection (IP) process. Let \(z_{b0}\) be the initial parameters of BM and \(p_0(x, h; \xi_{b0})\) be the corresponding stationary distribution.

For \(i = 0, 1, 2, \ldots\), it has the following.

1) Put \(q_{i+1}(x, h) = \Gamma_H(p_i(x, h; z_{b0}^i))\).

2) Put \(p_{i+1}(x, h; z_{b0}^{i+1}) = \Gamma_B(q_{i+1}(x, h))\)

where \(\Gamma_H(p)\) denotes the projection of \(p(x, h; \xi_{b0})\) to \(H_q\), and \(\Gamma_B(q)\) denotes the projection of \(q(x, h)\) to \(B\). The iteration ends when we reach the fixed points of the projections \(p^*\) and \(q^*\), that is \(\Gamma_H(p^*) = q^*\) and \(\Gamma_B(q^*) = p^*\). The IP process is shown in Fig. 3. The convergence property of this iterative algorithm is guaranteed using Proposition 6.

\textbf{Proposition 6:} The monotonic relation holds in the iterative learning algorithm

\begin{align*}
D[q_{i+1}, p_i] \geq D[q_{i+1}, p_{i+1}] \geq D[q_{i+2}, p_{i+1}]
\end{align*}

where the equality holds only for the fixed points of the projections.

\textbf{Proof:} See Appendix G.

The next two propositions show how the projection \(\Gamma_H(p)\) and \(\Gamma_B(q)\) are obtained.

\textbf{Proposition 7:} Given a distribution \(p(x, h; \xi_{p}) \in B\), the projection \(\Gamma_H(p) \in H_q\) that gives the minimum divergence \(D(H_q, p(x, h; \xi_{p}))\) from \(H_q\) to \(p(x, h; \xi_{p})\) is the \(q(x, h; \xi_{bm}) \in H_q\) that satisfies \(\xi_{bm} = \xi_{p}\).

\textbf{Proof:} See Appendix H.

\textbf{Proposition 8:} Given \(q(x, h; \xi_{q}) \in H_q\) with mixed coordinates \([\xi_{h1}^1, \xi_{h1}^2, \xi_{h2}^1, \xi_{h2}^2, \theta_{2+}]\), the coordinates of the learned projection \(\Gamma_B(q) \in B\) are uniquely given by the tailored mixed coordinates

\begin{align*}
\left[\xi_{h1}^1, \xi_{h1}^2, \xi_{h2}^1, \xi_{h2}^2, \theta_{2+} = 0\right] = (\eta_i^1, \eta_j^2, \eta_{xj}^1, \eta_{xj}^2, \eta_{hi}^1, \eta_{hi}^2, \theta_{2+} = 0) \ (20)
\end{align*}

\textbf{Proof:} This proof comes in three parts.

1) The projection \(\Gamma_B(q)\) of \(q(x, h)\) on \(B\) is unique.

2) This unique projection \(\Gamma_B(q)\) can be achieved by minimizing the divergence \(D[q(x, h), B]\) using the gradient descent method.

3) The mixed coordinates of \(\Gamma_B(q)\) are exactly the one given in (20).

See Appendix I for the detailed proof.

2) \textbf{Interpretation for BM Learning via CIF:} The IP learning gives us an alternative way to investigate the learning process of BM. Based on the CIF principle in Section III, we can see that the process of the projection \(\Gamma_B(q_i)\) can be interpreted using CIF, i.e., highly confident coordinates \([\eta_i^1, \eta_j^2, \eta_{xj}^1, \eta_{xj}^2, \eta_{hi}^1, \eta_{hi}^2]\) of \(q_i\) are preserved, while lowly confident coordinates \([\theta_{2+}]\) are set to neutral value zero, given in 20.

In summary, the essential parts of the real distribution that can be learned by BM (with and without hidden units) are exactly the confident coordinates indicated by the CIF principle. As a special kind of BM, the commonly used RBM can be analyzed similarly.

