Markov Chain Monte Carlo

Introduction, Comparison & Analysis

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ABSTRACT

Markov Chain Monte Carlo (MCMC) is a technique to make an estimation of a statistic by simulation in a complex model. Restricted Bolztmann Machine(RBM) is a crucial model in the field of Machine Learning. However, training a large RBM model will include intractable computation of the partition functions, i.e. $Z(\theta)$. This problem has aroused interest in the work of estimation using a MCMC methods. In this paper, we first conduct Metropolis-Hastings Algorithm, one of the most prevalent sampling methods, and analyze its correctness & performance, along with the choice of the accepting rate. We then implement three algorithms: TAP, AIS, RTS, to estimate partition functions of an RBM model. Our work not only give an introduction about the available algorithms, but systematically compare the performance & difference between them. We seek to provide an overall view in the field of MCMC.

1. INTRODUCTION

Markov Chain A Markov Chain is a special stochastic process in which the current state only depends on its previous state, we call this property the Markov property or memorylessness. i.e.

$$P(X_{t+1} = x | X_t, X_{t-1}, \dots) = P(X_{t+1} = x | X_t)$$
 (1)

Given a Markov Chain, if a vector π , has the property:

$$\pi = \pi P \tag{2}$$

then we call π the stationary distribution, it denotes the final state distribution of the stochastic process in a Markov Chain.

Markov Chain has a wide application in many fields in the real world, such as social science[1], econmics & finance[7] and of course, computer science[17], etc.

Markov Chain Monte Carlo If we are given a probability distribution p(x), it would be a great thing if we could generate some samples of it by a simple method, so here comes the Markov Chain Monte Carlo(MCMC) method. If we could construct a Markov Chain which its stationary distribution π just equals to p(x), then we could use this Markov Chain to sample from this distribution p(x). And this is the main idea of MCMC.

In this paper, we implement the Metropolis-Hastings Algorithm[15, 8], which is one of the most widely used sampling method. We also deal with the accepting rate, which I will introduce later, for previous work[19] have shown that the accepting rate may influence the result of the experiment

and there is theoretical support in choosing an optimal accepting rate.

Restricted Boltzmann Machine A Restricted Bolztmann Machine (RBM)[14] is a significant work bringing theory in statistical physics to computer science. By stacking several layers of RBM, we will get a fundamental model, Deep Belief Network[9], in the field of Deep Learning, which is nowadays the hottest class of algorithms used in Machine Learning.

Estimating Partition functions In the process of training an RBM, however, will include incontractable computation of the partition function. When the model grows large, the complexity of this work will be incompletable. The good news is, researches have shown that there are ways to avoid this by using an MCMC approach instead, to estimate it.

In this paper, we implement three prevalent MCMC methods of estimating a partition function, Thouless-Anderson-Palmer Sampling(TAP)[5], Annealed Importance Sampling (AIS)[16, 20], Rao-Blackwellized Tempered Sampling(RTS)[3], respectively, and give an overall comparison on the theory & performance between them.

2. METROPOLIS-HASTINGS

In the Introduction, we have shown the main idea of an MCMC sampling method. In this section, we will introduce the Metropolis Hastings Algorithm and conduct an experiment.

2.1 Algorithm¹

2.1.1 Detailed Balance Condition

Before stepping further into the MH Algorithm, We would first introduce a theorem called the Detailed Balance Condition.

In the introduction, we said that we want to construct a Markov Chain which its stationary distribution $\pi(x)$ just equals to the required probability distribution p(x).

At first, a theorem is needed.

Theorem 2.1 (Detail Balance Condition). Given a non periodic Markov Chain, if

$$\pi(i)P_{ij} = \pi(j)P_{ji}$$
 for all i, j (3)

then $\pi(x)$ is the stationary distribution of this Markov Chain.

So, the key question will be how to construct a Markov Chain which satisfy this Detail Balance Condition.

