Subset Selection in High-Dimensional Genomic Data using Hybrid Variational Bayes and Bootstrap priors

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Abstract. In this study, the Variational Bayes (VB) approach was hybridized with the bootstrap prior procedure to improve the accuracy of subset selection as well as optimizing the algorithm time in modelling high-dimensional genomic data with inherent sparse structure. The new hybrid VB approach is shown to yields a minimal sufficient statistic which under mild regularity conditions converges to the true sparse structure. Simulation and real-life high-dimensional genomic data experiments revealed comparable empirical performance with other competing frequentist and Bayesian methods. In addition, a new fast algorithm that illustrates the procedure was developed and implemented in the environment of R statistical software as package “VBbootprior”.

Key Words: Bayesian Inference, Big data, Variational Bayes, Bootstrap prior, R package

1. Introduction

Variable selection is one of the fundamental aspects of statistical modelling as evidenced by numerous authors. The classes of model selection procedures include criteria based approaches [1 - 3], penalized regression approaches [4 - 6] and Bayesian modelling approaches [7 - 10]. There have been several improvements in the area yet there is still no standard view on how to effectively perform subset selection, especially in high-dimensional settings. The primary focus of this paper is to develop a robust strategy for selecting variables in high-dimensional genomic data with inherent classification problem.

The Bayesian model selection strategy is more advantageous due to its ability to incorporate many types of variation as well as the incorporation of prior information. The weak aspect of the procedure is its sensitivity to the knowledge and thus proper specification of prior is highly necessary. One of the common prior used in Bayesian model selection involving linearity of variables and parameters is Zellner’s g-prior [11].

In low dimensional setting, Bayesian model selection is performed by exploring all models involved vis-à-vis Bayesian inference. In contrast, for high-dimensional setting the Markov Chain Monte Carlo (MCMC) methods are employed. MCMC for moderate to large scale problems can be computationally
inefficient [12-18]. For this reason, an enormous amount of effort has been put into developing MCMC and similar stochastic search based methods which can be used to explore the model space in a computationally efficient manner [19-23].

The Mean-field variational Bayes (VB) is a computationally efficient but approximate alternative to MCMC for Bayesian inference [12]. Although, it is difficult to fairly compare MCMC and VB as they differ theoretically. The VB approach is deterministic while MCMC is a stochastic approach, but in most cases, VB tends to be faster than MCMC, especially in high-dimensional settings [24-25].

2. Variational Bayes Bootstrap Prior (VBprior) for Subset Selection in Binary Classification

Suppose we have a binary class problem defined as \( \{ y \in [0, 1] \mid z \} \) governed by the logistic discriminant model:

\[
p(y = 1 \mid z) = \frac{\exp(z\beta)}{1 + \exp(z\beta)}
\]

where \( y \) is the response variable, \( z \) is the matrix of predictors and \( \beta \) is the weight vector.

The classifier often return estimated probability \( \hat{p}(y = 1 \mid z) \) which is then converted to \([0, 1]\) using the scheme below:

\[
\psi = \begin{cases} 
1, & p(y = 1 \mid z) \geq \text{cutpoint} \\
0, & p(y = 1 \mid z) < \text{cutpoint} 
\end{cases}
\]

The cut-point probability is often fixed at 0.5.

Let \( x \in \mathbb{N} \) be the subset of \( z \) that is correctly related to \( y \). Thus we define the probability distribution of \( x \) as Poisson with parameter \( \lambda \), thus;

\[
x \sim \text{Poisson}(\lambda)
\]

with hierarchical prior; \( \lambda \sim \text{Gamma}(a, \theta) \) and \( \theta \sim \text{Gamma}(b, c) \). We want to estimate the posterior of \( p(\lambda, \theta \mid x) \) given the distribution \( q(\lambda, \theta) \). The first step is to obtain the factorization method;

So we choose:

\[
p(\lambda, \theta \mid x) \approx q(\lambda, \theta) = q(\lambda)q(\theta)
\]

From the general approach we can obtain;

\[
q(\theta) \propto \exp \{ E_{q(\lambda)}[\ln p(x \mid \lambda, \theta)] + \ln p(\lambda) + \ln p(\theta) \}
\]

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\]

Since the expectation is only over \( \lambda \), hence it doesn’t affect \( \ln p(\theta) \). Thus;

\[
q(\theta) \propto \exp \{ E_{q(\lambda)}[\ln p(x \mid \lambda, \theta)] \} p(\theta)
\]

\[
p(x \mid \lambda, \theta) = p(x \mid \lambda)p(\lambda \mid \theta) = \frac{\exp(-\lambda)\lambda^x}{x!} \times \frac{\theta^a}{\Gamma a} \lambda^{a-1} \exp(-\theta \lambda).
\]

