Mixture of Multilayer Perceptron Regressions

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Abstract: This paper investigates mixture of multilayer perceptron (MLP) regressions. Although mixture of MLP regressions (MoMR) can be a strong fitting model for noisy data, the research on it has been rare. We employ soft mixture approach and use the Expectation-Maximization (EM) algorithm as a basic learning method. Our learning method goes in a double-looped manner; the outer loop is controlled by the EM and the inner loop by MLP learning method. Given data, we will have many models; thus, we need a criterion to select the best. Bayesian Information Criterion (BIC) is used here because it works nicely for MLP model selection. Our experiments showed that the proposed MoMR method found the expected MoMR model as the best for artificial data and selected the MoMR model having smaller error than any linear models for real noisy data.

1 INTRODUCTION

Mixture models have been widely used in econometrics, marketing, biology, chemistry, and many other fields. The book by McLachlan and Peel (McLachlan and Peel, 2000) contains a comprehensive review of finite mixture models.

When data arise from heterogeneous contexts, it is reasonable to introduce mixture of regressions as a class of mixture models. In mixture of regressions, since the introduction by Goldfeld and Quandt (Goldfeld and Quandt, 1973), mixture of linear regressions (MoLR) has been focused (Bishop, 2006; Qian and Wu, 2011) and implemented as library programs (Leisch, 2004; NCSS, 2013). Around that time, Bayesian approaches to mixture of regressions were vigorously investigated using Markov chain Monte Carlo (MCMC) methods (Hurn et al., 2003).

Since this world is full of nonlinear relationships, mixture of nonlinear regressions may have the great potential. The research on the topic, however, has been relatively few. Huang, Li, and Wang (Huang et al., 2013) investigated mixture of nonlinear regressions by employing kernel regression, but they assumed that explanatory variable is univariate and the extension to multivariate will suffer from curse of dimensionality; this can be a serious limitation.

As another approach, modal regression (Chen et al., 2016) estimates the local modes of the distribution of a dependent variable given a value of an explanatory variable. Modal regression, however, will not give us an explicit representation and the extendability to multivariate data seems not clear.

Since multilayer perceptron (MLP) is a popular powerful nonlinear model, mixture of MLP regressions (MoMR) will be quite a reasonable model of mixture of nonlinear regressions; however, MoMR has hardly been addressed so far.

This paper investigates MoMR. There can be two types of mixture: hard and soft. In hard mixture a data point is exclusively classified, while in the latter a data point belongs probabilistically to every class. Since soft mixture is more natural for modeling and more appropriate for computation, we employ soft mixture approach, and use the Expectation-Maximization (EM) algorithm (Dempster et al., 1977).

This paper is organized as follows. Section 2 reviews the background of our research, and Section 3 explains modeling, EM solver, and model selection of MoMR. Then Section 4 describes our experiments performed to examine how our MoMR works using a two-class artificial dataset and a noisy real dataset.

2 BACKGROUND

2.1 EM Algorithm

The EM algorithm is a general-purpose iterative algorithm for maximum likelihood (ML) estimation in in-
complete data problems (Dempster et al., 1977). The EM and its variants have been applied in many applications (McLachlan and Peel, 2000).

Suppose that a data point \((x, z)\) is generated with the density \(p(x, z|\theta)\), where only \(x\) is observable and \(z\) is hidden. Here \(\theta\) denotes a parameter vector, and let \(p(x|\theta)\) be the density generating \(x\). In the EM context, \(\{x^\mu, z^\mu\}\) is called complete data, and \(\{x^\mu\}\) is called incomplete data, where \(\mu = 1, \cdots, N\).

