gamselBayes: Bayesian generalized additive model selection including a fast variational option

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1 Introduction

The R package gamselBayes implements algorithms for Bayesian generalized additive model selection developed by and described in He & Wand (2022). The underlying model is similar to, and based on, that developed by Chouldechova & Hastie (2015), and implemented in the R package gamsel (Chouldechova & Hastie, 2018).

The default fitting and inference algorithm in gamselBayes is a Markov chain Monte Carlo scheme. For problems of moderate size, the run time with this default approach is likely to be acceptable. However, it is well-known that Markov chain Monte Carlo approaches do not scale well to very large problems. With such circumstances in mind, the gamselBayes package also also provides the option of using a faster mean field variational Bayes algorithm for generalized additive model selection.

In Sections 2 -7 of this vignette we work through some illustrative examples involving simulated and actual data. Section 8 describes limitations of the gamselBayes and contains some trouble-shooting advice.

2 Illustrations with Simulated Data

We start with two data sets simulated from Gaussian response additive models. Such examples have the advantage of explaining the use of gamaelBayes in simple terms, and also allowing comparison with true functions from which the data are simulated.

2.1 All Candidate Predictors Continuous

The following code generates data corresponding to the additive model

$$y_i \stackrel{\text{ind}}{\sim} N\left(\sum_{j=1}^6 f_j(x_{ji}), \sigma^2\right), \quad 1 \le i \le n,$$
 (1

with n = 500 and $\sigma = 0.15$:

```
> sigmaTrue <- 0.15

> fiTrue <- function(x) return(0.96*x)

> f2True <- function(x) return(0.5*(pnorm(6*x - 3) + 1))

> f3True <- function(x) return(0.5*(sin(3*pi*x) + 1))

> f4True <- function(x) return(0.5*(0.04*cosh(x^3 - 9*x^2 + 4) + 1))

> set.seed(1); n <- 500

> xl <- runif(n); x2 <- runif(n); x3 <- runif(n)

> x4 <- runif(n); x5 <- runif(n); x6 <- runif(n)

> y <- rnorm(n,f1True(x1) + f2True(x2) + f3True(x3) + f4True(x4),sigmaTrue)

> Kgeneral <- data.frame(x1,x2,x3,x4,x5,x6)
```

The data for all 6 candidate predictors x_1, \ldots, x_6 are stored in the data frame Xgeneral. The response data are stored in the vector y.

These data are simulated from f_i functions such that:

 f_1 is linear: f_2 , f_3 , and f_4 are non-linear: f_5 and f_6 are zero.

2.1.1 Analysis Using the Default Method: Markov Chain Monte Carlo

The main function in the gamselBayes package is gamselBayes(). The default gamselBayes() fit object, labeled fit1, is obtained via:

> library(gamselBayes) : fit1 <- gamselBayes(v = v.Xgeneral = Xgeneral)

A quick look at the estimated effect types is achieved via:

> effectTynesVector(fit1)

To obtain a tabulated version of the same information, issue:

```
> effectTypes(fit1)
```

```
Predictor selected as having a linear effect:
```

Predictors selected as having non-linear effects:

x2 x3 x4

This output shows that, for this example, gamselBayes() correctly estimates the effect of x_1 to be linear, the effects of x_2 , x_3 and x_4 to be non-linear and the effects of x_5 and x_6 to be zero.

A Bayesian inferential summary of the linear effect coefficient is achieved using:

> summarv(fit1)

```
posterior mean 95% credible interval
x1 0.97625 0.9268 1.0263
```

and is in keeping with the true value of the coefficient, 0.96, that generated the data.