\section*{V. EXPERIMENTAL STUDY ON SAMPLE-SPECIFIC CIF}

In section III and IV, when \(p_s\) is unknown and the \(l\)-mixed \(z\)-coordinates \([\eta]\) is adopted, we have shown that there exist globally optimal solutions of CIF for \(k = \sum_{i=1}^l C_n^i\)
Algorithm 1: Adaptive Network Design for BM

Input: Samples $D = \{d_1, d_2, \ldots, d_N\}$; Significance level $\alpha$; Nodes $V = \{x_1, x_2, \ldots, x_N\}$; Edges $U = \{U_{ij}, \forall x_i, x_j\}$; Output: Set of confident edges $U_{conf} \subset U$

$U_{conf} \leftarrow \{\}$

for $U_{ij} \in U$ do

Estimate marginal distribution $p(x_i, x_j)$ from samples

** parameterize to $\zeta$-coordinates: $[\zeta]$**

$\eta_i \leftarrow E_p[x_i]$; $\eta_j \leftarrow E_p[x_j]$ 

$\theta_{ij} \leftarrow \log p_{00} - \log p_{01} - \log p_{10} + \log p_{11}$

$[\zeta] \leftarrow \{\eta_i, \eta_j, \theta_{ij}\}$

** Fisher information of $\theta_{ij}$ in $[\zeta]$**

$g \leftarrow \left(\frac{1}{p_{00}} + \frac{1}{p_{01}} + \frac{1}{p_{10}} + \frac{1}{p_{11}}\right)^{-1}$

** confidence of $\theta_{ij}$ in $[\zeta]$**

$\rho_{ij} \leftarrow \theta_{ij} \cdot g \cdot \theta_{ij}$

** hypothesis test: $\rho_{ij} = 0$ against $\rho_{ij} \neq 0$**

$\pi \leftarrow cd f_{\theta}(N\rho_{ij})$

if $(1 - \pi) \cdot 2 < \alpha$ then

** reject null hypothesis: $\rho_{ij} = 0$**

$U_{conf} \leftarrow U_{conf} \cup \{U_{ij}\}$

end if

end for

return $U_{conf}$

in terms of the optimization problem in 1. For $l = 2$, we further show that the optimal submanifold $M$ derived by CIF is exactly the BM.

Given specific samples, can CIF further recognize less-confident parameters in BM and reduce them properly? For VBM, we will investigate how to use CIF to modify the topology of VBM by reducing less-confident connections among visible units with respect to given samples. We will empirically investigate this sample-specific CIF principle as a heuristic method to decide the priority order of parameters, so as to improve the search efficiency for the model selection of BM without degrading the quality of model selection results.

A. Sample-Specific CIF: Adaptive Model Selection of BM

The data constrain the state of knowledge about the unknown distribution. Let $q(x)$ denote the sampling distribution (representing the data). In order to force the estimate of our probabilistic model [denoted by $p(x; \zeta)$] to meet the data, we could incorporate the data into CIF by recognizing the confidence of parameters $\zeta$ in terms of $q(x)$. Then, parametric reduction procedure can be further applied to modify the topology of VBM adaptively according to the data, as shown in Algorithm 1 and explained as in the following.

As a graphical model, the VBM comprises a set of vertices $V = \{x_1, x_2, \ldots, x_n\}$ together with a set of connections $U = \{U_{ij}, \forall x_i, x_j, i \neq j\}$. The confidence for each connection parameter $U_{ij}$ can be assessed by the parameter choice criterion in CIF, i.e., the contribution to the FID. Based on [33, Th. 1], $U_{ij}$ could be expressed as follows:

$$U_{ij} = \log \frac{p(x_i = x_j = 1|A) \cdot p(x_i = x_j = 0|A)}{p(x_i = 1, x_j = 0|A) \cdot p(x_i = 0, x_j = 1|A)}$$

where the relation hold for any conditions $A$ on the rest variables. However, it is often infeasible for us to calculate the exact value of $U_{ij}$ because of data sparseness. To tackle this problem, we propose to approximate the value of $U_{ij}$ by using the marginal distribution $p(x_i, x_j)$ to avoid the effect of condition $A$. To estimate $p(x_i, x_j)$, we need to go through all $N$ samples and count the number of samples for each assignment of $x_i$ and $x_j$. For example, $p(x_i = 0, x_j = 0) = (\text{count}(x_i = 0, x_j = 0)/N$, and so on.

Let $[\zeta]_{ij} = (\eta_i, \eta_j, \theta_{ij})$ be the mixed-coordinates for the marginal distribution $p(x_i, x_j)$ of VBM. Note that each $\theta_{ij}$ corresponds to one connection $U_{ij}$. Since $\theta_{ij}$ is orthogonal to $\eta_i$ and $\eta_j$, the FID between two distributions can be decomposed into two independent parts: the information distance contributed by $[\eta_i, \eta_j]$ and $[\theta_{ij}]$. For the purpose of parameter reduction, we consider the two close distributions $p_1$ and $p_2$ with coordinates $\zeta_1 = [\eta_i, \eta_j, \theta_{ij}]$ and $\zeta_2 = [\eta_i, \eta_j, 0]$, respectively. The confidence of $\theta_{ij}$, denoted as $\rho(\theta_{ij})$, can be estimated by its contribution to FID between $p_1$ and $p_2$.