For iid samples;
\[
p(x_1, \ldots, x_N | \lambda, \theta) = \prod_{i=1}^{N} \frac{\exp(-\lambda) x_i^\lambda}{\Gamma(\lambda)} \times \frac{\theta^a}{\Gamma(a)} \lambda^{a-1} \exp(-\theta \lambda)
\]

\[
lnp(x_1, \ldots, x_N | \lambda, \theta) = \sum_{i=1}^{N} \ln \left( \frac{\exp(-\lambda) x_i^\lambda}{\Gamma(\lambda)} \times \frac{\theta^a}{\Gamma(a)} \lambda^{a-1} \exp(-\theta \lambda) \right)
\]

\[
q(\theta) \propto \exp \left\{ E_q(\lambda) \left[ \sum_{i=1}^{N} \ln \left( \frac{\exp(-\lambda) x_i^\lambda}{\Gamma(\lambda)} \times \lambda^{a-1} \exp(-\theta \lambda) \right) \right] \right\} \lambda^{-a} \exp(-\theta \lambda)
\]

\[
p(\theta) = \frac{c}{b} \theta^{-b} \exp(-c \theta)
\]

\[
q(\theta) \propto \exp \left\{ E_q(\lambda) \left[ \sum_{i=1}^{N} \ln \left( \frac{\exp(-\lambda) x_i^\lambda}{\Gamma(\lambda)} \times \lambda^{a-1} \exp(-\theta \lambda) \right) \right] \right\} \theta^{-b} \exp(-c \theta)
\]

\[
q(\theta) \propto \left[ \prod_{i=1}^{N} \frac{\exp(-\lambda) x_i^\lambda}{\Gamma(\lambda)} \right] \left[ x_i \right]^{a-1} \left( 1 + \theta \right) \exp \left[ E_q(\lambda) \left[ -\lambda \right] \right] \theta^{-b} \exp(-c \theta)
\]

\[
q(\theta) = \text{Gamma}(\theta | a', b')
\]

\[
a' = b + a + \sum_{i=1}^{N} x_i
\]

\[
b' = c + E_q(\lambda) \left[ -\lambda \right]
\]

Next to obtain optimal \( \lambda; \)

\[
q(\lambda) \propto \exp \left\{ E_{q(\theta)} [lnp(x|\lambda, \theta)] + ln p(\lambda) \right\}
\]

\[
q(\lambda) \propto \exp \left\{ E_{q(\theta)} [lnp(x|\lambda, \theta)] \right\} p(\lambda)
\]

\[
p(x|\lambda, \theta) = p(x|\lambda)p(\lambda|\theta) = \frac{\exp(-\lambda) x^\lambda}{\Gamma(\lambda)} \times \frac{\theta^a}{\Gamma(a)} \lambda^{a-1} \exp(-\theta \lambda)
\]

\[
q(\lambda) \propto \exp \left\{ E_{q(\theta)} \left[ \sum_{i=1}^{N} \ln \left( \frac{\exp(-\lambda) x_i^\lambda}{\Gamma(\lambda)} \times \lambda^{a-1} \exp(-\theta \lambda) \right) \right] \right\} \lambda^{-a} \exp(-\theta \lambda)
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q(\lambda) \propto \exp \left\{ E_{q(\theta)} \left[ \sum_{i=1}^{N} \ln \left( \frac{\exp(-\lambda) x_i^\lambda}{\Gamma(\lambda)} \times \lambda^{a-1} \exp(-\theta \lambda) \right) \right] \right\} \lambda^{-a} \exp(-\theta \lambda)
\]

\[
q(\lambda) \propto \left[ \prod_{i=1}^{N} \frac{\exp(-\lambda) x_i^\lambda}{\Gamma(\lambda)} \right] \exp(-\lambda) \exp \left[ E_{q(\theta)} \left[ \theta \right] \right] \lambda^{-a} \exp(-\theta \lambda)
\]

\[
q(\lambda) = \text{Gamma}(\lambda | c', d')
\]

\[
c' = b + a + \sum_{i=1}^{N} x_i
\]
\[ d' = E_q(\theta) \]

The next step is to calculate variation inference objective which is the likelihood \[ L = L(a', b', c', d') \].

**Algorithm 1: Variational Bayes Bootstrap Prior for Subset Selection in Binary Classification**

Define the data as input for \( q(\theta) = \text{Gamma}(\theta|a', b') \) and \( q(\lambda) = \text{Gamma}(\lambda|c', d') \)

Obtain \( a', b', c', d' \) as output.

