

Sankhyā : The Indian Journal of Statistics
Special Issue on Statistics in Biology and Health Sciences
2007, Volume 69, Part 3, pp. 514-547
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Gene Expression-Based Glioma Classification Using Hierarchical Bayesian Vector Machines

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Abstract

This paper considers several Bayesian classification methods for the analysis of the glioma cancer with microarray data based on reproducing kernel Hilbert space under the multiclass setup. We consider the multinomial logit likelihood as well as the likelihood related to the multiclass Support Vector Machine (SVM) model. It is shown that our proposed Bayesian classification models with multiple shrinkage parameters can produce more accurate classification scheme for the glioma cancer compared to several existing classical methods. We have also proposed a Bayesian variable selection scheme for selecting the differentially expressed genes integrated with our model. This integrated approach improves classifier design by yielding simultaneous gene selection.

AMS (2000) subject classification. Primary 62G08, 62H30, 68T05, 68T10. Keywords and phrases. Gibbs sampling, Markov chain Monte Carlo, Metropolis-Hastings algorithm, microarrays, reproducing kernel Hilbert space, shrinkage parameters, support vector machines.

1 Introduction

There are two main types of brain tumours: those that start in the brain (primary) and those that spread from cancer somewhere else in the body

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(metastasis). Gliomas are the largest group of primary brain tumours. According to the American Brain Tumor Association (http://www.abta.org), primary brain tumours occur at a rate of 14 per 100,000 people. Although people of any age can develop a brain tumour, the problem seems to be most common in children with ages 3 to 12 and in adults with ages 40 to 70. In the United States, approximately 2,200 children younger than age 20 are diagnosed annually with brain tumours (http://www.abta.org/kids/learning /facts.htm). In the past, physicians did not think about brain tumours in elderly people. Due to increased awareness and better brain scanning techniques, people, who are 85 years old and older, are now being diagnosed and treated. The modern clinical practice of neuro-oncology depends on accurate tumour classification. This classification of the tumour type is the basis on which clinicians make critical therapeutic recommendations to the patients. For example, among high-grade gliomas, anaplastic oligodendrogliomas (AO) have a more favourable prognosis than glioblastomas (GM) (Kleihues and Cavenee, 2000). Moreover, though GMs are resistant to most available therapies, AO are often chemosensitive, with approximately two-thirds of cases responding to procarbazine and vincristine (Cairncross et al., 1994). Hence, treatment of brain tumours is dictated by histological diagnosis, and there is a critical need for an objective, clinically relevant method for glioma classification.

Most of the current glioma classifications are derived from the seminal system of Bailey and Cushing (1928). They drew parallels between the histological appearances of glial tumours and putative developmental stages of glia. They reasoned that the cells of astrocytomas microscopically most closely resembled astrocytes and those of oligodendrogliomas mostly mimicked oligodendrocytes. As these tumours become more malignant, they resembled less differentiated precursor cells, hence malignant astrocytomas were dubbed "astroblastomas". It is already confirmed that both at the ultrastructural level and at the immunohistochemical level, many astrocytomas are comprised of cells that exhibit astrocytic differentiation.

Two widely used current histological systems of brain tumour classification are that of the WHO (Kleihues and Cavenee, 2000) and St. Anne-Mayo. Gliomas are classified according to defined histological features that are characteristic of the presumed normal cell of origin. Tumours of classic histology clearly display these features and resemble typical depictions. These cases would be diagnosed similarly by nearly all pathologists. Unfortunately, in several situations, the use of the WHO or St. Anne-Mayo classification system is problematic, primarily because pathological diagnosis remains subjective. Intra-tumoural histological variability is common, and high-grade gliomas can display little cellular differentiation, thereby lacking defining histological features. The diagnosis of tumours with such non-classic histology is controversial. Consequently, diagnosis accuracy and reproducibility are jeopardized, and significant inter-observer variability can occur. Hence, these primary brain tumours have come under intense scientific scrutiny in recent years.

The discovery that cancer is a genetic disease, arising when defects occur in growth-regulatory genes, has revolutionized our understanding of tumourigenesis. Inquiries into the genetic basis of gliomas have yielded large amounts of information about specific genetic events that underlie the formation and progression of human gliomas (Kleihues and Cavenee, 2000). Specific molecular alterations are associated with astrocytic gliomas, and other genetic changes with oligodendrogliomas. However, particular genetic changes may occur in some subtypes of histologically defined astrocytoma or oligodendroglioma. Given the likely biological differences brought about by such genetic variety, each subtype requires a specific and unique set of treatments. For example, clinical testing for chromosome 1p loss in patients diagnosed with anaplastic oligodendrogliomas has been suggested. Testing for 19q loss, CDKN2A/p16 deletion, EGFR gene amplification, and TP53 mutation also provide useful information.

These molecular sub-typing approaches have primarily focused on relatively few but presumably casual tumourigenic events. The advent of expression microarray techniques now allow simultaneous analysis of thousands of genes. There is an increasing interest in changing the basis of tumour classification from morphological to molecular, using microarrays which provide expression measurements for thousands of genes simultaneously (Schena et al., 1995; DeRisi et al., 1997), a key goal being to perform classification via different expression patterns. Several studies using microarrays to profile colon, breast and other tumours have demonstrated the potential power of expression profiling for classification (Alon et al., 1999; Hedenfalk et al., 2001, Mallick et al., 2005). The majority of the methods employed treat binary classification problems. When there are more than two tumour subtypes, as with glioma, multi-class molecular classification is desirable. Multiclass error rates tend to be higher, especially because often there will be a small number of samples in each class.

Support vector machines (SVM) has shown their popularity in microarray literature specially for binary classification problem. Usually there are two ways to handle multicategory problems using SVM: (i) by solving the multicategory problem through a series of binary problems, as suggested by Dietterich and Bakiri (1995), and Allwein, Schapire, and Singer (2000), and (ii) by considering the classes all at once as proposed by Vapnik (1995). Bredensteiner and Bennett (1999), and Crammer and Singer (2001). Constructing pairwise classifier or one-versus-rest classifiers is popular among the first approaches though they have a possible disadvantage of inflating the variance since smaller samples are used to learn each classifier. The second approach is a more algorithmic extension of the binary approach, without much connection with decision rule and uncertainty. Recently, Lee, Lin, and Wahba (2004) proposed a coherent decision theoretic approach for the multicategory support vector machine. It involves a data adaptive tuning criterion for the smoothing parameters using generalized approximate cross validation (GACV) (Wahba et al., 2002).

Recently, Bayesian model based approaches have been used to characterize gene expression pattern for tasks such as gene selection, classification and clustering (Lee et al., 2003; Newton et al., 2001, 2002; Do et al., 2004; Parmigiani et al., 2002; Medvedovic and Sivaganesan, 2002). We develop a fully probabilistic model-based approach, specifically Bayesian multicategory support vector machines based on reproducing kernel Hilbert space (RKHS) (Lee et al., 2004) and also a RKHS based Bayesian multinomial logit model for multicategory classification. We construct a hierarchical model, where the unknown smoothing parameters will be interpreted as shrinkage parameters (Denison et al., 2002). We will assign a prior distribution to these parameters, and obtain the posterior distribution via the Bayesian paradigm. In this way, we obtain not only the point predictors, but find also the associated measures of uncertainty. Furthermore, we will extend the model to incorporate multiple smoothing parameters, leading to significant improvements in the prediction for the example considered.

In many published studies, the number of selected genes is large, (e.g., 495 genes (Khan et al., 1998) and 2000 genes (Alon et al., 1999)). Even in studies that obtained smaller numbers of genes, the numbers are often excessive when compared to the small number of sample points (microarrays) (e.g., 50 genes (Golub et al., 1999) or 96 genes (Khan et al., 2001)). An overly excessive number of genes in conjunction with very few samples is not advisable because it can create an unreliable selection process (Dougherty, 2001).

With a limited sample size, it is common for the expected error to decrease and then increase for increasingly large feature sets. On account of this peaking phenomenon (Hughes, 1968), it is necessary to select a set of features from the full collection of potential features. In principle, there is an optimal number of features for a classification rule, feature-label distribution, and sample size (Hua et al., 2005), but in practice the distribution is unknown. A huge hurdle confronting high-dimensional classification is the combinatorial nature of feature selection; in order to select a subset of kfeatures from a set of n potential features and be assured that it provides an optimal classifier with minimum error among all optimal classifiers for subsets of size k, all k-element subsets must be checked unless there is distributional knowledge that mitigates the search requirement – a condition rarely satisfied in practice (Cover and van Campenhout, 1977). Hence, suboptimal feature-selection methods are required. Suboptimality results not only from the lack of a full search, but also from the need to estimate featureselection criteria from sample data (Sima et al., 2005). Feature selection is often split into two categories. In the *filter* method, features are selected without recourse to classifier design, for instance, by choosing features most correlated with the labels or via mutual information. In the *wrapper* method, features are selected in conjunction with a classifier design. When the number of features is very large, such as in the case of gene expressions on a microarray, the two methods can be used in conjunction. First, one uses filtering, and then one uses some selection method involving classification on the preliminary reduced set. Using a filtering method only involves the danger of selecting many redundant features and also missing features that perform poorly in isolation but work well in combination.