The purpose of ML estimation is to maximize the following log-likelihood from incomplete data.

\[
L(\theta) = \sum_\mu \log p(x^\mu|\theta).
\]

The EM performs ML estimation by iteratively maximizing the following \(Q\)-function, where \(\theta^{(t)}\) is the estimate obtained after the \(t\)-th iteration.

\[
Q(\theta|\theta^{(t)}) = \sum_\mu \sum_{z^\mu} P(z^\mu|x^\mu, \theta^{(t)}) \log p(x^\mu, z^\mu|\theta),
\]

where \(P(z^\mu|x^\mu, \theta^{(t)}) = \frac{p(x^\mu, z^\mu|\theta^{(t)})}{\sum_{\mu'} p(x^\mu, z^\mu'|\theta^{(t)})}\).

The EM algorithm goes as below.

**EM Algorithm**

1. Initialize \(\theta^{(0)}\) and \(t \leftarrow 0\).
2. Iterate the following EM-step until convergence.
   - **E-step:** Compute \(Q(\theta|\theta^{(t)})\) by computing the posterior \(P(z^\mu|x^\mu, \theta^{(t)})\).
   - **M-step:** \(\theta^{(t+1)} = \text{argmax}_\theta Q(\theta|\theta^{(t)})\) and \(t \leftarrow t + 1\).

It can be shown that the EM iteration makes the likelihood \(L(\theta)\) increase monotonically: that is, \(L(\theta^{(t+1)}) \geq L(\theta^{(t)})\), which means \(\theta^{(t)}\) converges to a local maximum.

### 2.2 MLP Learning Methods

In this paper we employ three MLP learning methods described below. Hereafter MLP\((J)\) indicates MLP having \(J\) hidden units and one output unit.

The BP algorithm (Rumelhart et al., 1986) is well-known method of MLP learning. BP uses only the gradient and goes in an online mode. BP is beautifully simple and easily adaptable to many layers, used even for deep learning (Goodfellow et al., 2016).

Although BP is widely used, its learning speed is usually very slow and its capability to find excellent solutions is quite limited; thus, to accelerate the convergence and improve the limited capability, several methods were proposed (Luenberger, 1984). Here we employ quasi-Newton method called BPQ (BP based on quasi-Newton) (Saito and Nakano, 1997). BPQ uses the Broyden-Fletcher-Goldfarb-Shanno (BFGS) update to get a search direction, and uses 2nd-order approximation to get a suitable step length. Getting a suitable step length usually requires a lot of time, but 2nd-order approximation is carried out very quickly.

Recently, singularity stairs following (SSF) has been proposed as a very powerful learning method of single MLPs (Satoh and Nakano, 2013). SSF successively learns MLPs to stably and systematically find excellent solutions, making good use of singular regions generated by using the optimal solution of one-step smaller model MLP\((J-1)\), and guaranteeing monotonic decrease of training errors.

### 2.3 Model Selection

Since we consider many candidates of mixture models, we need a criterion to evaluate the desirability of each candidate. For this purpose we make use of information criterion, which indicates a trade-off between learning error and model complexity. Although many information criteria have been proposed so far, we employ the Bayesian information criterion BIC (Schwarz, 1978), because BIC stably showed nice performance on MLP model selection (Satoh and Nakano, 2017).

Let \(p(x|\theta)\) be a learning model with parameter vector \(\theta\). Given data \(\{x^\mu, \mu = 1, \cdots, N\}\), the log-likelihood is defined as shown in eq.(1) Let \(\hat{\theta}\) be a maximum likelihood estimate. BIC is obtained as an estimator of free energy \(F(D)\) shown below, where \(p(D)\) is called evidence and \(p(\theta)\) is a prior of \(\theta\).

\[
F(D) = - \log p(D),
\]

\[
p(D) = \int p(\theta) \prod_{\mu=1}^{N} p(x^\mu|\theta) \, d\theta
\]

BIC is derived using the asymptotic normality and Laplace approximation.

\[
\text{BIC} = -2L(\hat{\theta}) + M \log N
\]

\[
= -2 \sum_\mu \log p(x^\mu|\hat{\theta}) + M \log N (6)
\]

BIC is calculated using only one point ML estimate \(\hat{\theta}\), where \(M\) is the number of parameters.