$$\rho(\theta_{ij}) = (\zeta_1 - \zeta_2)^T G_{\zeta}(\zeta_1 - \zeta_2) = \theta_{ij} \cdot g_{\zeta}(\theta_{ij}) \cdot \theta_{ij}$$

where $G_{\zeta}$ is the Fisher information matrix in Proposition 3 and $g_{\zeta}(\theta_{ij})$ is the Fisher information for $\theta_{ij}$. Note that the second equality holds, since $\theta_{ij}$ is orthogonal to $\eta_i$ and $\eta_j$.

For a given number of free parameters $k$, we can directly decide the optimal subset of connections for VBM by selecting the top-$k$ confident parameters. In principle, this solution could be globally optimal in the sense of preserving the Fisher–Rao distance, since the orthogonality of the fractional mixed coordinates [45]. However, a global solution is time-consuming. In this paper, we use a greedy heuristic to “approximate” the global solution. Our heuristic separately evaluates the confidence of each parameter $U_{ij}$ with respect to the marginal distribution $p(x_i, x_j)$, instead of the whole joint distribution $p(x_1, x_2, \ldots, x_n)$. Then, to decide whether the FID in the coordinate direction of $\theta_{ij}$ is significant or negligible, we set up the hypothesis test for $\rho$, i.e., null hypothesis $\rho = 0$ versus alternative $\rho \neq 0$. Based on the analysis in [46], we have $N\rho \sim \chi^2(1)$ asymptotically, where the $\chi^2(1)$ is chi-square distribution with degree of freedom 1 and $N$ is the sampling number. In this way, by setting the significant level $\alpha$, we can simply determine the threshold for $\rho$, i.e., only those parameters with confidence value greater than the threshold are selected. For example, when $\alpha = 0.05$ and $N\rho > 5.024$, we can ensure that the FID in the direction of $\theta_{ij}$ (with respect to the, marginal distribution $p(x_i, x_j)$) is significant with at least 95% confidence. This model selection method is shown in Algorithm 1, called the CIF + $\alpha$.

B. Experiments With VBM

In this section, we investigate the density estimation performance of CIF-based model selection methods for VBM,
i.e., CIF + AIC and CIF + α. Two comparative model selection methods are used.

- **StepwiseAIC**: The backward elimination approach is adopted to select the subset of connections with the optimal AIC: starting with all candidate connections, testing AIC after the deletion of each connection, deleting the connection with the lowest AIC, and repeating this process until AIC stops decreasing.

- **L1-norm**: A full VBM is trained to minimize the $L_1$-regularization objective: $-\log L(\xi) + \lambda \|\xi\|_1$, where $\xi$ is the set of parameters, $\log L(\xi)$ denotes the log likelihood, $\lambda$ is the regularization coefficient, and $\|\cdot\|_1$ represents the $L_1$ norm. The optimal coefficient $\lambda_{opt}$ is chosen via fivefold cross validation. Then, those connections with significant absolute values, i.e., $|\xi| > \epsilon$, are selected. Here, $\epsilon$ is certain small positive threshold.

In this paper, we adopt the AIC as $-2 \log L(\xi_k) + 2k$, where $k$ is the number of selected parameters and $L(\xi_k)$ represent the ML of samples. For $L_1$-norm, the threshold $\epsilon$ is searched in the interval from $10^{-3}$ to $10^{-4}$, and it generally achieves good performance at $10^{-3}$.

1) **Experiment on Artificial Data Set—Density Estimation**: The artificial binary data set is generated as follows: we first randomly select the target distribution $q(x)$, which is randomly chosen from the open probability simplex over $n$ random variables using the Jefferys prior [47]. Since the distribution over $n$ binary variables belongs to the family of multinomial distribution, we adopt the Jefferys prior for the $p$-coordinates of $q(x)$, i.e., the Dirichlet distribution with all alpha parameters set to $0.5$. Then, the data set with $N$ samples is generated from $q(x)$. For computation simplicity, the artificial data set is set to be 10-D. The CD learning algorithm is used to train the VBM.

The Full-VBM, i.e., the VBM with full connections is used as baseline. The KL divergence is adopted to measure the performances of the VBM trained by various algorithms. For each sample size $N$, 20 distributions are randomly generated and the averaged KL divergence is reported. Note that this experiment studies on the case that the variable number $n=10$, which is relatively small. Because it is feasible to efficiently evaluate the KL divergence analytically and hence study the proposed algorithms in more details. Changing the number of variables only offers a trivial influence for experimental results, since we obtained similar observations on various variable numbers (not reported here).