There has been a number of feature-selection procedures proposed in the context of gene expression. Dudoit et al. (2000) have proposed a method for the identification of singly differentially expressed genes by considering a univariate testing problem for each gene and then correcting for multiple testing using adjusted p-values. Tusher et al. (2001) have created *Significance Analysis of Microarray (SAM)*, which assigns a score to each gene on the basis of change in gene expression relative to the standard deviation of repeated measurements. Given genes with scores greater than an adjustable threshold, SAM uses permutations of the repeated measurements to estimate the percentage of genes identified by chance. Hastie et al. (2000) have suggested gene shaving, a new class of clustering methods that tries to identify subsets of genes with coherent expression patterns with large variation across conditions. Kim et al. (2002) have proposed to design analytically

low-dimensional linear classifiers from a probability distribution resulting from spreading the mass of the sample points to make classification more difficult, while maintaining sample geometry. The algorithm is parameterized by the variance of the spreading distribution. By increasing the spread, the algorithm finds gene sets, whose classification accuracy remains strong relative to greater spreading of the sample.

Among Bayesian contributions in gene selection, Ibrahim et al. (2002) proposed a Bayesian univariate selection method, for binary responses only, that primarily models gene expressions of individual genes given disease status. Lee et al. (2004) and Bae et al. (2005) proposed stochastic search algorithms for binary responses. Sha et al. (2004) extended variable selection for multicategory data. Zhou et al. (2004) have proposed a Bayesian approach to nonlinear probit gene selection and gene selection using logistic regressions based on the AIC, BIC, and MDL criteria.

Apart from the methods of Zhou et al. (2004), the usual practice of gene selection for a nonlinear classification model is, first to exploit an ad hoc variable selection method to select the significant genes; and then use these genes in the nonlinear model. As the genes are not selected from the same classification model, there is a chance of possible bias. In this article, we develop a simultaneous variable selection approach (wrapper) with the nonlinear classification model using a unified hierarchical Bayesian model.

We compare our procedure with some highly sophisticated methods like classical support vector machine (CSVM), neural network (NN) and random forest (RF). Almost all of these methods except random forest has an inherent weakness in handling high-dimensional data with limited sample sizes. Hence, we have used the "between square vs. within square" (BWS/BSS) technique proposed by Dudoit et al. (2002) to order the full set of genes and then select only a few from the top to include in the models.

Section 2 describes the materials and the data collected. Section 3 introduces a RKHS-based Bayesian multinomial logit model. Section 4 develops a Bayesian SVM model for multicategory classification. Section 5 provides a Bayesian gene-selection scheme. Section 6 describes how to classify a new sample and identify the differentially expressed genes. Section 7 demonstrates an application of our models on glioma cancer. Section 8 explains the biological importance of the genes selected by our models in glioma cancer, and Section 9 provides some concluding remarks.

2 Materials and Data

All primary glioma tissues came from Brain Tumor Tissue Bank of the University of Texas M.D. Anderson Cancer Center. Tissue bank specimens were frozen shortly after the surgical removal at -80° C. Nothing is known about any possible effect of time delay between tumour removal and tumour freezing on gene expression, but all of the tumour tissue samples were handled in an identical way. Thus, the tumour harvesting procedure should have a similar effect on all samples. H&E-stained frozen tissue sections are routinely prepared from all tissue bank specimens for screening purposes. All tissue specimens for cDNA array analysis were screened by a neuropathologist, and the diagnosis were independently confirmed by a second neuropathologist. The glioma tissue blocks were specifically selected for densest and purest tumour, and they were all comparatively and uniformly "pure". There was minimal contamination by normal brain parenchyma and minimal variation between samples in this regard.

In this study, the gliomas are termed according to the St. Anne-Mayo nomenclature as oligodendroglioma (OL), anaplastic oligodendroglioma (AO), anaplastic astrocytoma (AA) and low grade gliblastoma (GM). Oligodendroglioma (OL) develops from cells called oligodendrocytes that produce the fatty covering of nerve cells. This type of tumour is normally found in the cerebrum, particularly in the frontal or temporal lobes. Anaplastic oligodendroglioma (AO) is a kind of faster growing oligodendroglioma. Anaplastic astrocytoma (AA) is the most common type of glioma and develops from a type of star-shaped cell called an astrocyte. It can occur in most parts of the brain and occasionally in the spinal cord, and glioblastoma multiforme (GM) usually develops in the cerebral hemispheres, more often in the frontal lobes than the temporal lobes or basal ganglia but almost never in the cerebellum.

The cDNA microarray containing fragments representing 597 human genes with known functions and known tight transcriptional controls was used for the experiments. After a high-stringency wash, the hybridization pattern was analysed by autoradiography and quantified by phosphorimaging. So at the end, we have a set of gene expression profile data derived from 25 human glioma surgical tissue samples and expression information on 597 genes (Kim et al., 2002). We have 4 samples in AA, 5 in AO, 6 in OL, and 10 in GM class.

3 RKHS Based Bayesian Multinomial Logit Model

We are interested in classification of glioma cancer with 4 known classes. Hence the response is a categorical variable with more than two categories, and the covariates are the gene expression microarray measurements. Let $\mathbf{t} = (t_1, \ldots, t_n)$ indicates n observed response data, i.e., the type of glioma. This t_i represents one of the J possible categories $(1, \ldots, J)$ J > 2 (here J = 4). Let $p_{ij} = P(t_i = j)$, for $i = 1, \ldots, n$ and $j = 1, \ldots, J$, denote the probability that the *i*th observation falls into the *j*th category. We make an alternate representation of the response t_i by introducing a vector of indicator variables $\mathbf{y}_i \equiv (y_{i1}, \ldots, y_{iJ})^T$, where

$$y_{ij} = \begin{cases} 1 & \text{if } t_i = j, \\ 0 & \text{otherwise;} \end{cases}$$
(3.1)

for i = 1, ..., n and j = 1, ..., J. The multinomial likelihood is

$$f(\boldsymbol{y}|\boldsymbol{p}) \propto \prod_{i=1}^{n} p_{i1}^{y_{i1}} \dots p_{iJ}^{y_{iJ}}.$$
(3.2)

These probabilities are related to a set of p gene expressions (covariates) $X_{n \times p}$ through a logistic link function and a hierarchical model as

$$p_{ij} = \frac{\exp(z_{ij})}{\sum_{k=1}^{J} \exp(z_{ik})}.$$
(3.3)

We relate z_{ij} to $f_j(\boldsymbol{x}_i)$ by $z_{ij} = f_j(\boldsymbol{x}_i) + \epsilon_{ij}$, where ϵ_{ij} is the residual random effect. The f_j 's are unknown functions, which connect the gene expressions with the tumour types. Sha et al. (2004) considered similar multinomial model but they modelled the random latent variable z_{ij} using just a linear function. Taking the negative of the log of the multinomial likelihood (3.2), we get the corresponding loss function

$$L = -\sum_{i=1}^{n} \sum_{j=1}^{J} y_{ij} \log(p_{ij}).$$
(3.4)

This loss function is equivalent to the Kullback-Leibler (KL) directed divergence measure between y_{ij} and p_{ij} , given by

$$L_{KL} = \sum_{i=1}^{n} \sum_{j=1}^{J} y_{ij} \log(y_{ij}/p_{ij}) = \sum_{i=1}^{n} \sum_{j=1}^{J} y_{ij} \log(y_{ij}) - \sum_{i=1}^{n} \sum_{j=1}^{J} y_{ij} \log(p_{ij}).$$
(3.5)

Maximizing the multinomial likelihood (3.2) will be equivalent to minimizing the KL loss function (Bernardo and Smith, 1994). To avoid over-fitting, we can add a penalty function. Thus, casting the whole problem in the regularization framework, we have the following minimization problem

$$\min_{f \in \mathcal{H}_K} \left[\sum_{i=1}^n L_{KL}(y_i, \boldsymbol{f}(\boldsymbol{x}_i)) + \lambda \parallel f \parallel_{\mathcal{H}_K}^2 \right],$$
(3.6)

where $L_{KL}(,)$ is the KL loss function described above, $|| f ||_{\mathcal{H}_K}^2$ is the squared norm penalty functional, λ is the smoothing parameter, and $\mathbf{f} = (f_1, \ldots, f_J)^T$ is the *J*-tuple classification function. We assume that f_j is generated from a reproducing kernel Hilbert space (RKHS) with a positive definite kernel function K(.,.). Then the theory of RKHS as described in Kimeldorf and Wahba (1971), and Wahba et al. (2002) leads to a finite dimensional representation of f_j as

$$f_j(\boldsymbol{x}_i) = \beta_{0j} + \sum_{k=1}^n \beta_{kj} K(\boldsymbol{x}_i, \boldsymbol{x}_k | \boldsymbol{\theta}).$$
(3.7)