We consider another important measure for regression: goodness of fit. Total sum of squares (TSS) indicates how much variation the data have, residual sum of squares (RSS) indicates the discrepancy between the data and the estimates, and explained sum of squares (ESS) indicates how well a regression model represents the data. Given data \(\{x^\mu, y^\mu, \mu = 1, \cdots, N\}\), TSS, RSS, and ESS are given as below, where \(x\) are explanatory variables, \(y\) is a dependent
variable, $f_\mu = f(x_\mu)$ is an estimate obtained by a regression function, and $\bar{y}$ is a mean of $y$.

$$TSS = \sum_\mu (y_\mu - \bar{y})^2, \quad RSS = \sum_\mu (f_\mu - y_\mu)^2$$

(7)

Thus, ESS/TSS ($= 1 - \text{RSS/TSS}$) is an important measure indicating goodness of fit of a regression model. It is also called coefficient of determination in the linear regression context.

3 Mixture of MLP Regressions

3.1 Modeling of MoMR

This subsection formalizes the MoMR model.

Let $x = (x_1, \ldots, x_E)^T$ be $K$ explanatory variables, and $y$ be a dependent variable. In this paper $a^T$ denotes the transpose of $a$.

Given data $\{(x_\mu, y_\mu), \mu = 1, \ldots, N\}$, we consider a mixture of $C$ regression functions. Let $f(x|\theta_c)$ be a regression function of class $c = 1, \ldots, C$, where $\theta_c$ is the weight vector. Since each regression function is supposed to have a constant term, we extend a vector of explanatory variables to get $\tilde{x} = (1, x_1, \ldots, x_E)^T$.

MLP of class $c$ has $K$ input units, $L$ hidden units, and one output unit. It also has weight vectors $w^{(c)}_j$ between all input units and hidden unit $j = 1, \ldots, L_c$, and weights $v^{(c)}_j$ between hidden unit $j = 0, 1, \ldots, L_c$ and an output unit. Then its regression function is defined as follows.

$$f(x|\theta_c) = s_0^{(c)} + \sum_{j=1}^{L_c} v^{(c)}_j \sigma \left((w^{(c)}_j)^T \tilde{x}\right)$$

(9)

Here $\theta_c = (s_0^{(c)}, v^{(c)}_1, \ldots, v^{(c)}_L, (w^{(c)}_1)^T, \ldots, (w^{(c)}_{L_c})^T)^T$ for $c = 1, \ldots, C$, and $\sigma(h)$ denotes the sigmoid activation function. When $L_c = 1$, we consider the following linear regression function instead of MLP($L_c = 1$).

$$f(x|\theta_c) = w^{(c)}_1 \tilde{x}$$

(10)

We assume the value of $y$ is generated by adding a noise $\varepsilon_c$ to a value of $f(x|\theta_c)$; here, $\varepsilon_c$ is supposed to follow the Gaussian with mean 0 and variance $\sigma_c^2$.

$$\varepsilon_c \sim \mathcal{N}(0, \sigma_c^2)$$

(11)

Then, the dependent variable $y$ follows the following distribution.

$$y \sim \mathcal{N}(f(x|\theta_c), \sigma_c^2)$$

(12)

Let $\pi_c$ be the mixing coefficient of class $c$. Then, the density of complete data is described as follows.

$$p(y, c|\theta) = \pi_c \ g_c(y|f(x|\theta_c), \sigma_c^2)$$

(13)

Here $g(u|m, s^2)$ denotes a density function where $u$ follows one-dimensional Gaussian with mean $m$ and variance $s^2$.

$$g(u|m, s^2) = \frac{1}{\sqrt{2\pi} s} \exp \left( -\frac{(u - m)^2}{2 s^2} \right)$$

(14)

The density of incomplete data is written as follows.