¹Available at https://github.com/lzhbrian/MCMC/blob/master/metropolis_hastings/metropolis_hasting.R in R[18]

2.1.2 MCMC sampling method

Suppose we already have a transition matrix Q for a Markov Chain, q(i,j) denote the probability of transition from state i to state j. For the general case,

$$p(i)q(i,j) \neq p(j)q(j,i)$$

That is to say, we do not have the detailed balance condition (Theorem 2.1) So we introduce an $\alpha(i, j)$ s.t.

$$p(i)q(i,j)\alpha(i,j) = p(j)q(j,i)\alpha(j,i)$$
(4)

By sysmetrical characteristic, we choose:

$$\alpha(i,j) = p(j)q(j,i) \quad \alpha(j,i) = p(i)q(i,j) \tag{5}$$

So the new Markov Chain Q' would have the property of which its stationary distribution is p(x)

$$p(i)\underbrace{q(i,j)\alpha(i,j)}_{Q'(i,j)} = p(j)\underbrace{q(j,i)\alpha(j,i)}_{Q'(j,i)} \tag{6}$$

We call the $\alpha(i,j)$ we introduced, accepting ratio. It means that, in the original Markov Chain Q, when state i transits to state j with a probability of q(i,j), we accept this transition with a probability of $\alpha(i,j)$

Now, we have derived the MCMC sampling method.

2.1.3 Metropolis-Hastings Algorithm

The MCMC sampling method is a marvellous work. However, it has a critical drawback that if $\alpha(i,j)$ & $\alpha(j,i)$ are too small, we would seldom accept the transition.

A solution is that we multiply both $\alpha(i, j)$ & $\alpha(j, i)$ with a constant to make sure that the larger one between them equals 1. By doing so, we change the accepting ratio to

$$\alpha(i,j) = \min\left\{\frac{p(j)q(j,i)}{p(i)q(i,j)}, 1\right\}$$
 (7)

and now, we get Metropolis-Hastings Algorithm[8].

I would like to further introduce one more concept called accepting rate(not accepting ratio), which denotes the statistic ratio of accepting the transition. i.e. If we request 10 transition and we accept 8 times, then the accepting rate would be 0.8. This concept is crucial when we are dealing with a continual Markov Chain to use the MH algorithm.

2.1.4 Symmetric Case

In a Markov Chain whose transition matrix is symmetric, we have

$$q(i,j) = q(j,i)$$

so the accepting ratio could be simplified to

$$\alpha(i,j) = \min\left\{\frac{p(j)}{p(i)}, 1\right\} \tag{8}$$

which is also known as the Metropolis Algorithm[15].

2.1.5 Continual Case

In a continual Markov Chain, such as the experiment we are going to do in the next subsection, we have a vague definition of transition matrix Q. So we introduce a concept called the proposal jump size, sd.T.

The method we get x_{k+1} from x_k is to add a sampled point of a normal distribution with a variance of the jump size and $\mu = 0$. For a two dimension example, we have:

$$x_{k+1} = x_k + sd.T \begin{pmatrix} norm_1 \\ norm_2 \end{pmatrix}$$
 (9)

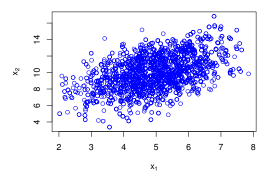


Figure 1: Sampling result of 5,000 points correlation = 0.50009, set sd.T = 3.0

The pseudo code of this algorithm is shown below:

2.2 Sampling Experiment

For our experiment, we use an example of a bivariate Normal distribution, with

$$\mu = \begin{pmatrix} 5\\10 \end{pmatrix}, \Sigma = \begin{pmatrix} 1 & 1\\1 & 4 \end{pmatrix}$$

By theoretical computation, we can easily compute the the pearson correlation between the two dimensional value is 0.5.

$$\rho = 0.5$$

We then generate 10,000 samples using the MH algorithm and take the second half (i.e. the last 5,000 points), setting the standard deviation of proposal to 3.0. We can see from the result (Figure 1) that we have derived 5,000 sampled points whose pearson correlation value $\rho=0.50009$, which matches the theoretical value.