The non-linear predictor z_{ij} is thus treated as a random latent variable so that the model is now

$$z_{ij} = \beta_{0j} + \sum_{k=1}^{n} \beta_{kj} K(\boldsymbol{x}_i, \boldsymbol{x}_k | \boldsymbol{\theta}) + \epsilon_{ij} = \boldsymbol{K}_i^T \boldsymbol{\beta}_j + \epsilon_{ij}, \qquad (3.8)$$

where the ϵ_{ij} 's are iid $N(0, \sigma^2)$, $\mathbf{K}_i = (1, K(\mathbf{x}_i, \mathbf{x}_1 | \theta)^T, \dots, K(\mathbf{x}_i, \mathbf{x}_n | \theta))$ and $\boldsymbol{\beta}_j = (\beta_{0j}, \dots, \beta_n j)^T$, $i = 1, \dots, n$. In practice, only the first J - 1 elements of \mathbf{y}_i are used in fitting the RKHS so that the problem is of full rank. So z_{iJ} is set to 0 for all *i* for identifiability of the model. There are several possible choices of kernel functions. In this paper, we use only the Gaussian kernel $\mathbf{K}(\mathbf{x}_i, \mathbf{x}_j | \theta) = \exp\{-\frac{\|\mathbf{x}_i - \mathbf{x}_j\|^2}{\theta}\}.$

We assign hierarchical priors on the unknown parameters as follows.

$$\boldsymbol{\beta}_{j}|\boldsymbol{D}_{j},\sigma^{2} \stackrel{ind}{\sim} \mathrm{N}_{n+1}(\boldsymbol{0},\sigma^{2}\boldsymbol{D}_{j}^{-1}); \ \boldsymbol{D}_{j} = \mathrm{Diag}(\lambda_{0j},\ldots,\lambda_{nj});$$
 (3.9)

$$\sigma^2 \sim \mathrm{IG}(\gamma_1, \gamma_2); \tag{3.10}$$

$$\theta \sim U(a_L, a_U);$$
 (3.11)

$$\lambda_{ij} \stackrel{iid}{\sim} \operatorname{Gamma}(c,d),$$
 (3.12)

where $j = 1, \ldots, J - 1$, $i = 1, \ldots, n$. Denote $\boldsymbol{\lambda} = (\lambda_{01}, \ldots, \lambda_{n1}, \lambda_{02}, \ldots, \lambda_{n2}, \ldots, \lambda_{0(J-1)})^T$, and $\boldsymbol{\beta} = (\boldsymbol{\beta}_1^T, \ldots, \boldsymbol{\beta}_{J-1}^T)^T$, λ_{0j} are fixed at small

values to assign a large variance for the intercept terms β_{0j} , $j = \ldots, J - 1$. Here λ is the vector of smoothing parameters. In the above formulation, we have multiple smoothing parameters λ_{ij} . Often for the sake of simplicity, we assign only one smoothing parameter as $\lambda_{ij} = \lambda$, for all $j = 1, \ldots, J - 1$ and $i = 1, \ldots, n$.

The joint posterior is given by

$$\pi(\boldsymbol{z}, \sigma^{2}, \boldsymbol{\beta}, \boldsymbol{\lambda}, \boldsymbol{\theta} | \boldsymbol{y})$$

$$\propto \prod_{i=1}^{n} p_{i1}^{y_{i1}} \dots p_{iJ}^{y_{iJ}} \times \frac{\exp\{-\frac{1}{2\sigma^{2}} \sum_{i=1}^{n} \sum_{j=1}^{J-1} (z_{ij} - \boldsymbol{K}_{i}^{T} \boldsymbol{\beta}_{j})^{2}\}}{(\sigma^{2})^{n(J-1)/2}}$$

$$\times \frac{\exp\{-\frac{1}{2\sigma^{2}} \sum_{j=1}^{J-1} \boldsymbol{\beta}_{j}^{T} \boldsymbol{D}_{j} \boldsymbol{\beta}_{j}\}}{(\sigma^{2})^{(n+1)(J-1)/2} \prod_{j=1}^{J-1} |\boldsymbol{D}_{j}^{-1}|^{1/2}} \times \exp(-\gamma_{2}/\sigma^{2}) (\sigma^{2})^{-\gamma_{1}-1}$$

$$\times \prod_{i=1}^{n} \prod_{j=1}^{J-1} \exp(-d\lambda_{ij}) (\lambda_{ij})^{c-1}$$
(3.13)

The posterior distribution is intractable; and to generate samples from this posterior, we use MCMC sampling techniques like Gibbs sampling (Gelfand and Smith, 1990) and Metropolis-Hastings (MH) algorithm (Metropolis et al., 1953). To generate samples from the joint posterior, we use the full conditional distributions. The conditional distributions are listed as follows.

(i)
$$\boldsymbol{\beta}_{j}|\boldsymbol{\lambda}, \boldsymbol{z}, \sigma^{2}, \boldsymbol{\theta}, \boldsymbol{y} \sim N_{(n+1)}(\boldsymbol{\mu}_{\beta_{j}}^{*}, \boldsymbol{V}_{\beta_{j}}^{*}),$$

where $\boldsymbol{\mu}_{\beta_{j}}^{*} = \boldsymbol{V}_{\beta_{j}}^{*} \left(\sum_{i=1}^{n} \boldsymbol{K}_{i} z_{ij}\right), \boldsymbol{V}_{\beta_{j}}^{*} = \sigma^{2} \left(\sum_{i=1}^{n} \boldsymbol{K}_{i} \boldsymbol{K}_{i}^{\mathrm{T}} + \boldsymbol{D}_{j}\right)^{-1}$ for $j = 1, \dots, J-1;$

(ii)
$$\sigma^2 | \boldsymbol{\beta}, \boldsymbol{\lambda}, \boldsymbol{z}, \boldsymbol{\theta}, \boldsymbol{y} \stackrel{ind}{\sim} \operatorname{IG}(\gamma_1^*, \gamma_2^*)$$
, where $\gamma_1^* = (J-1)(2n+1)/2 + \gamma_1$ and
 $\gamma_2^* = \frac{\left\{\sum_{i=1}^n \sum_{j=1}^{J-1} (z_{ij} - \boldsymbol{K}_i^T \boldsymbol{\beta}_j)^2\right\} 2 + \left\{\sum_{j=1}^{J-1} \boldsymbol{\beta}_j^T \boldsymbol{D}_j \boldsymbol{\beta}_j\right\}}{2} + \gamma_2;$

- (iii) $\lambda_{ij}|\boldsymbol{\beta}, \boldsymbol{z}, \sigma^2, \theta, \boldsymbol{y} \stackrel{ind}{\sim} \text{Gamma}(c^*, d^*), \ j = 1, \dots, J 1, \ i = 1, \dots, n,$ where $c^* = c + 1/2$ and $d^* = \beta_{ij}^2/2 + d;$
- (iv) $p(\theta|\boldsymbol{\lambda}, \boldsymbol{\beta}, \sigma^2, \boldsymbol{z}, \boldsymbol{y}) \propto \exp\left\{-\frac{1}{2\sigma^2}\sum_{i=1}^n \sum_{j=1}^{J-1} (z_{ij} \boldsymbol{K}_i^T \boldsymbol{\beta}_j)^2\right\} \times I(a_L < \theta < a_U)$, where I() is an indicator variable;

(v)
$$p(\boldsymbol{z}|\boldsymbol{\theta}, \boldsymbol{\lambda}, \boldsymbol{\beta}, \sigma^2, \boldsymbol{y}) \propto \prod_{i=1}^n p_{i1}^{y_{i1}} \dots p_{iJ}^{y_{iJ}} \times \frac{\exp\left\{-\frac{1}{2\sigma^2}\sum_{i=1}^n \sum_{j=1}^{J-1} (z_{ij} - \boldsymbol{K}_i^T \boldsymbol{\beta}_j)^2\right\}}{(\sigma^2)^{n(J-1)/2}}$$

Generation from conditional distributions (i) to (iii) is easy as each represents a standard probability distribution. The conditional for θ given in (iv) does not represent any known distribution. Hence, we device a MH algorithm to sample from it. Let θ be the current state, then draw a candidate value θ^* from its prior U(a_L, a_U). Accept θ^* as a new value of θ with acceptance probability

$$\alpha = \min\left\{1, \frac{p(\theta^*|\boldsymbol{\lambda}, \boldsymbol{\beta}, \sigma^2, \boldsymbol{z}, \boldsymbol{y})}{p(\theta|\boldsymbol{\lambda}, \boldsymbol{\beta}, \sigma^2, \boldsymbol{z}, \boldsymbol{y})}\right\}.$$
(3.14)

The latent variables z_{ij} , i = 1, ..., n, j = 1, ..., J - 1, are sampled using the data augmentation technique suggested by Albert and Chib (1993) as follows.