$$p(y|\theta) = \sum_{c=1}^{C} p(y, c|\theta) = \sum_c \pi_c \ g_c(y|f(x|\theta_c), \sigma_c^2)$$

(15)

Here $\theta$ is a vector comprised of all parameters, where $\theta_c$ is a parameter vector of class $c$.

$$\theta = (\theta_1^T, \ldots, \theta_C^T, \ldots, \theta_L^T)^T, \quad \theta_c = (\pi_c, w^{(c)}_1, \sigma_c^2)^T$$

(16)

3.2 EM Solver of MoMR

Bishop describes the framework to solve soft mixture of linear regressions (Bishop, 2006). We extend Bishop’s framework to solve soft mixture of nonlinear regressions, including MoMR.

Since class $c$ is a latent variable and cannot be observed, we employ the EM algorithm as a basic learning method to solve the problem.

Posterior probability $P(c|y, \theta)$ indicates the probability that $y$ belongs to class $c$ under $\theta$.

$$P(c|y, \theta) = \frac{p(y, c|\theta)}{\sum_c p(y, c|\theta)}$$

(17)

Given data $D = \{(x_\mu, y_\mu), \mu = 1, \ldots, N\}$, the log-likelihood is defined as below.

$$L(\theta) = \sum_{\mu=1}^{N} \log p(y_\mu|\theta)$$

(18)

The Q-function to maximize is shown as below. Here $\hat{\theta}^{(t)}$ denotes the estimate obtained at the $t$-th step of the EM, and let $f^{(t)}_\mu \equiv f(x|\theta_c)$.

$$Q(\theta|\hat{\theta}^{(t)}) = \sum_{\mu=1}^{N} \sum_c P(c|y_\mu, \hat{\theta}^{(t)}) \log p(y_\mu, c|\theta)$$

$$= \sum_{\mu} \sum_c P_c^{(t)} \log(\pi_c \ g_c(y_\mu|f_\mu, \sigma_c^2))$$

$$= \sum_{\mu} \sum_c P_c^{(t)} \left( \log \pi_c - \frac{1}{2} \log(2\pi) - \log \sigma_c - \frac{(y_\mu - f_\mu)^2}{2\sigma_c^2} \right)$$

(19)
In the above, we use the following for brevity.

\[ P_c^{(t)} \equiv P(c | y^\mu, \theta^{(t)}) = \frac{\pi_c^{(t)} g_c^{(t)}}{\sum_c \pi_c^{(t)} g_c^{(t)}} \] (20)

where \( g_c^{(t)} \equiv g_c(y^\mu f_c^{(t)}, \sigma_c^2) \) (21)

When we maximize the Q-function, we use the Lagrange method since there is an equality constraint \( \sum \pi_c = 1 \). The Lagrangian function can be written as follows with \( \lambda \) as a Lagrange multiplier.

\[ J = Q(\theta | \theta^{(t)}) - \lambda \left( \sum_c \pi_c - 1 \right) \] (22)

The necessary condition for a local maximizer is shown below for \( c = 1, \cdots, C \).

\[ \frac{\partial J}{\partial \pi_c} = \sum_\mu P_c^{(t)} / \pi_c - \lambda = 0 \] (23)

\[ \frac{\partial J}{\partial w_c} = \sum_\mu P_c^{(t)} (y^\mu - f_c^\mu) / \sigma_c^2 \frac{\partial f_c^\mu}{\partial w_c} = 0 \] (24)

\[ \frac{\partial J}{\partial \sigma_c} = \sum_\mu P_c^{(t)} \left( -\frac{1}{\sigma_c} + \frac{(y^\mu - f_c^\mu)^2}{\sigma_c^2} \right) = 0 \] (25)

Since we have \( \lambda = N \) from eq.(23) and the equality constraint, a new estimate of \( \pi_c \) is given below.

\[ \pi_c^{(t+1)} = \frac{1}{N} \sum_\mu P_c^{(t)} \] (26)