2.3 Performance Analysis

2.3.1 Choice of proposal jump size

MH algorithm is an effective MCMC method for many diverse problems. However, for a continual case in MH algorithm, its performance somewhat depends on the selection of the proposal density. With the proposal jump size being small, the accepting rate would be very low and eventually stick to only one point(eg. the initial point); When the proposal jump size is too big, the accepting rate would be too high.

Roberts et al. have shown in previous work[19] that the optimal accepting rate of the MH algorithm should approximately be at 0.234 for the case of an N-dimensional Gaussian target distribution. We test the accepting rates in different proposal jump size(Figure 2) and find that the optimal value should be at appoximately 3.0 to acquire a model with accepting rate being close to 0.234. That is the reason why we choose 3.0 as our proposal jump size.

2.3.2 Efficiency

Due to the limit of the accepting rate, for a high dimensional condition, using the MH sampling methods may spend

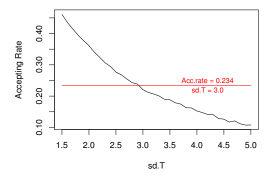


Figure 2: Accepting rate on different proposal jump size.

more time in traverse all of the possible states, which could sometimes be be less satisfying. Thus, many would switch to Gibbs Sampling Algorithm.

2.4 Gibbs Sampling

Gibbs Sampling is a special case of Metropolis Hastings Algorithm, by letting the accepting rate = 1, we will get a Gibbs Sampler. As the length & time limit, we will not specify more here. But it is worthy to notice that Gibbs sampling method is used more often than Metropolis-Hastings method in the real practice, probably because it has a slightly simpler process.

3. PARTITION FUNCTION ESTIMATION

3.1 Restricted Bolztmann Machine

Co-invented and enhanced largely[9] by Geoff Hinton, a Restricted Bolzmann Machine(RBM)[14] is a model which brings the idea of a physics concept to the field of computer science.

3.1.1 Introduction

An RBM is a two-layer undirected model (Figure 3). The first layer of the RBM is called visible layer, and the second is called the hidden layer. In the model, every visible units are connected to all hidden units and vice versa. For every given value of visible layer ${\bf v}$ & hidden layer ${\bf h},$ we can define an energy of this state.

$$E(\mathbf{v}, \mathbf{h}; \theta) = -\mathbf{v}^T \mathbf{W} \mathbf{h} - \mathbf{b}^T \mathbf{v} - \mathbf{a}^T \mathbf{h}$$
 (10)

where $\theta = \{W, \mathbf{b}, \mathbf{a}\}$ are the model configurations. W_{ij} represents the weight between visible unit v_i and hidden unit h_j . $\mathbf{b} & \mathbf{a}$ are biases for visible and hidden layer, respectively.

3.1.2 Training an RBM

On training an RBM, we want our RBM model to have a lowest scale of energy. By doing so, we have to calculate the joint distribution over the visible and hidden units, which is defined by:

$$p(\mathbf{v}, \mathbf{h}; \theta) = \frac{e^{-E(\mathbf{v}, \mathbf{h}; \theta)}}{Z(\theta)}$$
(11)

where

$$Z(\theta) = \sum_{\mathbf{v}} \sum_{\mathbf{h}} e^{-E(\mathbf{v}, \mathbf{h}; \theta)}$$
 (12)

is the partition function.

However, calculating partition functions has always been an intractable work since we have to traverse all the possible state of \mathbf{v} & \mathbf{h} . When the model grows large, this process will be very time & rescouces consuming and thus become unrealistic for the real practice.

So, we have to introduce methods to estimate the partition functions instead of just calculating it in brute force. Although some deviation may include in the estimation, but the efficiency along with them make them preferable. In fact, studies have shown that only few deviation is included that we could just ignore it since it does petty influence on our training.