STEP 1. Let the latent variable $z_i = (z_{i1}, \ldots, z_{iJ-1})$ be a vector corresponding to the *i*th subject.

STEP 2. The relationship between y_{ij} and z_{ij} is as follows.

$$y_{ij} = \begin{cases} 0 & \text{if } \max_{1 \le k \le J-1} \{z_{ik}\} \le 0\\ j & \text{if } \max_{1 \le k \le J-1} \{z_{ik}\} > 0 \text{ and } z_{ij} = \max_{1 \le k \le J-1} \{z_{ik}\}. \end{cases}$$
(3.15)

STEP 3. So $\mathbf{z}_{ij} \sim N(\mathbf{K}_i^T \boldsymbol{\beta}_j, \sigma^2)$ under the above constraint. If the *i*th subject belongs to the *k*th class, this can be simulated by repeated drawing from $N(\mathbf{K}_i^T \boldsymbol{\beta}_j, \sigma^2), j = 1, \ldots, J-1$, and accepting only when the *k*th component of \mathbf{z}_i is the maximum.

We can see that when the number of covariates p is much larger than the sample size n, as in a typical gene expression microarray experiment, using RKHS and Wahba representation, and can reduce the dimension from p to n automatically. Compared to the Bayesian multinomial models proposed by Sha et al. (2004), our RKHS based Bayesian multinomial logit model (BMLM) does not impose a linear model structure for modelling the random latent variable. We rather consider the underlying relationship between the latent variable and the covariates to be an unknown function f, and then use the RKHS theory to estimate that unknown function. This indeed produces a richer class of models than before.

4 Bayesian Multicategory Support Vector Machine

Lee et al. (2004) extended the two class SVM with the hinge loss function to the multicategory setup. In their paper, they generalized the hinge loss function for binary classification to a multivariate loss for the multicategory SVM. As in the previous section, let $\mathbf{t} = (t_1, \ldots, t_n)^T$ be the observed response data, where t_i takes one of the J possible values $1, \ldots, J$. To maintain the symmetry of class label as in the two class SVM, \mathbf{y}_i is coded as

$$y_{ij} = \begin{cases} 1 & \text{if } t_i = j \\ -1/(J-1) & \text{otherwise.} \end{cases}$$
(4.1)

The separating function here will be a *J*-tuple function $f(x) = (f_1(x), \ldots, f_J(x))$. As the sum of the components of the vector y_i is 0, so the *J*-tuple function f(x) will have a zero sum constraint, $\sum_{j=1}^J f_j(x) = 0$. Let $f_j(x) = \beta_{0j} + h_j(x)$, and $h_j(x) \in \mathcal{H}_{K_j}$, for $j = 1, \ldots, J$, where \mathcal{H}_{K_j} denotes a reproducing kernel Hilbert space of functions and h_j denotes any unknown function in that function space. Then $f(x) = (f_1(x), \ldots, f_J(x)) \in \prod_{j=1}^J (\{1\} + \mathcal{H}_{K_j})$, the product space of *J* reproducing kernel Hilbert spaces. Hence, solving the multicategory SVM is equivalent to finding the appropriate f(x) by minimizing the penalized multicategory loss with the zero sum constraint

$$\sum_{i=1}^{n} L(\boldsymbol{y}_{i}) \cdot (\boldsymbol{f}(\boldsymbol{x}_{i}) - \boldsymbol{y}_{i})_{+} + \frac{1}{2}\lambda \sum_{j}^{J} \| h_{j} \|_{\mathcal{H}_{K_{j}}}^{2}, \qquad (4.2)$$

where, if $t_i = j$, then $L(\boldsymbol{y}_i)$ is a *J*-dimensional vector with 0 in the *j*th component and 1 elsewhere. Here, $(\boldsymbol{f}(\boldsymbol{x}_i) - \boldsymbol{y}_i)_+ = [(f_1(\boldsymbol{x}_i) - y_{i1})_+, \dots, (f_J(\boldsymbol{x}_i) - y_{iJ})_+]$, $(x)_+ = \max(0, x)$, and \cdot denotes the Euclidean inner product. Note that $L(\boldsymbol{y}_i) \cdot (\boldsymbol{f}(\boldsymbol{x}_i) - \boldsymbol{y}_i)_+$ is an extension of the hinge loss function (Lee et al., 2004) in a multicategory setup.

Lee et al. (2004) showed that, to find $f(\mathbf{x}) = (f_1(\mathbf{x}), \ldots, f_J(\mathbf{x})) \in \prod_{j=1}^{J} (\{1\} + \mathcal{H}_{K_j})$, with the zero sum constraint, minimizing (4.2) is equivalent to finding $(f_1(\mathbf{x}), \ldots, f_J(\mathbf{x}))$ with the form

$$f_j(\boldsymbol{x}_i) = \beta_{0j} + \sum_{k=1}^n \beta_{kj} K(\boldsymbol{x}_i, \boldsymbol{x}_k | \boldsymbol{\theta}) \text{ for } j = 1, \dots, J$$

$$(4.3)$$

satisfying the constraint $\sum_{j=1}^{J} f_j(\boldsymbol{x}_i) = 0$, for $i = 1, \ldots, n$. If the kernel function K is strictly positive definite, the zero sum constraint over the data points can be replaced by the zero sum constraint on the intercept and the

coefficients as

$$\sum_{j=1}^{J} \beta_{0j} = 0; (4.4)$$

$$\sum_{j=1}^{J} \beta_{kj} = 0, \text{ for all } k = 1, \dots, n.$$
 (4.5)

Viewing the loss as the negative of the log likelihood, we construct our Bayesian multicategory SVM (BMSVM). The conditional distribution of y_i given the latent variable z_i is given as

$$p(\boldsymbol{y}_i|\boldsymbol{z}_i) \propto \exp\left\{-L(\boldsymbol{y}_i).(\boldsymbol{z}_i-\boldsymbol{y}_i)_+\right\},\tag{4.6}$$

where the z_i 's is the random J component latent vector, and the y_i 's are conditionally independent of the z_i 's. As in the previous section, the latent vector z_i is connected to the unknown separating functions $f(x) = (f_1(x), \ldots, f_J(x))$ as

$$\boldsymbol{z}_i = \boldsymbol{f}(\boldsymbol{x}_i) + \boldsymbol{\epsilon}_i, \tag{4.7}$$

where $\boldsymbol{\epsilon}_i = (\epsilon_{i1}, \ldots, \epsilon_{iJ})^T$ is the residual random vector, and $\boldsymbol{\epsilon}_i \sim N_J(0, \sigma^2 \boldsymbol{I}_J)$. Assuming that $\boldsymbol{f}(\boldsymbol{x})$ belongs to a product space of J RKHS and with a strictly positive definite kernel K, from (4.3) and (4.7) we can write

$$z_{ij} = \beta_{0j} + \sum_{k=1}^{n} \beta_{kj} K(\boldsymbol{x}_i, \boldsymbol{x}_k | \boldsymbol{\theta}) + \epsilon_{ij} = \mathbf{K}'_i \boldsymbol{\beta}_j + \epsilon_{ij}$$
(4.8)

with the restrictions given in (4.4) and (4.5). We put hierarchical priors on the unknown intercepts and the regression coefficients as follows.

$$\boldsymbol{\beta}_{j}|\boldsymbol{D}_{j},\sigma^{2} \sim N_{n+1}(\boldsymbol{0},\sigma^{2}\boldsymbol{D}_{j}^{-1});\boldsymbol{D}_{j} = \text{Diag}(\lambda_{0j},\ldots,\lambda_{nj})$$
 (4.9)

with the constraints given in (4.4) and (4.5);

$$\sigma^2 \sim \mathrm{IG}(\gamma_1, \gamma_2); \tag{4.10}$$

$$\theta \sim U(a_L, a_U);$$
 (4.11)

$$\lambda_{ij} \stackrel{iid}{\sim} \operatorname{Gamma}(c,d), \text{ where } j = 1, \dots, J, \ i = 1, \dots, n.$$
 (4.12)

We notice that unlike what happens in the case of the multinomial logit model, here the β_j 's are not independent. The dependence is due to the zero

sum constraints imposed on them. The joint posterior is given by

$$\pi(\boldsymbol{z}, \sigma^{2}, \boldsymbol{\beta}, \boldsymbol{\lambda}, \boldsymbol{\theta} | \mathbf{y})$$

$$\propto \exp\left\{-\sum_{i=1}^{n} L(\boldsymbol{y}_{i}).(\boldsymbol{z}_{i} - \boldsymbol{y}_{i})_{+}\right\} \times \frac{\exp\left\{-\frac{1}{2\sigma^{2}}\sum_{i=1}^{n}\sum_{j=1}^{J-1}(z_{ij} - \boldsymbol{K}_{i}^{T}\boldsymbol{\beta}_{j})^{2}\right\}}{(\sigma^{2})^{nJ/2}}$$

$$\times \frac{\exp\left\{-\frac{1}{2\sigma^{2}}\sum_{j=1}^{J}\boldsymbol{\beta}_{j}^{T}\boldsymbol{D}_{j}\boldsymbol{\beta}_{j}\right\}}{(\sigma^{2})^{(n+1)J/2}\prod_{j=1}^{J-1}|\boldsymbol{D}_{j}^{-1}|^{1/2}} \times I\left(\sum_{j=1}^{J}\boldsymbol{\beta}_{kj} = 0, k = 0, \dots, n\right)$$

$$\times \exp\left(-\gamma_{2}/\sigma^{2}\right)(\sigma^{2})^{-\gamma_{1}-1} \times \prod_{i=1}^{n}\prod_{j=1}^{J}\exp\left(-d\lambda_{ij}\right)(\lambda_{ij})^{c-1}.$$
(4.13)