**Results and Summary:**

The averaged KL divergences between VBM and the real distribution are shown in Fig. 4. We can see that all model selection methods could improve density estimation results of VBM, especially when the sample size is small ($N = 100–900$). With relatively large samples, the effect of parameter reduction gradually becomes marginal.

For the two methods that use AIC criterion, CIF + AIC and StepwiseAIC achieve a similar performance, as shown in Fig. 4. The StepwiseAIC is a strong baseline, since it extensively searches candidate VBM topologies in a stepwise approach. The result indicates that CIF could generally produce the set of connections which are also preferred in the

![Fig. 4. Averaged density estimation results of VBM for artificial data set.](image-url)

StepwiseAIC. To show this, we make comparison between the topology of VBM selected by both methods in the first row of Fig. 5. For all sample sizes (from 100 to 3000), we can see that the majority of preserved connections are shared for both methods. We also show how the AIC score changes when different numbers of connections are preserved in the second row of Fig. 5. Both methods achieve optimal AIC performance at the (almost) same $k$. In terms of time complexity, CIF + AIC is a linear time algorithm, while StepwiseAIC is $O(K^2)$.

For the two methods that use thresholds to control the choice of parameters, CIF + α shows a better performance than $L_1$-norm when sample size is small. We check the connection weights in $L_1$-norm, and our experiment shows a significant consistency between CIF + α and $L_1$-norm. That is, the parameters selected by CIF + α would also have higher significant absolute values in $L_1$-norm. This observation can be regarded as a mutual confirmation of the CIF principle and the $L_1$-regularization method. Note that, there is a lack of an analytical method to adaptively set the threshold $\epsilon$ in $L_1$-norm for different sample sizes and parameters. That maybe the main reason that the more principled CIF + α achieves superior performance.

We also compare CIF and $L_1$-norm when the number of free parameters $k$ is fixed. For CIF, we simply sort the confidence of connections in descend order, select the top $k$ connections as free parameters and set the weight of residual connections to zero. For $L_1$-norm, we first train the VBM using $L_1$-regularization and then select the $k$ connections with top absolute weights. The result is shown in Fig. 6. We can see that the CIF outperforms $L_1$-norm on most model complexities $k$, which indicates the effectiveness of CIF.

For the two CIF-based method, CIF + α shows a better performance than CIF + AIC when sample size $N$ is small and gradually achieves similar performance along with the increasing $N$. We observe that the CIF + α tends to preserve less connections than CIF + AIC when the sample size is relatively small. This indicates that the AIC criterion may need to increase the penalty term for small sample sizes so as
Fig. 5. Illustration on the behavior of CIF + AIC and StepwiseAIC for different sample sizes ($N = \{100, 900, 3000\}$). The first row shows the topology of VBM selected by both methods; the second row compares the AIC criterion of both methods in different model complexities.

Fig. 6. Density estimation results of CIF and $L_1$-norm when the number of parameters is fixed for different sample sizes ($N = \{500, 1000, 3000\}$).

to achieve a better performance, which will not be explored further in this paper.

2) Experiments on Real Data Sets—Information Retrieval:
In this section, we empirically investigate how the CIF-based model selection algorithm works on real-world data sets in the context of information retrieval. In particular, we use the VBM to learn the underlying probability density of terms in the document [48], which is further used to rank document based on the query likelihood in information retrieval.

Experiments are conducted on three standard TREC collections: WSJ8792 (topics 151–200), ROBUST04 (topics 601–700) and WT10G (topics 501–550). The WSJ and ROBUST collections are relatively small and consist of news articles, science and technology reports, and government documents, whereas WT10G is a larger Web collection. Collections are indexed by Indri 5.3 with Porter stemming and stopwords removed. For each topic, both the title field and desc field are used to generate queries.

We compare six document models: VBM (baseline), VBM + Smooth (use the smoothing approach in [48]), VBM + $L_1$-norm, VBM + StepwiseAIC, and VBM + CIF (apply CIF + AIC or CIF + $\alpha$). Mean average precision is used as the evaluation metric, which is the mean of average precision scores over all the queries. The result is shown in Table II.

For short queries (title field), CIF-based VBM shows a similar performance compare to the VBM without model selection. This is because short query usually contains only 1–4 terms which leads to relatively simple VBMs with respect to training samples in one document, and there is no need to do model selection.