From eq.(25) a new estimate of \( \sigma_c^2 \) is given below.

\[ (\sigma_c^2)^{(t+1)} = \frac{\sum_\mu P_c^{(t)} (y^\mu - f_c^\mu)^2}{\sum_\mu P_c^{(t)}} \] (27)

From eq.(24) we obtain a new estimate of \( w_c \) by solving the following.

\[ \sum_\mu P_c^{(t)} (y^\mu - f_c^\mu) \frac{\partial f_c^\mu}{\partial w_c} = 0 \] (28)

Note that the condition eq.(28) is equal to the following optimal condition of \( E_c(w_c) \).

\[ \frac{\partial E_c(w_c)}{\partial w_c} = 0. \] (29)

Here the following is sum-of-squares error of class \( c \).

\[ E_c(w_c) = \frac{1}{2} \sum_\mu P_c^{(t)} (f_c^\mu - y)^2 \] (30)

Residual sum of squares (RSS) in MoMR is given as below.

\[ \text{RSS} = 2 \sum_c E_c(w_c) = \sum_c \sum_\mu P_c^{(t)} (f_c^\mu - y^\mu)^2 \] (31)

In \( E_c(w_c) \), squared error \((f_c^\mu - y^\mu)^2\) for data point \( \mu \) is weighted by posterior \( P_c^{(t)} \). Thus, in MLP learning of class \( c \), the gradient for data point \( \mu \) should be weighted by posterior \( P_c^{(t)} \). This modification should be embodied in MLP learning methods.

The learning of MoMR is carried out in a double-looped manner: the outer loop is controlled by the EM and the inner loop is controlled by MLP learning BP or BPQ.

### 3.3 Model Selection of MoMR

This subsection describes how BIC is calculated in MoMR.

The density of incomplete data is given by eq.(15). Then, log-likelihood at the optimal point \( \hat{\theta} \) is given as follows.

\[ L(\hat{\theta}) = \sum_{\mu=1}^N \log P(y^\mu | \hat{\theta}) \] (32)

Hence, BIC in MoMR is obtained as below.

\[ \text{BIC} = -2 \sum_c \sum_\mu \hat{\pi}_c g_c(y^\mu f(x^\mu | \hat{w_c}), \hat{\sigma}_c^2) + M \log N \] (33)

Here \( M \), the number of parameters, is calculated as follows. We should not forget to count two parameters \( \pi_c \) and \( \sigma_c \) in calculating \( M_c \).

\[ M = \sum_c M_c, \quad M_c = \begin{cases} K + 3 & \text{if } J_c = 1 \\ J_c(K + 2) + 3 & \text{if } J_c \geq 2 \end{cases} \] (34)

TSS, RSS and ESS in MoMR are shown below, where each data point \( \mu \) is weighted by posterior \( P_c^{(t)} \).

\[ \text{TSS} = \sum \sum_\mu P_c^{(t)} (y^\mu - \hat{y})^2 \] (35)

\[ \text{RSS} = \sum \sum_\mu P_c^{(t)} (f_c^\mu - y^\mu)^2 \] (36)

\[ \text{ESS} = \text{TSS} - \text{RSS} \] (37)

Goodness of fit ESS/TSS in MoMR is calculated using the above.

### 4 EXPERIMENTS

#### 4.1 Design of Experiments

The following 26 models are considered for each dataset. Models are given numbers, which are used in the figures and explanations shown later.
(a) Models 1 to 10: 10 single MLP(J) regressions: \( J = 1, \ldots, 10, \)

(b) Models 11 to 16: 6 mixtures of MLP(J1) and MLP(J2) regressions: \( (J_1, J_2) = (1,1), (1,2), (1,3), (2,2), (2,3), (3,3), \)