Comparing the posteriors (3.13) and (4.13), we can see the main difference is in the likelihood. In the Bayesian multinomial logit model, we have the multinomial likelihood, whereas in the Bayesian multicategory SVM, we have the likelihood corresponding to the multivariate hinge loss. Also in (4.13), zero sum constraint on the regression coefficients is imposed using the indicator function $I(\sum_{j=1}^{J} \beta_{kj} = 0, k = 0, ..., n)$. Hence the posterior in (4.13) has a truncated support. The conditional distributions of σ^2 , λ_{ij} , and θ are the same as those in (ii), (iii), and (iv) in the earlier section. The conditional posterior of β_j , j = 1, ..., J, will follow multivariate normal distribution as in (i) of Section 3, but with the zero sum constraint imposed by (4.4) and (4.5). As the underlying likelihood is different in the multicategory SVM, the conditional distribution (v) is now changed to $p(\boldsymbol{z}|\theta, \boldsymbol{\lambda}, \boldsymbol{\beta}, \sigma^2, \boldsymbol{y}) \propto \exp\left\{-\sum_{i=1}^{n} L(\boldsymbol{y}_i).(\boldsymbol{z}_i - \boldsymbol{y}_i)_+\right\} \times \exp\{-\frac{1}{2\sigma^2} \sum_{i=1}^{n} \sum_{j=1}^{J} (z_{ij} - \boldsymbol{K}_i^T \boldsymbol{\beta}_j)^2\}$. We can generate samples from the posterior (4.13) by the following steps.

- STEP 1. Generate λ_{ij} , σ^2 and θ in the same way as in the Bayesian multinomial logit model of the previous section.
- STEP 2. Generate β_j , j = 1, ..., J from the multivariate normal distribution satisfying the constraints (4.4) and (4.5).
- STEP 3. The conditional distribution of the latent vector \boldsymbol{z}_i is not standard; so we update it by blocks of \boldsymbol{z}_i as suggested by Roberts and Sahu (1997). When \boldsymbol{z}_i is in the present state, draw a candidate value \boldsymbol{z}_i^* from $N(\boldsymbol{K}_i^0\boldsymbol{\beta},\sigma^2 I_J)$, where $\boldsymbol{K}_i^0 = I_q \bigotimes \boldsymbol{K}_i^{\mathrm{T}}, \boldsymbol{\beta} = (\boldsymbol{\beta}_1^{\mathrm{T}},\ldots,\boldsymbol{\beta}_q^{\mathrm{T}})^{\mathrm{T}}$. Accept the update with the acceptance probability

$$\alpha_{i} = \min\left\{1, \frac{\exp\left\{-\sum_{i=1}^{n} L(\boldsymbol{y}_{i}).(\boldsymbol{z}_{i}^{*}-\boldsymbol{y}_{i})_{+}\right\}}{\exp\left\{-\sum_{i=1}^{n} L(\boldsymbol{y}_{i}).(\boldsymbol{z}_{i}-\boldsymbol{y}_{i})_{+}\right\}}\right\}.$$

As in our previous model, the RKHS gives us a low n-dimensional representation for a much higher p-dimensional problem, without making an additional projection on the feature space.

5 A Bayesian Gene Selection Scheme

The two models proposed in the previous two sections have an in-built dimension reduction technique, where a successful dimension reduction is made via RKHS. Consequently, instead of dealing with p covariates, we deal with n different kernel functions. In a typical microarray experiment, we have gene expression data on 5000 - 10,000 genes for less than 100 tumour samples. Many genes do not contain information that is useful for determining the differences between the samples. These genes should not be used for classification: indeed, sometimes they may even contain noise that can lead to incorrect classification. Although the RKHS formulation enables us to keep all the genes in our model, yet an improved classification can be obtained if only differentially expressed genes are included in the model. A simple method as proposed by Dudoit et al. (2002) may be used to rank the full set of genes and select the top few genes or the genes with the most marginal relevance. But their method does not consider the possible interaction effect between several genes. In this section, we propose a Bayesian variable selection technique (George and McCulloch, 1993) for our models in Sections 3 and 4.

Let $X_{n \times p}$ be a matrix of gene expression data, where each column represents a gene and each row represents a sample. To do the gene selection, introduce $\boldsymbol{\gamma} = (\gamma_1, \ldots, \gamma_p)^T$, a $p \times 1$ vector of indicators, such that

$$\gamma_k = \begin{cases} 0 & \text{the } i\text{th gene is not selected,} \\ 1 & \text{the } i\text{th gene is selected.} \end{cases}$$
(5.1)

For a particular combination of genes or the choice of γ , X_{γ} denotes the reduced gene expression matrix with only those columns of the full gene expression matrix X, which correspond to those the elements of γ that are equal to one. Hence the dimension of X_{γ} is $n \times p_{\gamma}$, where $p_{\gamma} = \sum_{k=1}^{p} \gamma_k$, or the number of nonzero components in the vector γ .

We put independent Bernoulli prior on γ_k as follows.

$$\gamma_k \stackrel{iid}{\sim} \text{Bernoulli}(\omega), \quad i = 1, \dots, p.$$
 (5.2)

The value of ω is chosen to be small to restrict the number of genes in the model. We can also include the prior knowledge of some of the genes, which are more important than others by assigning different ω_k for different γ_k .

The inclusion of γ , the indicator vector, will result in the computation of the kernel function K on the basis of the reduced expression matrix $X\gamma$ and is denoted by $K\gamma(,)$. The posterior (3.13) from the RKHS based Bayesian multinomial logit model will now change to

$$\pi(\boldsymbol{z}, \sigma^{2}, \boldsymbol{\beta}, \boldsymbol{\lambda}, \boldsymbol{\theta}, \boldsymbol{\gamma} | \boldsymbol{y})$$

$$\propto \prod_{i=1}^{n} p_{i1}^{y_{i1}} \dots p_{iJ}^{y_{iJ}} \times \frac{\exp\{-\frac{1}{2\sigma^{2}} \sum_{i=1}^{n} \sum_{j=1}^{J-1} (z_{ij} - \boldsymbol{K}_{\boldsymbol{\gamma}_{i}}^{T} \boldsymbol{\beta}_{j})^{2}\}}{(\sigma^{2})^{n(J-1)/2}}$$

$$\times \frac{\exp\{-\frac{1}{2\sigma^{2}} \sum_{j=1}^{J-1} \boldsymbol{\beta}_{j}^{T} \boldsymbol{D}_{j} \boldsymbol{\beta}_{j}\}}{(\sigma^{2})^{(n+1)(J-1)/2} \prod_{j=1}^{J-1} |\boldsymbol{D}_{j}^{-1}|^{1/2}} \times \exp(-\gamma_{2}/\sigma^{2}) (\sigma^{2})^{-\gamma_{1}-1}}$$

$$\times \prod_{i=1}^{n} \prod_{j=1}^{J-1} \exp(-d\lambda_{ij}) (\lambda_{ij})^{c-1} \times \prod_{k=1}^{p} \omega^{\gamma_{k}} (1-\omega)^{1-\gamma_{k}}.$$
(5.3)

In order to generate samples from (5.3), in addition to all previous steps for sampling $\boldsymbol{z}, \sigma^2, \boldsymbol{\beta}, \boldsymbol{\lambda}$, and $\boldsymbol{\theta}$, an additional step is required to sample $\boldsymbol{\gamma}$. We use the conditional distribution

(vi)
$$p(\boldsymbol{\gamma}|\boldsymbol{z}, \sigma^2, \boldsymbol{\beta}, \boldsymbol{\lambda}, \boldsymbol{\theta}, \mathbf{y}) \propto \exp\left\{-\frac{1}{2\sigma^2} \sum_{i=1}^n \sum_{j=1}^{J-1} (z_{ij} - \boldsymbol{K}_{\boldsymbol{\gamma}_i}^T \boldsymbol{\beta}_j)^2\right\} \times \prod_{k=1}^p \omega^{\gamma_k} (1-\omega)^{1-\gamma_k}.$$