In the next subsection, we will discuss about three methods available, which each have their pros and cons in doing this complex estimation.

3.2 Algorithms

3.2.1 Thouless-Anderson-Palmer Sampling²

Algorithm Thouless-Anderson-Palmer Sampling (TAP)[5] is a very efficient and easy-to-practice iterative procedure based on an improved mean field method from statistical physics called Thouless-Anderson-Palmer approach.

The main idea of this method is to iteratively compute the magnetization vector m^v, m^h , and then input the values into the Legendre transform of the free energy $F = log(Z(\theta))$ to compute it.

The Legendre transform of F to the second order is:

$$\Gamma(\mathbf{m}^{v}, \mathbf{m}^{h}) \approx -S(\mathbf{m}^{v}, \mathbf{m}^{h}) - \sum_{i} a_{i} m_{i}^{v} - \sum_{j} b_{j} m_{j}^{h}$$

$$- \sum_{i,j} \left(W_{i,j} m_{i}^{v} m_{j}^{h} - \sum_{i} \left(W_{i,j} m_{i}^{v} m_{j}^{h} - \left(W_{i,j}^{h} m_{i}^{v} - \left(W_{i,j}^{h} \right)^{2} \right) \right)$$

$$- 0.5 W_{ij} \left(m_{i}^{v} - \left(W_{i,j}^{v} \right)^{2} \right) \left(m_{j}^{h} - \left(W_{j}^{h} \right)^{2} \right)$$

$$(13)$$

where $S(\mathbf{m}^v, \mathbf{m}^h)$ indicates the entropy:

$$S(\mathbf{m}^v, \mathbf{m}^h) = -\sum_{i} \left(m_i^v log m_i^v + (1 - m_i^v) log (1 - m_i^v) \right)$$
$$-\sum_{j} \left(m_j^h log m_j^h + (1 - m_j^h) log (1 - m_j^h) \right)$$
(14)

Practice In the real practice, in the next subsection, we see TAP method can obtain a converged result in a very short time, but has less accuracy. And sometimes, the converged results are periodic, which is not what we want by us

The pseudo code of this algorithm is shown below:

3.2.2 Annealed Importance Sampling³

 $^{^2}$ Available at https://github.com/lzhbrian/MCMC/blob/master/rbm/TAP.m in Matlab

³Available at https://github.com/lzhbrian/MCMC/blob/master/rbm/AIS.m in Matlab

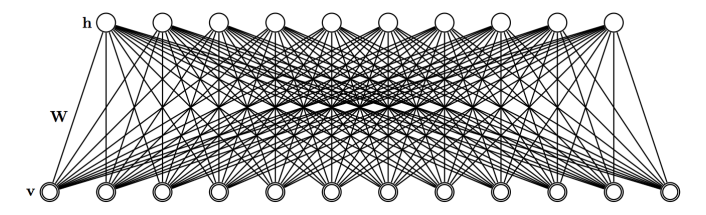


Figure 3: A Restricted Boltzmann Machine

Algorithm Annealed Importance Sampling (AIS) [16, 20] is probably one of the most preferable estimating methods avaible.

Previous work [13] have shown that if P_A and P_B in the SIS method is not close enough, the estimator would be very poor.

Based on SIS, the main idea of this algorithm is to gradually alter the value from an known Z_A to our required Z_B (or Z_K), by the following identity:

$$\frac{Z_K}{Z_0} = \frac{Z_1}{Z_0} \frac{Z_2}{Z_1} \dots \frac{Z_K}{Z_{K-1}} \tag{15}$$

where

$$\frac{Z_K}{Z_{k+1}} = \frac{1}{M} \sum_{i=1}^{M} \frac{P_{k+1}^*(\mathbf{x}^{(i)})}{P_k^*(\mathbf{x}^{(i)})} \quad where \ x^{(i)} \sim P_k$$
 (16)

in which we can get x_{k+1} from:

$$p(h_{j}^{A} = 1|\mathbf{v}) = sigmoid\left((1 - \beta_{k})\left(\sum_{i} W_{ij}^{A} v_{i} + a_{j}^{A}\right)\right)$$

$$p(h_{j}^{B} = 1|\mathbf{v}) = sigmoid\left(\beta_{k}\left(\sum_{i} W_{ij}^{B} v_{i} + a_{j}^{B}\right)\right)$$

$$p(v_{i}' = 1|\mathbf{h}) = sigmoid\left((1 - \beta_{k})\left(\sum_{j} W_{ij}^{A} h_{i}^{A} + b_{i}^{A}\right)\right)$$

$$+ \beta_{k}\left(\sum_{j} W_{ij}^{B} h_{j}^{B} + b_{i}^{B}\right)\right)$$

$$(17)$$

this procedure is shown in Figure 4.

Note that model A indicates an initial model which we can easily compute all its configurations. Commonly, we choose an RBM model with $\theta = \{0,0,0\}$

 β in the above equations is defined by users as a set of inverse temperatures $\{0 = \beta_1 < \beta_2 < ... < \beta_K = 1\}$, which can define a sequence of

$$P_k(\mathbf{x}) \propto P_A^*(\mathbf{x})^{1-\beta_k} P_B^*(\mathbf{x})^{\beta_k} \tag{18}$$

where

$$P_k^*(\mathbf{v}) = \sum_{h^A h^B} e^{(1-\beta_k)E(\mathbf{v}, \mathbf{h}^A; \theta_A) + \beta_k E(\mathbf{v}, \mathbf{h}^B; \theta_B)}$$
(19)

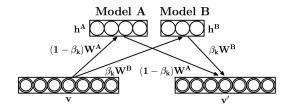


Figure 4: The transition process from x_k to x_{k+1} which leaves $P_k(\mathbf{v})$ invariant.

Initialize Z_A with dataset In [20], Ruslan also notice a method to make Z_A near Z_B . As the length & time limit, we will not specify the process here.

Originally, we initialize model A with a configuration of $\theta = \{0, 0, 0\}$. This method can use the training data to initialize the visible bias **b** to a desired value s.t. we can get a better outcome of the estimation.

In our real practice, we find that this method take less than 0.005 second to initialize **b** even for a very big model (i.e. 784 visible and 500 hidden units), but have strongly improved the result as we will mention in the next subsection.

The pseudo code of this algorithm is shown below:

3.2.3 Rao-Blackwellized Tempered Sampling⁴

Algorithm Similar to AIS, Rao-Blackwellized Tempered (RTS)[3] Sampling also has a set of inverse temperatures $\{0 = \beta_1 < \beta_2 < ... < \beta_K = 1\}$, which can define a sequence of

$$f_k(\mathbf{x}) \propto f(\mathbf{x})^{\beta_k} p_1(\mathbf{x})^{1-\beta_k}$$
 (20)

Different from AIS, we do not traverse β . Instead we sample a β^* every loop, from the β set with the distribution $(\beta|x)$.

Subsequently, we sample from x_k to x_{k+1} by the probability of $q(x|\beta^*)$ just like what we did in AIS, shown in Figure 4. However, what also different from AIS is that, we have to iterate from x_k to x_{k+1} many times(i.e. 50 times in[3]) for the sake of getting a better x_{k+1} .

At the last of every loop, we update the lower variance

 $^{^4{\}rm Available}$ at https://github.com/lzhbrian/MCMC/blob/master/rbm/RTS.m in Matlab

estimator $\hat{\mathbf{c}}$ by

$$\hat{c}_k = \hat{c}_k + \frac{1}{N} q(\beta_k | x) \tag{21}$$

Finally, we get Z_k by

$$\hat{Z}_k^{RTS} = \hat{Z}_k \frac{r_1 \hat{c}_k}{r_k \hat{c}_1}, \quad k = 2, ..., K$$
 (22)

in which what we do care is $Z_B \approx \hat{Z}_K^{RTS}$.