(c) Models 17 to 26: 10 mixtures of MLP(J1), MLP(J2) and MLP(J3) regressions: \( (J_1, J_2, J_3) = (1,1,1), (1,1,2), (1,1,3), (1,2,2), (1,2,3), (1,3,3), (2,2,2), (2,2,3), (2,3,3), (3,3,3). \)

Note that MLP(J=1) is replaced with linear regression here. Thus, Model 1 is a simple linear regression, Model 11 is a mixture of two linear regressions, and Model 17 is a mixture of three linear regressions. Model 12 is a mixture of one linear regression and one MLP(J=2), and so on. As for the learning of mixture of linear regressions (MoLR), refer to (Nakano and Satoh, 2018). A single MLP(J) regression is learned by SSF or BP if \( J \geq 2. \)

Parameters of BP and BPQ are selected through our preliminary experiments, as shown in Table 1. Very weak regularization of weight decay is employed to prevent weight values from getting huge. Note that the maximum of sweeps per EM loop needs not be large since posterior may change during EM learning. For SSF, maximum of search tokens is set to be 20. We used a PC with Xeon(R)E5 8GB memory for computation.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>BP</th>
<th>BPQ</th>
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<tbody>
<tr>
<td>max sweeps/EM loop (MoMR)</td>
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<td>500</td>
</tr>
<tr>
<td>learning rate (MoMR)</td>
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<td>adaptive</td>
</tr>
<tr>
<td>weight decay coeff (MoMR)</td>
<td>( 10^{-1} )</td>
<td>( 10^{-6} )</td>
</tr>
<tr>
<td>max sweeps (Single)</td>
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<td>5000</td>
</tr>
<tr>
<td>learning rate (Single)</td>
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<tr>
<td>weight decay coeff (Single)</td>
<td>( 10^{-7} )</td>
<td>( 10^{-6} )</td>
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</tbody>
</table>

### 4.2 Experiments using Artificial Data

We generated one-dimensional 2 class artificial data. The following two parabolas were used to generate 51 data points for each class by adding Gaussian noise \( \mathcal{N}(0, 0.035^2) \). The range of \( x_1 \) is \([0.1, 1.0]\).

\[
\begin{align*}
    y_1 &= -4(x_1 - 0.6)^2 + 2.0 \quad (38) \\
    y_2 &= -2(x_1 - 0.6)^2 + 1.5 \quad (39)
\end{align*}
\]

Figure 1 shows two parabolas and 102 data points. Since MLP(J \( \geq 2 \)) can fit a parabola well, two MLPs(J=2) are expected to fit this artificial data well as the minimal model.

Figure 2 compares BIC of each model for artificial data. Horizontal axis indicates model number.

BIC obtained by EM+BPQ was always smaller (better) than the corresponding BIC by EM+BP except pure linear Models 1, 11 and 17. This was caused by BP’s weak capability to find excellent solutions.

BIC obtained by EM+BPQ selected Model 14, two MLPs(J=2), as the best among all the models, which we expected. Figure 3 depicts Model 14. We can see these two curves are very close to the original parabolas.

BIC obtained by EM+BP selected Model 20, one linear and two MLPs(J=2), as the best, whose BIC is larger than Model 14. Figure 4 shows Model 20.

As the best mixture of linear regressions, BIC selected Model 11, which is composed of two lines. Figure 5 depicts Model 11. Its BIC was larger (worse) than that of the best single MLP model (Model 2).

Among single regression models, BIC obtained by SSF selected Model 2, MLP(J=2), while BIC obtained by BP selected wrong Model 1, linear regression. Figure 6 shows Model 2, which runs in a middle empty space between two parabolas.

Figure 7 compares residual sum of squares (RSS) of each model for artificial data. It can be seen that the
solid line (EM+BPQ) always indicates smaller RSS than the dotted line (EM+BP) except pure linear Models 1, 11 and 17. RSS of Model 14, the best model obtained by EM+BPQ, was 0.1046 and thus its goodness of fit $1 - \frac{\text{RSS}}{\text{TSS}}$ was very high 0.9867 since TSS = 7.8436 for artificial data. Moreover, the solid line indicates that mixture models achieved much smaller RSS than single models. Among mixture models, the solid line also indicates that MoMR models had much smaller RSS than mixture of pure linear regressions, Models 11 and 17. Hence we can say MoMR effectively improved goodness of fit compared with single regression models or mixtures of linear regressions.