Visualisation of the non-linear estimated effects is obtained using:

```
> plot(fit1)
```

and is shown in Figure 1. Issuing code such as the following can be used to compare the estimates shown in Figure 1 with the true functions:

```
> par(mfrow=c(2,2)) ; xg <- seq(0,1,length = 1001)
> f2g <- f2True(xg) ; f3g <- f3True(xg) ; f4g <- f4True(xg)
```

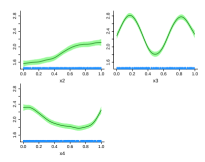


Figure 1: The plots produced by the command plct(fit1) for the first gamselBayes() fit, stored as fit1, to the data simulated according to (1). The curves are estimates of f2, f3 and f4. The convention of plct() for gamselBayes() is to display relevant slices of the additive model fit, for which each of the other functions are evaluated at the median value of their predictor. The shaded regions correspond to pointwise 95% credible intervals. The rugs at the base of each plot show values of each predictor.

Whenever the default Bayesian inference method is used then it is important to keep in mind that the results depend on behaviour of the Markov chain Monte Carlo samples, also known as chains. The function checkChains() facilitates a cursory graphical check of the particular chains and the command:

> checkChains(fit1)

leads to the graphic shown in Figure 2. This graphic shows that key chains for each predictor are well-behaved and that the resultant Bayesian inference is quite trustworthy.

Details on the graphics produced by the checkChains() function are provided by the command:

> help(checkChains)

2.1.2 Analysis Using a Faster Alternative Method: Mean Field Variational Bayes

The default method for Bayesian inference is Markov chain Monte Carlo. For large sample sizes and dimensions, this approach can be slow to compute. To help mitigate this problem gamselBayes() offers and alternative faster approach based on mean field variational Bayesian approximate inference. The relevant argument specification is method = "MFVB" and is illustrated for this section's example via:

The estimated effect types are obtained using:

predictor	trace	lag 1	acf
x1	Many disease and the con-		
x2	And the state of	- Page 1	Herm
х3	and her profiled godin.		
x4	and all property of the second		

Figure 2: The graphic produced by the command check(Chains(fit1) for the gamselBayes() fit object fit1. For the predictor, x₁, selected as having a linear effect the chain is the coefficient of this predictor. For the predictors, x₂, x₃, x₄, selected as having non-linear effects the chains corresponds to vertical slices of the non-linear fits at the median of each predictor. The second column is a time series (trace) plot of the chain. The third column is a scatterplot of the chain values against their previous (lag 1) values. The fourth column is the sample autocorrelation function (acf.) based on the R function acf().

```
Predictor selected as having a linear effect:

x1

Predictors selected as having non-linear effects:
```

This output shows that, for this example, gamselBayes() with method = "MFVB" correctly estimates the effect of x_1 to be linear, the effects of x_2 , x_3 and x_4 to be non-linear and the effects of x_5 and x_6 to be zero.

A Bayesian inferential summary of the linear effect coefficient is achieved using:

```
posterior mean 95% credible interval
```

Visualisation of the non-linear estimated effects is obtained using:

```
> plot(fit2)
```

> summarv(fit2)

and is shown in Figure 3.

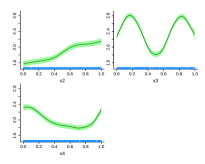


Figure 3: The plots produced from the command plot(fit2) for the gamselBayes() fit with method = "MFVB", stored as fit2, to the data simulated according to (1). The curves are vertically centred estimates of f₂, f₃ and f₄. The shaded regions correspond to pointwise 95% credible intervals. The rugs at the base of each plot show values of each predictor.

2.2 Candidate Predictors Continuous Both Continuous and Binary

We now consider a simulated data example where not all of the predictors are continuous.