The posterier distribution $q(\beta_k|x)$ in the above equations is defined by:

$$q(\beta_k|x) = \frac{f_k(x)r_k/\hat{Z}_k}{\sum_{k'=1}^K f_{k'}(x)r_{k'}/\hat{Z}_{k'}}$$
(23)

Practice In the paper[3], Carlson et al. note an initializing method to initialize Z_k , whose procedure is just like the above process. The only difference is that they sampled β_k by uniform distribution in every loop, not by the distribution $(\beta|x)$. They claim that after doing such initializing work, then we conduct the algorithms above would acquire a better result.

In our real practice, we directly use the initializing method mentioned above by selecting β_k with a uniform distribution in every loop. We also initialize the value of Z_A by the method we have mentioned in the AIS section using the dataset. And we have found that the result is already satisfying, there is no need to conduct more loops with β_k sampled by $(\beta|x)$.

Also, we found that we have to conduct the procedure above for several times s.t. we can acquire our desired partition function value.(i.e. We did it for 100 times, that is to say we update \mathbf{Z} for 100 times).

The pseudo code of this algorithm is shown below:

3.2.4 Other method

There are many other methods which can also estimate the partition functions. Such as Self-adjusted mixture sampling(SAMS)[21] proposed a method to estimate multiple partition functions together to improve the efficiency. As the length & time limit, we only implement 3 methods here in this paper.

3.3 Estimating Results

We estimate the results of the three algorithms using 4 models with 10, 20, 100, 500 hidden units respectively, all 4 models have 784 visible units. The models are trained by the MNIST handwritten digit dataset[12].

Note that we only calculate the real value of the partition function in the model with 10 hidden units $(log Z(\theta) = 226.11)$, due to my poor laptop has broken down several times when calculating the model with 20 hidden units and maltab doesn't even allow to calculate the other two models because they require unimaginable quantity of memories.

AIS By setting number of iteration at 100 times, and β uniformly sampling from 0 to 1 for 10,000 points.

From Figure 5, we can see that the estimation results will differ a lot if we use an initialization method to initialize the visible bias **b**, as mentioned in the previous subsection. In a model with 10 hidden units. We see AIS with init almost obtain the real value (226.11) without error.

RTS

TAP When running a TAP, we see we can get converged

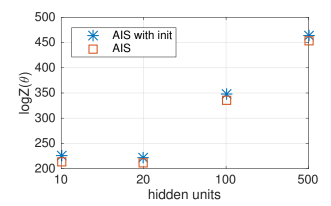


Figure 5: Comparison of AIS method, with or without an initialization of b

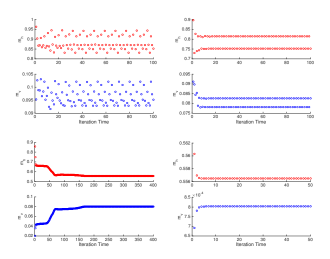


Figure 6: m_h , m_v convergence condition using TAP method. From left to right, up to down, the figure indicates an RBM model with 10, 20, 100, 500 hidden units. The convergence time is approximately 40, 20, 175, 5. By such few iteration time, all results can be obtained in less than 1 second.

 $m_v \& m_h$ in very few iteration time (Figure 6, convergence time is approximately 40, 20, 175, 5 for the model with 10, 20, 100, 500 hidden units respectively). Given this observation, the computing time of TAP is negligeable (we obtain all results in less than a second).

However we do notice that the converged results are sometimes not consistent but periodic. (eg. Figure 6, when there are 10, 20 hidden units), this is not a good news because even if we have more resources to compute the iterations, we would not have a better result.

And disappointingly, compared to the result of other algorithms (Figure ??), TAP usually have a lower estimation value, which is not preferable.

If we don't consider time & resources, we can get good results for all three methods (Figure x).

3.4 Performance Analysis

Since TAP's result is relatively irrelevant to the run time and resources. We use the value estimated in the last sub-

section.