The above conditional distribution is not of a standard form. Hence we again deploy a MH algorithm where the components of the new update γ^* , γ_i^* can be drawn from the Bernoulli(ω) distribution independently. The new γ^* is accepted with probability

$$\alpha = \min\left\{1, \frac{\exp\{-\frac{1}{2\sigma^2}\sum_{i=1}^n \sum_{j=1}^{J-1} (z_{ij} - \boldsymbol{K}_{\gamma_i}^T \boldsymbol{\beta}_j)^2\}}{\exp\{-\frac{1}{2\sigma^2}\sum_{i=1}^n \sum_{j=1}^{J-1} (z_{ij} - \boldsymbol{K}_{\gamma_i}^T \boldsymbol{\beta}_j)^2\}}\right\}.$$

Similarly, we can also incorporate the Bayesian model selection technique in our Bayesian multicategory support vector machine model. The use of the indicator vector γ will only change the kernel matrix. The other part of the BMSVM model as explained in Section 4 remains same. The posterior (5.2) from the Bayesian multicategory SVM model will now take the form

 $\pi(\boldsymbol{z}, \sigma^{2}, \boldsymbol{\beta}, \boldsymbol{\lambda}, \boldsymbol{\theta}, \boldsymbol{\gamma} | \boldsymbol{y})$ $\propto \exp\left\{-\sum_{i=1}^{n} L(\boldsymbol{y}_{i}).(\boldsymbol{z}_{i} - \boldsymbol{y}_{i})_{+}\right\} \times \frac{\exp\left\{-\frac{1}{2\sigma^{2}}\sum_{i=1}^{n}\sum_{j=1}^{J}(z_{ij} - \boldsymbol{K}_{\boldsymbol{\gamma}i}^{T}\boldsymbol{\beta}_{j})^{2}\right\}}{(\sigma^{2})^{nJ/2}}$ $\times \frac{\exp\{-\frac{1}{2\sigma^{2}}\sum_{j=1}^{J}\boldsymbol{\beta}_{j}^{T}\boldsymbol{D}_{j}\boldsymbol{\beta}_{j}\}}{(\sigma^{2})^{(n+1)J/2}\prod_{j=1}^{J}|\boldsymbol{D}_{j}^{-1}|^{1/2}} \times I\left(\sum_{j=1}^{J}\boldsymbol{\beta}_{kj}=0, k=0,\ldots,n\right)$ $\times \exp\left(-\gamma_{2}/\sigma^{2}\right)(\sigma^{2})^{-\gamma_{1}-1} \times \prod_{i=1}^{n}\prod_{j=1}^{J}\exp\left(-d\lambda_{ij}\right)(\lambda_{ij})^{c-1}$ $\times \prod_{k=1}^{p}\omega^{\gamma_{k}}(1-\omega)^{1-\gamma_{k}}.$ (5.4)

As in Section 4, here too we will follow exactly same steps to generate samples from the joint posterior (5.4). The extra step needed to sample γ , the vector of indicators, is incorporated by sampling from the conditional posterior

(vi)
$$p(\boldsymbol{\gamma}|\boldsymbol{z}, \sigma^2, \boldsymbol{\beta}, \boldsymbol{\lambda}, \boldsymbol{\theta}, \mathbf{y}) \propto \exp\left\{-\frac{1}{2\sigma^2} \sum_{i=1}^n \sum_{j=1}^J (z_{ij} - \boldsymbol{K}_{\boldsymbol{\gamma}_i}^T \boldsymbol{\beta}_j)^2\right\} \times \prod_{k=1}^p \omega^{\gamma_k} (1-\omega)^{1-\gamma_k}.$$

From the conditional posterior distribution of γ it is clear that although the priors of the components γ_i , $i = 1, \ldots, p$ are assumed to be i.i.d. Bernoulli(ω), the posterior is not independent. Hence, we can establish some dependency among the genes, which was not possible to establish by the Dudoit et al. (2002) criterion.

6 Classification of Future Cases and Identifying the Significant Genes

When we have J different classes of cancer tumours, and J > 2. For a new sample, whose corresponding gene expression measurement is denoted by \boldsymbol{x}_{new} , the classification rule is induced by

$$\phi(\boldsymbol{x}_{new}) = \arg \max_{j} p(t_{new} = j | \boldsymbol{x}_{new}, \boldsymbol{t}_{old}), \qquad (6.1)$$

where

$$p(t_{new}=j|\boldsymbol{x}_{new},\boldsymbol{t}_{old}) = \int_{\Theta} p(t_{new}=j|\boldsymbol{x}_{new},\boldsymbol{t}_{old},\Theta)\pi\left(\Theta|\boldsymbol{t}_{old}\right)d\Theta; \quad j=1,\ldots,J$$
(6.2)

is the posterior predictive probability that the tumour belongs to the *j*th class and $\Theta = (\boldsymbol{z}, \sigma^2, \boldsymbol{\beta}, \boldsymbol{\lambda}, \boldsymbol{\theta}, \boldsymbol{\gamma})$, the set of all parameters in the model. A Monte Carlo estimate of (6.2) is given by

$$\hat{p}(t_{new} = j | \boldsymbol{x}_{new}, \boldsymbol{t}_{old}) = \sum_{t=1}^{B} \hat{p}(t_{new} = j | \boldsymbol{x}_{new}, \boldsymbol{t}_{old}, \Theta^{t}); \quad j = 1, \dots, J, \quad (6.3)$$

where Θ^t is t-th MCMC sample from the posterior, and B is the total number of Monte Carlo samples used for estimation after the initial burn in. Hence for the new tissue sample, we compute its posterior predictive probability of being in class j for all the classes $j = 1, \ldots, J$, and finally assign the new sample to that class for which this probability is maximum.

For identifying the significant genes from the dormant ones, we run the MCMC chain for a long time, and after discarding sufficient samples to account for the burn in period, we calculate the relative number of times each gene appeared in the sample. This will serve as an estimate of the posterior probability that a single gene is included in the model, and can be used as a measure for identifying the differentially expressed genes from the inactive ones. Since we assume that only the differentially expressed genes are responsible for the variation in types of tumours, those should be included in our model more frequently to maximize the posterior distribution. It is to be noted now that instead of a two step procedure of first gene selection and then classification, our model can simultaneously select important genes and do the classification.

7 Analysis of the Glioma Data

In this section, we have modelled the glioma cancer data using our BMLM and BMSVM models. The gliomas are termed according to the St. Anne-Mayo nomenclature as low grade OL, AO, AA and GM, so J = 4. We have in total 597 genes, and tissue samples are collected from 25 patients. As the number of patients is very small, we cannot split the data further into training and test sets and evaluate the performance of our classifier on the test set. Leave one out cross-validation technique is often not accurate and suffers badly from high outliers (Braga-Neto and Dougherty, 2004). To

evaluate the performance of our classifiers, we use .632 bootstrap technique (Efron, 1983). We draw a sample of n observations from our data of n observations with replacement, and make this the training set. The sample, which are left out or not included in the training set, are kept as the test set. We denote the proportion of samples classified incorrectly in the test set by e_{test_i} and the proportion of samples classified incorrectly in the training set by e_{train_i} . We do this kind of bootstrapping B times. At the end, the average gives us the bootstrap estimate of the error. The Bootstrap estimate of the misclassification error given by the formula

$$\frac{1}{B} \sum_{i=1}^{B} \left(0.632 e_{test_i} + 0.368 e_{train_i} \right). \tag{7.1}$$

We have fixed B = 5000 in our case.

The Bayesian gene selection criterion as developed in Section 5, and also the gene ordering criterion as suggested by Dudoit et al. (2002), are used to include only the important genes. For both examples, we have also considered three standard nonlinear classification models: (i) Neural Network (NN), (ii) Random Forest (RF) and (iii) Classical Support Vector Machine (CSVM). None of the models, except the random forest, is equipped to deal with such a high dimensional problem. In each case, we use the BWS/BSScriterion to order the genes and then select the top few genes for each of our models. Next, we fit our (iv) RKHS based Bayesian multinomial logit model (BMLM) and (v) Bayesian multicategory SVM (BMSVM). For each method, we make an initial gene selection according to the BWS/BSS criterion before running our model. Lastly, we fit our (vi) Bayesian RKHS based multinomial logit model integrated with Bayesian gene selection technique (Section 6) (BMLM + BGS) and (vii) Bayesian multicategory SVM integrated with Bayesian gene selection technique (BMSVM + BGS). Our models (vi) and (vii) are different from (iv) and (v) in the sense that in our last two models, we can simultaneously predict the class label and the differentially expressed genes in a full Bayesian setup. In contrast, in (iv) and (v), the gene selection was made on the basis of BWS/BSS, which is a strong frequentist idea similar to an F-statistic, while the classification is made on the basis of our Bayesian models.