Figure 8 indicates how RSS decreased through EM learning in the best Model 14. The error decreased very smoothly and monotonically.

The CPU time required to get the results for artificial data is compared below. As average CPU time required to learn 16 MoMR models per initialization, EM+BPQ required 1m 12s, while EM+BP required 7m 16s. Although BPQ computes more information than BP, its average CPU time was smaller because it converged faster for this dataset.

4.3 Experiments using Real Data

As real data we used Abalone dataset from UCI Machine Learning Repository. We selected this dataset because any single powerful regression model cannot fit well. The dataset has seven numerical explanatory variables and the number of data points $N = 4177$.

Figure 9 compares BIC of each model for Abalone data. It can be seen that BIC obtained by EM+BPQ was always much smaller (better) than the corresponding BIC by EM+BP except three pure linear models. BIC(EM+BPQ) selected Model 20, one linear and two MLPs ($J=2$), as the best, while BIC(EM+BP) selected inadequate Model 17, mixture of two linear regressions, as the best. Note that Model 20 had smaller BIC than any single model or any mixture of linear regressions. Among single models MLP($J=7$) is the best single model.
model among all the models obtained by EM+BPQ, was 727.32, then the goodness of fit was $1 - \frac{\text{RSS}}{\text{TSS}} = 0.8258$, showing nice fitting. RSS of Model 17, the best mixture of linear regressions, was 865.32, and its goodness of fit was 0.7928, a bit worse than the best model. Model 24 had the smallest RSS 656.00 among all the models, and its goodness of fit was 0.8429.

Goodness of fit for Abalone data can be improved to this level by using MoMR.

The CPU time required to get the results for Abalone data is shown here. As average CPU time required to learn 16 MoMR models per initialization, EM+BPQ required 6h 7m 40s, while EM+BP required 4h 46m 37s.

4.4 Considerations

The experimental results may suggest the following.

Figure 11 indicates how RSS decreased through EM learning in the best Model 20. The error decreased very smoothly and monotonically.

The CPU time required to get the results for Abalone data is shown here. As average CPU time required to learn 16 MoMR models per initialization, EM+BPQ required 6h 7m 40s, while EM+BP required 4h 46m 37s.

4.4 Considerations

The experimental results may suggest the following.
(a) MoMR worked well, selecting the expected model MLPs($J = 2$) as the best for artificial data, and selecting the model composed of one linear and two MLPs as the best for Abalone data. These best models show smaller BIC and RSS values than those of any mixture of linear regressions or any single MLP regression. 

(b) The learning of MoMR goes in a double loop; the EM controls the outer loop and MLP learning method controls the inner loop. As for MLP learning, a quasi-Newton method called BPQ worked well for MoMR, while BP worked rather poorly, frequently finding rather poor solutions, having larger (worse) RSS than BPQ, selecting inadequate models different from those by BPQ. This tendency was caused by BP’s weak capability to find excellent solutions.

(c) MoMR using EM+BPQ is expected to improve goodness of fit for data having poor fit by any single regression model or mixture of linear regressions.

5 CONCLUSIONS

This paper proposes modeling and learning of mixture of MLP regressions (MoMR). The learning of MoMR goes in a double loop; the outer loop is controlled by the EM and the inner by MLP learning. As for MLP learning in MoMR, a quasi-Newton worked satisfactorily, while BP did not work. Our experiments showed MoMR worked well for artificial and real datasets. In the future we plan to apply MoMR using EM+BPQ to more data to show MoMR can be a useful regression model for noisy data.

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