1. Initialize \( a_0', b_0', c_0', d_0' \)
2. For iteration \( t = 1, ..., T \)
   - Update \( q(\theta) \) by setting;
     \[
     a_t' = b + a + \sum_{i=1}^{N} x_i \\
     b_t' = c_{t-1}' + E_q(\lambda)[-\lambda] \\
     b_t' = c_{t-1}' - \frac{c_{t-1}'}{d_{t-1}'}
     \]
   - Update \( q(\lambda) \) by setting;
     \[
     c_t' = b_{t-1}' + a_{t-1}' + \sum_{i=1}^{N} x_i \\
     d_t' = E_q(\theta)[\theta] \\
     d_t' = \frac{a_{t-1}'}{b_t - 1'}
     \]
3. Evaluate \( L(a', b', c', d') \) to assess convergence.

### 3. Data Calibration

This section presents the application of VB on published real data. Each dataset represents a microarray study of thousands of gene expression profiles (p) measured on (n) individuals. The dataset was extracted from the Gene Expression Omnibus (GEO) database a subsection of National Center Biotechnology Information (NCBI). The data were partitioned using 5-folds cross-validation. One-fifth of each dataset was used to test while the VB approached was developed on the remaining part of the dataset. The datasets description is summarized in Table 1.

| Author of Data | Cancer type     | \( n \) | \( p \) |
|---------------|-----------------|---------|--------|
| [26]          | Colon Cancer    | 62      | 2000   |
| [27]          | Lymphoma Cancer | 58      | 6817   |
| [28]          | Breast Cancer   | 168     | 2905   |
| [29]          | Lung Cancer     | 181     | 12533  |
4. Results

This section presents the results of the train and test data for each of the datasets. Table 2. Presents the results of best subsets after applying the VB algorithm. The best subsets were then later used to perform classification of the disease as shown in the subsequent Tables.

| Cancer type      | p   | x  |
|------------------|-----|----|
| Colon Cancer     | 2000| 3  |
| Lymphoma Cancer  | 6817| 3  |
| Breast Cancer    | 2905| 3  |
| Lung Cancer      | 12533| 2 |

Table 2. Number of Best Subsets of Predictors

| Performance Metrics                              | Sensitivity | Specificity | Positive Predictive Value | Negative Predictive Value | Accuracy | Balance Accuracy | Prevalence | Detection Rate | Area Under the ROC Curve (AUC) |
|--------------------------------------------------|-------------|-------------|---------------------------|---------------------------|----------|-----------------|------------|----------------|-------------------------------|
| Cancer type                                      | Colon       | Lymphoma    | Colon                    | Lymphoma                  |          |                 |            |                |                               |
| Train                                            | 85.18       | 98.72       | 91.55                    | 90.06                     | 90.06    | 35.49           | 30.27      | 90.06          | 90.06                         |
| Test                                             | 84.67       | 96.67       | 91.56                    | 90.00                     | 90.00    | 35.64           | 29.10      | 89.48          | 92.50                         |

Table 3. Performance metrics results (%) for Colon and Lymphoma cancer

Table 2 showed that out of many predictors 2 or 3 of them can guarantee accuracy in the prediction of test data of at least 85%.

| Performance Metrics                              | Sensitivity | Specificity | Positive Predictive Value | Negative Predictive Value | Accuracy | Balance Accuracy | Prevalence | Detection Rate | Area Under the ROC Curve (AUC) |
|--------------------------------------------------|-------------|-------------|---------------------------|---------------------------|----------|-----------------|------------|----------------|-------------------------------|
| Cancer type                                      | Breast      | Lung        | Breast                    | Lung                      |          |                 |            |                |                               |
| Train                                            | 100.00      | 90.00       | 100.00                    | 100.00                    | 100.00   | 51.04           | 51.04      | 100.00         | 100.00                        |
| Test                                             | 89.33       | 99.31       | 86.67                     | 91.43                     | 91.56    | 51.33           | 47.33      | 88.00          | 88.00                         |

Table 4. Performance metrics results (%) for Breast and Lung cancer
| Cancer type      | VB    | RF    |
|-----------------|-------|-------|
| Colon Cancer    | 90.26 | 85.48 |
| Lymphoma Cancer | 94.92 | 89.46 |
| Breast Cancer   | 91.56 | 61.83 |
| Lung Cancer     | 97.98 | 99.44 |
| Average         | 93.68 | 84.05 |

The results in Table 3 and Table 4 showed that overfitting is not a problem with VB procedure since the test data results are not better than the train data results except for prevalence and detection rate which are a function of the sample rather than the algorithm. One of the outstanding results can be observed with Lung cancer data where only 2 subsets of the 12533 gene expression profile taken resulted in almost 98% accuracy. Table 5 showed the side by side comparison with RF, the accuracies of VB were found to be larger than RF except for Lung cancer data.

5. Conclusion

In this paper, a new stage-wise feature selection named VB (Accuracy Based Feature Selection) was developed and its performance in the area of biomarker gene identification was achieved. The performance results revealed high accuracy in both test and train data. A comparative analysis of VB and Random Forest (RF) by [30] established the supremacy of the proposed methods for 75% of the data used.

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