Practice 3.4.1

Complexity For horizontal comparison on complexity, we let the correctness & stability to be approximately the same (i.e. the correctness the AIS could achieve in 2 seconds), and compare the run time for three algorithms.

We have conducted 20 experiments for the acquistion of

From our practice, we have found that xxx gives the most agreeable result while TAP completely underestimate the

Correctness & Stability For horizontal comparison on the correctness & stability, we allow the same run time for AIS and RTS algorithms to compute (i.e. 10 seconds per experiment). We have conducted 20 experiments for the acquistion of $Z(\theta)$.

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We also notice that the variance of ... is the smallest which indicates it has the best stability.

3.4.2 **Theoretical**

RTSFrom a theretical perspective, we have proven that the bias & the variance of the RTS method are to be:

$$E[log\hat{Z}_k^{RTS}] - E[Z_k] \approx \frac{1}{2} \left[\frac{\sigma_1^2}{\hat{c}_1^2} - \frac{\sigma_k^2}{\hat{c}_k^2} \right]$$
 (24)

$$Var[log\hat{Z}_{k}^{RTS}] \approx \frac{\sigma_{1}^{2}}{\hat{c}_{1}^{2}} + \frac{\sigma_{k}^{2}}{\hat{c}_{k}^{2}} - \frac{2\sigma_{1k}}{\hat{c}_{k}\hat{c}_{1}}$$
 (25)

where $\sigma_k^2 = Var[\hat{c}_k]$ and $\sigma_{1k} = Cov[\hat{c}_1, \hat{c}_k]$

This has shown that the bias of RTS has no definite sign.

However, in AIS, Neal and Jarzynski et al.[16, 11] have shown that if we want the result to be unbiased, we would have to let M=1 in the iteration, which by doing so have lost the advantage of AIS. That is to say, on the other hand, if M > 1, we would have a negative bias due to Jenson Inequality.

Although TAP shows the best efficiency, its results are the most disapointing. Apparantly TAP has underestimate the value of the partition function.

We did not analyze deep on the reason why it failed to perform a satisfying result, but our intuition tell that maybe it is because the Legendre transform. In our practice, we only took the Legendre transform to the 2nd order, which might result in the underestimation.

Performance Analysis

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 (26)

$$Var[log\hat{Z}_{k}^{RTS}] \approx \frac{\sigma_{1}^{2}}{\hat{c}_{1}^{2}} + \frac{\sigma_{k}^{2}}{\hat{c}_{k}^{2}} - \frac{2\sigma_{1k}}{\hat{c}_{k}\hat{c}_{1}}$$
 (27)

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RELATED WORK

In this paper, we do not dig very deep into a specific field of the algorithms, but provide an overall view of the MCMC method. There are brilliant seniors who have done many marvellous works in different specific field.

Metropolis-Hastings algorithm Gilks[6] and Chib et al.[4] did a significant job in giving a clear introduction and deeper explanation to the Metropolis-Hastings Algorithm. Beskos et al.[2] made a great work in analysing the complexity of the Algorithm.

Partition function estimation Hubbard[10] is probably the earliest reseacher managing to reduce the complexity of calculating a partition function. Jarzynski[11] systemetically analyzed the bias between the estimation and the real value of the partition functions in an sampling process.

Due to my limited knowledge, there might be some mistakes and flaws in this paper, please don't hesitate to contact and correct me.

CONCLUSION

In this paper, we discuss about the Markov Chain Monte Carlo method which are now undoubtedly one of the most important sampling methods.

We comprehensively introduce the concept of Metropolis-Hastings Algorithm and conduct an experiment to verify its correctness. We also make some analysis about how accepting rate would interfere the sampling result.

We systematically compare three methods of partition function estimation which are crucial works in training a Restricted Bolztmann Machine or a Deep Belief Network.

As future work, we would like to join more methods to the comparison and if could, propose some improvement to the algorithms available.

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