

Asymmetric Gaussian Mixtures with Reversible Jump MCMC

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Abstract—We propose a fully Bayesian learning approach using reversible jump Markov chain Monte Carlo (RJMCMC) for asymmetric Gaussian mixtures (AGM). Compared to classic Gaussian mixture model, AGM doesn't imply that target data is symmetric which brings flexibility and better fitting results. This paper also introduces a RJMCMC learning implementation based on Metropolis-Hastings (MH) within Gibbs sampling method. As an improvement of traditional sampling-based MCMC learning, RJMCMC has no assumption concerning the number of components and, therefore, the AGM model itself could be transferred between iterations. For better evaluating models with different mixture components numbers, the model selection is achieved by calculating integrated likelihood using Laplace approximation to figure out the best-fit components number. We selected both synthetic and challenging spam filtering dataset to show the merits of the proposed model.

Index Terms—Asymmetric Gaussian Mixture, Metropolis-Hastings, Gibbs sampling, RJMCMC, Laplace approximation, Spam Filtering

I. INTRODUCTION

Statistics reveal a crucial fact that more than 59% of worldwide e-mail traffic is considered as unsolicited messages, also well known as spams, in 2017 [1]. Most spams are irritating and resource-consuming, and some of them are extremely dangerous in terms of phishing scam, fee fraud, job offer scam, etc.. Since the damages of spam are persistent and significant not only for individuals but also for governments, companies and organizations, many spam filtering technologies have been proposed to address this issue and eliminate unwanted e-mails automatically over recent decades.

Most modern machine learning-based filtering approaches can be classified into two categories: supervised and unsupervised. As an important solution for spam filtering, supervised approaches [2] perform well under some circumstances, but compared to unsupervised learning methods, they have significant limitations and drawbacks because supervised classifiers cannot identify new spam patterns not presented in their training datasets. Once new patterns are discovered, model adjustment will be needed. Furthermore, poor training datasets could cause inductive bias and overfitting problems which will affect the accuracy of the models. Therefore, unsupervised solution has been increasingly drawing attention because of its flexibility and robustness.

As widely deployed unsupervised learning approaches, mixture models can be viewed as an improvement of independent methodologies which superimpose a finite number of components while respecting the dependency between data clusters, demonstrating outstanding suitability and generality especially for complex high-dimensional datasets. More precisely, for Gaussian-like datasets, Gaussian mixture model (GMM) [3] is proven as an effective learning approach in several domains such as computer vision, pattern recognition and data mining. In this paper, we show the merits of asymmetric Gaussian mixture (AGM) model [4] for clustering because of its two variance parameters for left and right parts of each distribution in the mixture which brings more accuracy of fitting real datasets which could be asymmetric or even non-Gaussian.

Estimating the parameters of mixture models could be a challenging task. The maximum-likelihood-based expectation maximization (EM) [5] algorithm is one of the most popular parameter learning approaches. However, the disadvantages of EM algorithm are also obvious. Given the fact that EM approximates values of mixture parameters in a deterministic way this could cause slow convergence and compromise the usability of the algorithm. Furthermore, bad initialization and overfitting problems [6] [7] will also significantly affect its accuracy. Therefore, fully Bayesian learning algorithms, such as Markov Chain Monte Carlo (MCMC) based implementations, are found to be useful to eliminate overfitting problems in mixture parameter learning by introducing prior and posterior distributions for mixture parameters. In this paper, the learning process is accomplished by a hybrid MCMC algorithm, which is well known as Metropolis-Hastings within Gibbs sampling [6], based on both Metropolis-Hastings (Hastings, 1970) [8] and Gibbs sampling (Geman and Geman, 1984) [9] methods because the main difficulty of classic MCMC method is that, under some circumstances, direct sampling is not always straightforward. Moreover, we reinforce the learning algorithm by introducing reversible jump MCMC (RJMCMC) [7] methodology to increase the flexibility of AGM model by allowing model transfer throughout iterations via increasing (component birth/split step) and decreasing (component death/merge step) mixture components. Because of the stochastic sampling-based learning process, learning iterations could end up with different number of components so

we choose marginal likelihood [6] to perform model selection in order to evaluate fitting results between models.

The rest of this paper is organized as follows. In the next Section, we present our Bayesian model. Section III is devoted to the experimental results. Section IV concludes the paper.