Choice of priors plays an important role in our analysis, and we have used near-diffused proper priors. The near diffuseness guarantees that our prior choice is as flat as possible but proper, which ensures the propriety of the posterior along with the objectivity of our analysis. We followed the

following combination of hyperparameters: $\gamma_1 = 1$, $\gamma_2 = 10$, $c = 10^{-8}$, $d = 10^{-5}$, $a_L = 0$ and $a_U = 100$. This choice of hyperparameters is also suggested by Mallick et al. (2005), and produces a near-diffused but proper prior. We used the Gaussian kernel as we found empirically that it produces better classification result than the polynomial kernel. We chose $\omega = 0.05$ so that on an average only top 5% of the genes are to be included in the models producing sparsity. For all our models (models (iv) to (vii)), we have tried both multiple smoothing parameters and single smoothing parameters. The results obtained with single smoothing parameters are denoted by "*".

We run a MCMC chain 200,000 times and discard the first half as the burn in. To ensure that we are not stuck in one of the many modes of the posterior distribution, we use multiple chains with different starting values. In both models, we used a total of 5 independent chains with widely different starting points and our final prediction is based on the pooled samples from these 5 chains. We have used neural network models with 20 hidden nodes. After 20 hidden nodes, we do not gain anything in terms of prediction accuracy compared to the cost of computational complexity. The *nnet* function in R with *softmax* option is used to fit the neural network model. For the random forest, we have used the *randomForest* function in R with 10000 boosted trees and all other default parameters.

Method		Top Genes			
	20	50	100	597	
NN CSVM RF	$0.1213 \\ 0.0909 \\ 0.1489$	$\begin{array}{c} 0.1338 \ 0.1554 \ 0.1691 \end{array}$	$\begin{array}{c} 0.1612 \\ 0.1921 \\ 0.1801 \end{array}$	- 0.3672 0.2839	
BMLM BMSVM BMLM* BMSVM*	$0.0702 \\ 0.0722 \\ 0.1258 \\ 0.1346$	$0.0802 \\ 0.0794 \\ 0.1428 \\ 0.1302$	0.0988 0.1002 0.2004 0.1444	$\begin{array}{c} 0.2138 \\ 0.2714 \\ 0.3814 \\ 0.3532 \end{array}$	

TABLE 1. BOOTSTRAP ERROR RATE OF MISCLASSIFICATION IN THE GLIOMA CANCER DATA. GENES SELECTED BY THE BWS/BSS criterion.

Table 1 reports the total .632 bootstrap estimate of the misclassification error using the formula (7.1). Initially, we used the BWS/BSS criterion as proposed by Dudoit et al. (2002) to order the genes. After we order the genes, we select the top 20, 50, 100 and 597 (i.e., all genes without any selection) genes and use them in the models to predict the class of the "out of bag" sample. The first row indicates the number of top genes included in

the model. As we have pointed out already that the classification of Glioma cancer is extremely difficult, we observe that most of the standard methods like neural network and random forest do equally poorly in classification. With top 20 genes selected by the BWS/BSS criterion, neural network and random forest give bootstrap misclassification errors as 0.1213 and 0.1489 respectively. As we went on adding genes to it, the performance of random forest decayed drastically. With 597 genes, the bootstrap error is nearly the double of what we obtained with top 20 genes. Neural network is much more stable in performance when more genes are added. But again it is impossible to fit a neural network with all the 597 genes. The CSVM worked very well with the top 20 genes, it gave 0.0909 bootstrap error, which is much lower than both the neural network and the random forest. But again, here also by increasing the number of genes to 50 and with any addition thereafter, the performance of the CSVM model diminishes drastically. Our BMLM and BMSVM models (the ones with multiple smoothing parameters) give much better results than all the previous standard models. With the top 20 genes, the bootstrap error is reduced by 20% to 50% in both of our models when compared to the three standard ones. Also, we see that as we increase the number of genes to 100 in our models, their performance is not as affected as the RF, CSVM and NN. But inclusion of all 597 genes gives very high error estimates for all the models, although BMLM and BMSVM continue to improve on CSVM and RF. A main limitation of our BMLM and BMSVM models is that they rely on a two step procedure of gene selection and subsequent class prediction. The main problem in the BWS/BSS criterion is that we do not know exactly how many top genes we should use, so that we keep on adding more genes in the models.

Methods	CV Error	No. of Genes Selected
$BMLM + BGS$ $BMSVM + BGS$ $PMLM^* + BCS$	0.0540 0.0596 0.0680	23 25 27
$BMSVM^* + BGS$	0.0652	23

TABLE 2. BOOTSTRAP ERROR RATE OF MISCLASSIFICATION GLIOMA CANCER DATA. Genes are adaptively selected by the models.

In Table 2, we report the results of our models BMLM+BGS and BMSVM +BGS, where we adaptively select the important genes and make the class predictions simultaneously. Both BMLM+BGS and BMSVM+BGS models reduce the bootstrap error by more than 20% than the BMLM and the BMSVM respectively. In BMLM+BGS model, on an average 23 genes are included, and in the BMSVM+BGS, on an average 25 genes are selected.



Figure 1. Glioma DNA data. (a) Marginal relevance of Genes by *BWS/BSS* criterion. (b) Relative number of times a gene is selected by our BMLM+BGS model. (c) Relative number of times a gene is selected by our BMSVM+BGS model.

Figure 1 plots the marginal relevance of each gene by the BWS/BSS criterion and also the relative number of times each gene appeared in our BMLM+BGS and BMSVM+BGS models. From the figure, we can see that there is an overlap of active genes as suggested by the BWS/BSS criterion and our Bayesian variable selection approach. In fact, there are 8 common important genes selected by both methods. Kim et al. (2002) developed an algorithm for identification of gene sets important genes, which are detected by both of our models and also found to be relevant by the algorithm of Kim et al. (2002). From Table 3, we get that 11 genes are found to match with

Kim's algorithm, while between the 23 genes selected by BMLM+BGS and the 25 genes selected by BMSVM+BGS, there is a match for 14 such genes. A heat map of the genes selected by our two models and the BWS/BSS criterion is also provided in Figure 2.

Three genes, namely cyclin-dependent kinase inhibitor 1C (CDKN1C), G2/mitotic-specific cyclin B1 (CCNB1) and cell division protein kinase 7 (CDK7), are marked as important by both models but are not selected by Kim's algorithm or BWS/BSS criterion. In Figure 3, we show the heat map of these three genes, and it appears that they might be important. It is well-known that inducible expression of CDKN1C in cell lines deficient in this cyclin-dependent kinase inhibitor reduces the motility and the invasiveness of malignant gliomas (Sakai et al., 2004). Presence of CCNB1 usually shows an increased growth rate of malignant glioma cell lines and plays a significant role in glioma tumorigenes (Weber et al., 2000). A further investigation may throw more light on the role of these three genes for glioma cancer.

From Tables 1 and 2, we see that we had a significant advantage of using multiple smoothing parameters over single smoothing parameters in terms of bootstrap misclassification error. In all the models with single smoothing parameter, we get higher bootstrap error than the corresponding model with multiple smoothing parameters.

TABLE 3. NAMES OF THE GENES SELECTED BY BOTH OF OUR MODELS BMLM+BGS AND BMSVM+BGS. THE GENES, WHICH ARE ALSO IDENTIFIED TO BE RELEVANT BY THE KIM ET AL. (2002) ALGORITHM, ARE MARKED BY A '*'.

Name of the gene	GeneBank access no.
* angiopoietin 1 receptor: TIE-2	L06139
* insulin-like growth factor-binding protein 2 (IGFBP2)	M35410
* FLT1: VEGFR1	X51602: U01134
* cell surface glycoprotein MUC18:	M28882
cyclin-dependent kinase inhibitor 1C (CDKN1C)	U22398
G2/mitotic-specific cyclin B1 (CCNB1)	M25753
STK1: CDK7	L20320
* guanine nucleotide-binding protein beta subunit 2	M36429
* guanine nucleotide-binding protein beta 1 subunit	
(GNB1)	M36430
* mitogen-activated protein kinase 10	
(MAP kinase 10; MAPK10; PRKM10)	U34819: U07620
* growth factor receptor-bound protein 2 (GRB2)	L29511; M96995
* tumour necrosis factor superfamily member 5 (TNFSF5)	L07414
* muscle-specific DNase I-like	
(DNase1L1; DNL1L); DNase X	X90392; L40817; U06846
* thymosin beta 10 (TMSB10; THYB10); PTMB10	M92381

criterion. zontal axis on the top, we have the Gene Bank Accession No. (c) Heatmap of 25 top genes selected by our BMSVM+BGS model. Figure 2. Glioma DNA data. (b) Heatmap of 23 top genes selected by our BMLM+BGS model. (a) Heatmap of 25 top genes selected by BWS/BSSOn the hori-







Samples





Figure 3. Heatmap of the 3 genes marked as significant by our two models but missed by the Kim's algorithm and the BWS/BSS criterion.

8 Biological Significance of the Selected Genes in Glioma Cancer

Our BMLM+BGS and BMSVM models have selected 23 genes and 25 genes as relevant respectively based on a marginal posterior probability of 0.01. There is a significant overlap in the gene lists between the two models. In total, 14 genes are found to be common in both the models and are listed in Table 3.