However, for long queries (desc field) with 8–16 terms, all model selection methods show significant improvements over VBM on WSJ and ROBUST04, except for WT10G. Note that the documents length of WT10G (about 380 words) are much longer than that of WSJ and ROBUST04 (about 250 words),
leading to a larger training set. It is the main reason that the improvement becomes less significant on WT10G.

Comparing CIF + AIC with StepwiseAIC, CIF + AIC achieves similar or slightly better performance on all collections. This indicates that CIF could effectively determine the set of connections that are also preferred by StepwiseAIC. CIF + α outperforms L1-norm on all collections, which shows that Algorithm 1 provides a reasonable way to balance between the model complexity and limited training samples.

3) Experiments on Real Data Sets—Density Estimation:

In particular, we use the VBM to learn the underlying probability density over 100 terms of the 20 News Groups binary data set, with different model complexities. There are 18,000 documents in 20 News Groups in total, which is partitioned into two sets: train set (80%) and test set (20%). The learning rate for CD is manually tuned in order to converge properly and set to 0.01. Since it is infeasible to compute the KL divergence due to the high dimensionality, the averaged Hamming distance between the samples in the test data set and those generated from VBM is used to evaluate the performance. Let \( D = \{d_1, d_2, \ldots, d_N\} \) denote the data set of \( N \) documents (each document \( d_i \) is a 100-D binary vector). For the VBM to be evaluated, we first randomly generate \( N \) samples from its stationary distribution \( p(x; \xi_{bm}) \), denoted as \( V = \{v_1, v_2, \ldots, v_N\} \). Then, the averaged Hamming distance \( D_{ham} \) is calculated as follows:

$$D_{ham}(D, V) = \frac{\sum_{d_i \in D} (\min_{v_j \in V} (\text{Ham}[d_i, v_j]))}{N}$$

where \( \text{Ham}[d_i, v_j] \) is the number of positions where the corresponding values are different.

Due to high dimensionality, it is infeasible to compute the AIC criterion. Therefore, we compare four kinds of VBMs.

1) Full-VBM: The VBM with all connections.
2) Rand: Randomly select \( k \) connections to build the VBM.
3) L1-norm: \( k \) connections are selected in descend order based on their absolute values of weights trained with L1-norm.
4) CIF: Select the \( k \) connections in descend order based on their confidences [see 21].

After training all VBMs on the training data set, we evaluate the trained VBMs on the test data set. We search for the best performances with respect to different model complexities, by varying \( k \) from 0 to 4950. We also mark the VBM that is automatically selected by CIF + α and the best \( k \) for L1-norm and CIF. The result is shown in Fig. 7.

We can see that the parameter reduction criteria (CIF, L1-norm, and Rand) significantly improve the performance of the Full VBM, which indicates the existence of overfitting. Comparing the best performances, CIF significantly outperforms Rand and achieves slightly a better performance than L1-norm. The performance of CIF + α is close to the optimal solution of CIF. These reflect the effectiveness of CIF to select suitable VBM with respect to given samples.

VI. CONCLUSION AND FUTURE WORK

In this paper, we studied the parametric reduction and model selection problem for BMs from both theoretical and application perspectives, and proposed a CIF principle as a general framework for the parametric reduction to maximally preserve the confident parameters and ruling out less confident ones. On the theoretical side, we showed that CIF could lead to an optimal submanifold for binary multivariate distributions in terms of 1. Furthermore, we illustrated that the BMs (with or without hidden units) can be derived from the general manifold based on the CIF principle. In the future, CIF could be the start of an information-oriented interpretation of deep learning models where BM is used as building blocks. For DBM [3], several layers of RBMs compose a deep architecture in order to achieve a sufficiently abstract representation at a certain level. The CIF principle can be used to describe how the information flows in the transformations of representation across layers. Each layer of DBM determines a submanifold \( M \) of \( S \), where \( M \) could maximally preserve the highly confident information on parameters. Then, the whole DBM can be seen as the process of repeatedly applying CIF in each layer, achieving a tradeoff between the abstractness of representation features and the intrinsic confidence of information preserved on parameters. The more detailed analysis and CIF-based designs on deep models will be left as future work.

On the application side, we proposed two sample-specific CIF-based model selection methods for BM, i.e., CIF + AIC.
and CIF + \( a \), which can independently select model parameters, or integrate with information criterion, e.g., AIC, by providing a heuristic way to decide the priority order of parameters and improve the search efficiency without degrading the quality of model selection results. In the future, we plan to incorporate the CIF into deep learning models (e.g., DBM) to modify the network topology, such that the most confident information in the data can be well captured.

\section*{References}


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