II. BAYESIAN MODEL

A. Asymmetric Gaussian Mixture Model

The likelihood function of AGM model [4] with M mixture components can be defined as follows:

$$p(\mathcal{X}|\Theta) = \prod_{i=1}^N \sum_{j=1}^M p_j p(X_i|\xi_j) \quad (1)$$

where $\mathcal{X} = (X_1, \dots, X_N)$ represents the dataset with N observations, $\Theta = \{p_1, \dots, p_M, \xi_1, \dots, \xi_M\}$ defines the mixture parameters set of AGM mixture model including component weight p_j ($0 < p_j \leq 1$ and $\sum_{j=1}^M p_j = 1$) and asymmetric Gaussian distribution (AGD) parameters set ξ_j for mixture component j . Assuming the dataset \mathcal{X} is d -dimensional, for each observation $X_n = (x_{n1}, \dots, x_{nd}) \in \mathcal{X}$, the probability density function [4] for j -th component of the model can be defined as follows:

$$p(X_n|\xi_j) \propto \prod_{k=1}^d \frac{1}{(\sigma_{ljk} + \sigma_{rjk})} \times \begin{cases} \exp\left[-\frac{(x_{nk} - \mu_{jk})^2}{2(\sigma_{ljk})^2}\right] & \text{if } x_{nk} < \mu_{jk} \\ \exp\left[-\frac{(x_{nk} - \mu_{jk})^2}{2(\sigma_{rjk})^2}\right] & \text{if } x_{nk} \geq \mu_{jk} \end{cases} \quad (2)$$

parameters set of component j is $\xi_j = (\mu_j, \sigma_{lj}, \sigma_{rj})$ where $\mu_j = (\mu_{j1}, \dots, \mu_{jd})$ is the mean, $\sigma_{lj} = (\sigma_{lj1}, \dots, \sigma_{lj d})$ and $\sigma_{rj} = (\sigma_{rj1}, \dots, \sigma_{rj d})$ are the left and right standard deviation vectors of AGD.

We introduce a M -dimensional membership vector Z for each observation $X_i \in \mathcal{X}$, $Z_i = (Z_{i1}, \dots, Z_{iM})$ which indicates to which specific component X_i belongs [10], such that:

$$Z_{ij} = \begin{cases} 1 & \text{if } X_i \text{ belongs to component } j \\ 0 & \text{otherwise} \end{cases} \quad (3)$$

in other words, $Z_{ij} = 1$ only if observation X_i has the highest probability of belonging to component j and accordingly, for other components, $Z_{ij} = 0$.

Therefore, the complete likelihood function can be derived by combining Eq. (1) and Eq. (3) as follows:

$$p(\mathcal{X}, Z|\Theta) = \prod_{i=1}^N \prod_{j=1}^M (p_j p(X_i|\xi_j))^{Z_{ij}} \quad (4)$$

B. Priors and Posteriors

As discussed before, MH-within-Gibbs based RJMCMC learning algorithm implementation defines priors and posteriors for mixture weights and parameters to avoid direct sampling. For a specific iteration t , since mixture weight p_j

satisfies $0 < p_j \leq 1$ and $\sum_{j=1}^M p_j = 1$, a nature choice of the prior is Dirichlet distribution [11] as follows:

$$\pi(p_j^{(t)}) \sim \mathcal{D}(\gamma_1, \dots, \gamma_M) \quad (5)$$

where γ_j is a known hyperparameter. By considering the membership vector Z as a condition, the posterior probability of mixture weight p_j is defined as follows:

$$p(p_j^{(t)}|Z^{(t)}) \sim \mathcal{D}(\gamma_1 + n_1^{(t)}, \dots, \gamma_M + n_M^{(t)}) \quad (6)$$

where n_j represents the number of observations belonging to component j which could be calculated by using membership vectors as follows:

$$n_j^{(t)} = \sum_{i=1}^N Z_{ij} \quad (j = 1, \dots, M) \quad (7)$$

The same idea applies to the sampling process of mixture parameters. The proposal posterior distribution is $\xi^{(t)} \sim q(\xi|\xi^{(t-1)})$. To be more specific, for parameters of AGM model $\xi^{(t)} = (\mu^{(t)}, \sigma_l^{(t)}, \sigma_r^{(t)})$, we choose d -dimensional Gaussian distributions as posterior distributions respectively:

$$\mu_j^{(t)} \sim \mathcal{N}_d(\mu_j^{(t-1)}, \Sigma), \sigma_{lj}^{(t)} \sim \mathcal{N}_d(\sigma_{lj}^{(t-1)}, \Sigma), \sigma_{rj}^{(t)} \sim \mathcal{N}_d(\sigma_{rj}^{(t-1)}, \Sigma) \quad (8)$$

where Σ is $d \times d$ identity matrix which makes the sampling a random walk MCMC process. Correspondingly, the priors are $\mu \sim \mathcal{N}_d(\eta, \Sigma)$ and $\sigma_l, \sigma_r \sim \mathcal{N}_d(\tau, \Sigma)$ given known hyperparameters η and τ .

C. Learning Algorithm

MH-within-Gibbs: As a sampling-based learning algorithm, MH-within-Gibbs method performs random sampling from posteriors of parameters, and then calculates the acceptance ratio r in order to make a decision whether the new samples should be accepted or discarded for the next iteration. Because of the usage of membership vector Z , the mixture weight p_j can be derived within Gibbs sampling part. Therefore, it will be excluded from the calculation of the acceptance ratio r which is defined as follows:

$$r = \frac{p(\mathcal{X}|\Theta^{(t)})\pi(\Theta^{(t)})q(\Theta^{(t-1)}|\Theta^{(t)})}{p(\mathcal{X}|\Theta^{(t-1)})\pi(\Theta^{(t-1)})q(\Theta^{(t)}|\Theta^{(t-1)})} \quad (9)$$

Once acceptance ratio r is derived, we compute acceptance probability $\alpha = \min[1, r]$ [12]. Then $u \sim U_{[0,1]}$ is supposed to be generated randomly. If $u < \alpha$, the proposed move should be accepted and parameters should be updated by $p^{(t)}$ and $\xi^{(t)}$ for the next iteration. Otherwise, we discard $p^{(t)}$, $\xi^{(t)}$ and set $p^{(t)} = p^{(t-1)}$, $\xi^{(t)} = \xi^{(t-1)}$.

RJMCMC moves: Traditional MH-within-Gibbs algorithm assumes that the components number M is given and persistent throughout the learning process. However, because of bad initialization or just leaking of information, components number M could be inaccurate or unknown. Under these circumstances, RJMCMC algorithm is found to be useful by providing extra four independent steps (birth/split steps and

death/merge steps) into learning process which could change components number M , therefore, brings more generalities.

Letting M_{min} and M_{max} denote the minimum and maximum number of components M , we assume that the probabilities of performing birth/split and death/merge steps are b_m and $d_m = 1 - b_m$ for $m = M_{min}, \dots, M_{max}$ respectively. Obviously, $b_{M_{max}} = 0$ and $d_{M_{min}} = 0$. Correspondingly, $d_{M_{max}} = 1 - b_{M_{max}} = 1$ and $b_{M_{min}} = 1 - d_{M_{min}} = 1$. For $m = M_{min} + 1, \dots, M_{max} - 1$, due to simplification reasons, we choose the same value for both b_m and d_m as $b_m = d_m = 0.5$. Within every iteration, we generate a random value $u' \sim U_{[0,1]}$ respectively for the four RJMCMC steps. If $b_m \geq u'$ or $d_m \geq u'$, birth/split or death/merge steps should be performed correspondingly. [3]

Merge and Split Steps: Randomly choose two components (j_1, j_2) satisfying that $\mu_{j_1} < \mu_{j_2}$ with no other μ_j in the interval $[\mu_{j_1}, \mu_{j_2}]$. The newly merged component j' will contain the observations that previously belong to both component j_1 and j_2 . Meanwhile, reduce current value of components number $m = m - 1$, then calculate mixture weight and parameters for j' as follows:

$$\begin{aligned} p_{j'} &= p_{j_1} + p_{j_2} \\ p_{j'} \mu_{j'} &= p_{j_1} \mu_{j_1} + p_{j_2} \mu_{j_2} \\ p_{j'} (\mu_{j'l}^2 + \sigma_{j'l}^2) &= p_{j_1} (\mu_{j_1l}^2 + \sigma_{j_1l}^2) + p_{j_2} (\mu_{j_2l}^2 + \sigma_{j_2l}^2) \\ p_{j'} (\mu_{j'r}^2 + \sigma_{j'r}^2) &= p_{j_1} (\mu_{j_1r}^2 + \sigma_{j_1r}^2) + p_{j_2} (\mu_{j_2r}^2 + \sigma_{j_2r}^2) \end{aligned} \quad (10)$$