Most of the selected genes carry a lot of biological significance and have putative roles in cancer biology. We focus on potential roles for a few of them. For example, Angiopoietin-2 (Ang2) induces human glioma invasion through the activation of matrix metalloprotease-2 and plays an important role in angiogenesis and tumour progression. Ang2 induces human glioma cell invasion. In invasive areas of primary human glioma specimens, upregulated expression of Ang2 was detected in tumour cells. Correspondingly higher levels of MMP-2 expression were present in Ang2-expressing tumour cells in these glioma, (Hu et al., 2003).

Another molecule that appears in the list is insulin-like growth factor binding protein 2 (IGFBP2). Wang et al. (2003) showed that IGFBP2 contributes to glioma progression in part by enhancing MMP-2 gene transcription and in turn tumour cell invasion.

There is also speculation that progression to glioma requires activation of angiogenesis and has stimulated significant efforts in the development of agents that will block this process. In particular, two pathways have received considerable attention. They are vascular endothelial growth factor (VEGF) and its receptors, VEGFR1 (Flt-1) and VEGFR2 (Flk-1); note that VEGFR appears on our list of genes in Table 3. VEGF has been shown to be critical for the earliest stages of vasculogenesis, promoting endothelial cell proliferation, differentiation, migration and tubular formation. Gene targeting studies have shown that deficiency of VEGF, Flt-1 or Flk-1 results in early embryonic lethality caused by defects in angiogenesis and vasculogenesis (Elizabeth et al., 2001).

Another gene from the list is MUC18, which is a cell adhesion molecule. It has been observed that increased levels of MUC18 results in an increased potential of the cell to grow and divide uncontrollably and thus spreading the glioma (Heimberger et al., 2005). The GNB1 gene included in both of our models is also considered by the biologists as one of the candidate genes for glioma tumour suppressor gene (Collins, 2004).

Some recent experiments support that extracellular signal regulated kinase (ERK), a mitogen activated protein kinase, MAPK2 might have a critical role in cell proliferation (Bhaskara et al., 2005). FACS analysis and immunofluorescence studies using monoclonal antibodies, which specifically recognized EGFR, demonstrated that EGFR was expressed predominantly at the cell surface, similar to wild-type (wt) EGFR and to the expression seen in primary biopsy-derived glioma cells (Cavenee, 2002). Phosphotyrosine residues in the carboxyl tail of wtEGFR provide sites for interaction with SRC homology 2 (SH2) domain-containing adaptor molecules such as SRC and GRB2. Immunoprecipitation studies have shown that EGFR is constitutively associated with phosphorylated SHC and GRB2 in several cell lines of different origins. This suggests that the low level constitutive activation of EGFR may cause coupling into unique pathways in these cells and may also point out an entry into interference therapies targeted at gliomas.

The tumour necrosis factor (TNF) superfamily member 5 is also selected by our model. At present, the anti-tumour activity of human recombinant TNF is being examined against various malignant tumours of human origin (Wakabayashi et al., 1997). In the study by Sawada et al. (2004), they reported the anti-tumour activity of recombinant human TNF against human malignant glioma cell lines in vitro and in vivo.

The three genes like CDKN1C, CCNB1 and cell CDK7, which are missed by Kim's algorithm but captured by us, also carries direct biological significance in glioma cancer as mentioned in the previous section.

9 Discussion

Gliomas are very complex cancers involving different growth characteristics and cell lineage features (Kleihuse and Cavenee, 2000). As the original clone of tumour cells may exist at any stage of cell differentiation and may have different transformation events, the boundaries between tumour grades and tumour lineages can be blurred. This is reflected in morphologically based tumour classification schemes that often mix cell lineage features with tumour growth characteristics. The results are subjective, and disagreement among pathologists regarding identity of the tumour are very common. The gene expression activities yielded by molecular and genomic biology are more objective to classify diseases as the usual belief is that cell phenotypes have genotypic origins. Recent success in subclassification of neoplasms within a

disease group using gene expression profiles (Golub et al., 1999, Hedenfalk et al., 2001) provide support for such a belief.

The major roadblock is the small sample size issue inherent to microarray based classification effort. Contributing to this are the limited number of human tissues for study and the cost of such gene expression profiling projects. We want to identify classifiers which: (i) are flexible to execute complex classifications efficiently, (ii) automatically reduce the dimension to accommodate the small sample size problem and (iii) can identify the significant genes. The method based on RKHS developed here satisfies all of these criteria.

The use of RKHS theory helps us to change the dimension of the problem from p to n. In cases when we use the gene expression covariates, p, the number of covariates, is much greater than n, the number of samples. Hence RKHS method provides us with an automatic dimension reduction from pto n. Earlier papers proposed Bayesian probit model approaches with latent variables for modelling cancer tumours with more than two classes. But all these methods are much restricted compared to ours in the sense that they used simple linear model to model the latent variables. In contrast, our model does not impose any kind of structure on the latent variables. The relationship between the latent variables and the microarray covariates is denoted by an unknown function f. and we assume that the function belongs to an abstract function space, the RKHS. This is the only assumption we make. Then maximize the penalized log likelihood or the posterior to estimate the unknown function. This type of function estimation helps us to come up with a more flexible class of models. Although the use of RKHS helps us to bypass the problem of gene selection by already reducing the dimension of the model from p to n, in real life applications, an initial gene selection is always recommended. From Table 1, it is clearly seen that all the standard models and our two models gave extremely poor performance when all genes are used. Rather than doing a two step model fitting, which can induce possible bias in the classifier, i.e, an initial gene selection and then fitting the models using only the selected genes, in Section 6, we suggested an integrated Bayesian gene selection and model fitting approach with the help of indicator variables.

The multicategory SVM proposed by Lee et al. (2004) makes use of the RKHS theory and an extension of the hinge loss. Our BMSVM model is an extension of their approach in the Bayesian paradigm. Lee et al. (2004) treated the whole problem of multiclass classification from an optimization

standpoint, whereas our method treats the whole problem in a probabilistic framework. Unlike the frequentist approach, in our method, the kernel parameter θ is not fixed, and we put a prior on it. By putting the prior on the kernel parameter, we gain as we eventually use a mixture of kernels. We can also obtain the full posterior predictive probability distribution of $p(t_{new} = j)$, for $j = 1, \ldots, J$, i.e., the probability that a new tumour belongs to the *j*th type. The full posterior predictive probability distribution contains much more information than just a point estimate, and we can easily construct a confidence interval based on it.

The use of near-diffused proper priors helps us to make our method less sensitive to the choice of prior parameters. It also ensures that the posterior is proper so that we can use all standard MCMC techniques to generate samples from it. The procedure is definitely sensitive to the choice of ω as it controls the number of genes each time it is included in the model. In both the examples, we have kept $\omega = 0.01$ as suggested by Lee et al. (2004), and Sha et al. (2004). It means that only 10% of the genes are expected to be included in the model, which indirectly implies that we are inducing sparsity. As an alternative, we can also assign a hierarchical Beta(a, b) prior on ω .

Our BMLM and BMSVM have lower misclassification errors than the standard methods like neural network, classical support vector machine and random forest in modelling the glioma cancer. Our methods gave better results than all three standard ones. Both BMLM and BMSVM, when integrated with the Bayesian variable selection technique, give improved performance. Hence we recommend either of our models with multiple smoothing parameters augmented with the Bayesian variable selection technique.

Identifying a particular class or type of glioma cancer is very important for its diagnosis and treatment. Targeting specific therapies to pathogenetically distinct tumour types is important for cancer treatment because it maximizes efficacy and minimizes toxicity (Golub et al., 1999). Toxicity plays a major role as the target area of the treatment is the brain or central nervous system, and any kind of toxic effect of the drug or treatment may lead to long lasting potential hazard to the patient. Diagnostic pathology has traditionally relied on macro- and microscopic histology and tumour morphology as the basis for tumour classification. Of all cancers, the gliomas are the hardest to classify and current methods are often unable to do the correct classification. In this paper, our proposed models are able to identify accurately different types of gliomas simultaneously with gene selection. From Section 8, we see that the genes marked as active or important by our models carry some special biological significance as they can lead to some new lines of investigation for the biologists and geneticists.

In a broader context, the approach applied in this study can be used to identify genes that contribute to the major differences between any two groups of samples analysed. In the process of this, some less understood phenotypes might be identified. For example, we might find significant genes that distinguish cancers with high metastatic potential from cancers with little or no metastatic potential or genes that identify cancers that will be sensitive to specific therapies versus those that will be resistant and continue to grow unabated through therapy. Current histology-based classification and grading systems can do neither of these. Identification of such significant genes may not only provide markers for diagnosis and disease management but may also provide novel potential targets for drug development. A method that could identify the strong features of cancer, both genotypically and phenotypically, would provide an ideal route to the heart of the problem and we will use our method for these future studies.

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Paper received March 2007; revised December 2007.