As a reverse of merge step, we split component j' into two (j_1 and j_2) with 3 degrees of freedom ($u_1 \sim Beta(2, 2), u_2 \sim Beta(2, 2), u_3 \sim Beta(1, 1)$) and, accordingly, increase $m = m + 1$. Therefore, mixture parameters for split components can be calculated as follows:

$$\begin{aligned} p_{j_1} &= p_{j'} u_1, p_{j_2} = p_{j'} u_2 \\ \mu_{j_1} &= \mu_{j'} - \frac{u_2(\sigma_{j'l} + \sigma_{j'r})}{2} \sqrt{\frac{p_{j_2}}{p_{j_1}}} \\ \mu_{j_2} &= \mu_{j'} + \frac{u_2(\sigma_{j'l} + \sigma_{j'r})}{2} \sqrt{\frac{p_{j_1}}{p_{j_2}}} \\ \sigma_{j_1l}^2 &= u_3(1 - u_2^2) \sigma_{j'l}^2 \frac{p_{j'}}{p_{j_1}} \\ \sigma_{j_1r}^2 &= u_3(1 - u_2^2) \sigma_{j'r}^2 \frac{p_{j'}}{p_{j_1}} \\ \sigma_{j_2l}^2 &= (1 - u_3)(1 - u_2^2) \sigma_{j'l}^2 \frac{p_{j'}}{p_{j_2}} \\ \sigma_{j_2r}^2 &= (1 - u_3)(1 - u_2^2) \sigma_{j'r}^2 \frac{p_{j'}}{p_{j_2}} \end{aligned} \quad (11)$$

In order to decide whether the merge and split steps should be accepted or not, we calculate the acceptance probability \mathcal{A} which is described in [3]. Therefore, the acceptance probability for merge step is $\min(1, \mathcal{A})$ and, correspondingly, for split step is $\min(1, \mathcal{A}^{-1})$.

Birth and Death Steps: Compared to merge and split steps, birth and death steps are relatively straightforward because the newborn and dead components are empty ones which means parameter re-calculation is not needed. Mixture weight p_{new} in

birth step can be obtained by sampling from Beta distribution $p_{new} \sim Beta(1, m)$ and mixture parameters can be derived from the priors as follows [13]:

$$\mu \sim \mathcal{N}(\xi, \kappa^{-1}), \quad \sigma_l^{-2}, \sigma_r^{-2} \sim \Gamma(\alpha, \beta), \quad \beta \sim \Gamma(g, h) \quad (12)$$

where hyperparameters κ, α, g and h are estimated by data and ξ is the midpoint of the observations. For death step, an empty component should be randomly selected and deleted among the existing components if there is any. Otherwise, this step will be skipped. After birth and death steps, mixture weight p_j should be re-scaled so that all weights sum to 1. Acceptance probability \mathcal{A}' for birth and death steps is also required as the one for merge and split steps. The probabilities of occurrence of birth and death steps are $\min(1, \mathcal{A}')$ and $\min(1, \mathcal{A}'^{-1})$ [3].

Finally, a typical reversible jump MH-within-Gibbs learning procedure for AGM model can be summarized as follows:

Input: Data observations \mathcal{X} and components number M

Output: AGM mixture parameter set Θ

1) Initialization

2) Step t : For $t = 1, \dots$

Gibbs sampling part

a) Generate $Z^{(t)}$ from Eq. (3)

b) Compute $n_j^{(t)}$ from Eq. (7)

c) Generate $p_j^{(t)}$ from Eq. (6)

Metropolis-Hastings part

d) Sample $\xi_j^{(t)}(\mu_j^{(t)}, \sigma_{lj}^{(t)}, \sigma_{rj}^{(t)})$ from Eqs. (8)

e) Compute acceptance ratio r from Eq. (9)

f) Generate $\alpha = \min[1, r]$ and $u \sim U_{[0,1]}$

g) If $u \geq \alpha$ then $\xi^{(t)} = \xi^{(t-1)}$

RJMCMC part

h) Generate $u' \sim U_{[0,1]}$. If $b_m \geq u'$, perform split or birth step, then calculate acceptance probability \mathcal{A} . If the step is accepted, set $m = m + 1$.

i) Generate $u' \sim U_{[0,1]}$. If $d_m \geq u'$, perform merge or death step, then calculate acceptance probability \mathcal{A}' . If the step is accepted, set $m = m - 1$.

III. EXPERIMENTAL RESULTS

A. Synthetic Data

We start by testing the performance of our model on a 3-D synthetic dataset with 400 observations composed of clusters. Hyperparameters related to the calculations of mixture

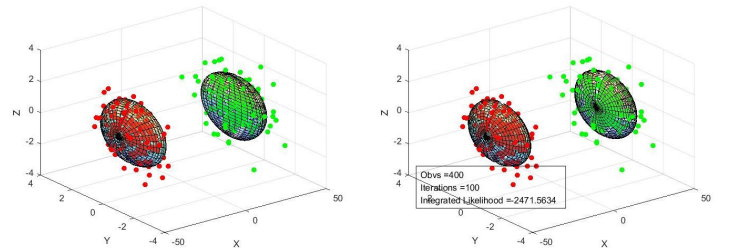


Fig. 1. Original synthetic data grouping and learning results

weights and parameters are set as $\gamma_j = 1$ and both η and τ are considered as d -dimensional zero vectors. Consequently, hyperparameters for priors of mixture parameters are set as follows [14]:

$$\kappa = \frac{1}{\mathcal{R}^2}, \quad \alpha = 3, \quad g = 0.3, \quad h = \frac{100g}{\alpha \mathcal{R}^2} \quad (13)$$

where \mathcal{R} is the interval of variation of observations.

Figure 1 compares original grouping and RJMCMC learning results. Euclidean distances between means of original and estimated clusters are 1.7496 and 1.6263 and standard deviations for σ_l and σ_r are [0.60272, 1.3511] and [1.3209, 0.78647], demonstrating a promising accuracy of AGM model.

B. Spam Filtering

A well organized Spambase dataset [15] is selected with attributes related to multiple spam textual features including spam word/character dictionaries and profiles of uninterrupted capital letter sequences. Data pre-processing includes Scaling-based data normalization which re-scales numerical values within the range between 0 and 1 and label extraction for generating confusion matrix.

To better evaluate the performance and accuracy of AGM model under different initial number of components, the integrated likelihood [6] values are given in Table I to identify the best-fit result. Obviously, the result with initial component number $m = 3$ has the largest integrated likelihood value (8.4238e5). Therefore, we select it as the best-fit result and make horizontal comparison with GMM. Statistics in Table II reveal the fact that comparing to GMM, AGM provides higher accuracy and precision, additionally, lower false positive rate and false negative rate indicate that AGM outperforms GMM. However, because of the nature of spambase, the performance of both mixture models is not satisfactory since most of spams

cannot be identified. Therefore, data-based adjustment of the model might lead to a better result in the future.

IV. CONCLUSION AND FUTURE WORK

A fully Bayesian analysis based on reversible jump MCMC of AGM model is proposed. The RJMCMC learning algorithm allows transfers between AGM models with different amount of mixture components which brings flexibility and generality. We show the merits of this model by applying it into a challenging spam filtering database and the horizontal comparison reveals that AGM outperforms GMM in terms of better statistical measurements. Meanwhile, model adjustments are needed which might lead to a better output in the future.

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TABLE I
AGM STATISTICS

Init. Comp. Number m	Accuracy	Integrated Likelihood
$m = 1$	55.64%	5.7074e5
$m = 2$	51.21%	4.0543e5
$m = 3$	58.99%	8.4238e5

TABLE II
CONFUSION MATRICES AND STATISTICS OF GMM AND AGM

GMM			AGM		
	NF ^a	F ^b		NF	F
NF	35	1778	NF	249	1564
F	295	2493	F	323	2465

	GMM	AGM
Accuracy	54.94%	58.99%
Precision	1.93%	13.81%
False Positive Rate	41.63%	38.81%
False Negative Rate	89.39%	56.46%

^aNon fault-prone, ^bFault-prone.