Three binary predictors, x₇, x₉ and x₉, are added so that the model becomes:

$$y_i \stackrel{\text{ind.}}{\sim} N\left(\sum_{i=1}^{9} f_j(x_{ji}), \sigma^2\right), \quad 1 \le i \le n,$$
 (2)

Data from model (2) are generated according to:

```
> fTTrue <- function(x) return(-0.78*x)
> f8True <- function(x) return(0.53*x)
> set.seed(1); n <- 500
> x7 <- rbinom(n,1,0.5); x8 <- rbinom(n,1,0.5); x9 <- rbinom(n,1,0.5)
> y <- rnorm(n,f1True(x1) + f2True(x2) + f3True(x3) + f4True(x4)
+ f7True(x7) + f8True(x8), sigmaTrue)
> Xgeneral <- data.frame(x1,x2,x3,x4,x5,x6)
> Xlinear <- data.frame(x7,x8,x9)
```

where the data frame Xlinear corresponds to a design matrix that contains predictors that can only have a zero or linear effect.

The functions $f_1, ..., f_6$ are the same as in the previous example and the code for f1True(), ..., f4True()

still applies. The functions f_7 and f_8 are linear and non-zero. The function f_9 is zero. The call to gamselBayes() should now use the Xlinear argument as follows:

> fit3 <- gamselBayes(y = y,Xlinear = Xlinear,Xgeneral = Xgeneral)

The estimated effect types are tabulated using:

> effectTypes(fit3)

```
Predictors selected as having linear effects:
```

Predictors selected as having non-linear effects:

v2 v3 v/

A Bayesian inferential summary of the linear effect coefficients is achieved using:

> summary(fit3)

interval	credible	95%	posterior mean	
1.02400	0.92812		0.97539	x1
-0.73743	-0.79282		-0.76608	x7
0.57137	0.51799		0.54394	x8

These are in keeping with the true value of the coefficients, $0.96,\,-0.78,\,0.53$ that generated the data.

The command:

> checkChains(fit3)

leads to the graphic shown in Figure 4. The Figure 4 graphic shows good behaviour of the

predictor	trace	lag 1	acf
x1	andersethingsoff,		
x2		A Company	
х3	And produced in plant		
x4	There exists to the		
x7	Construction of the Constr		
x8	atogodoron destrott Hepritoron deprio		

Figure 4: The graphic produced by the command checkChains(fit3) for the gamselBayes() fit object fit3. The Figure 1 caption provides full details on checkChain() graphics.

chains.

3 Analysis of California Schools Data

The R package Ecdat (Crossaint, 2020) contains a data frame named Caschool. The data frame contains data on 14 variables for 420 school districts in the state of California, U.S.A. The following code leads in the data and produces descriptions of each of the variables:

```
> library(Ecdat) : data(Caschool) : help(Caschool)
```

Two outcome variables are

mathser average mathematics score for the district.

readscr average reading score for the district.

We will use the first of these as a response variable. Ten of the remaining 11 variables in Caschool are continuous. These are:

enritot total enrolment

teachers number of teachers.

calword percentage of students qualifying for the CalWORKSs welfare programme.

mealoct percentage of students qualifying for reduced-price lunches.

computer number of computers.

compstu number of computers per student.

expostu expenditure per student

str student teacher ratio.

avginc average income of the district.

elact percentage of English learners.

The other variable, Caschool["grspan"] is categorical and codes whether the school has grades spanning from kindergarten to year 6 or from kindergarten to year 8. The following binary variable is obtained for use in gasselBayes():

```
> upToYear8 <- as.numeric(Caschool[, "grspan"] == "KK-08")
```

3.1 Analysis for Raw Data

We are ready to set up the response vector y and the Xlinear and Xgeneral predictor variable data frames, with the following commands:

Note that Xgeneral contains each of the available continuous predictors except for Caschool["teachers"] and Caschool["computer"], since they are simple functions of other predictors. The following default call to gamsolBayes() fit command:

```
> fit4 <- gamselBayes(y = y,Xlinear = Xlinear,Xgeneral = Xgeneral)
```

leads to the fit object being stored in fit4. Then:

> effectTynes(fit4)

leade to

```
Predictors selected as having linear effects:
```

```
avginc elect mealect
```

We see that, out of the 9 candidate predictors, 3 are chosen and each of them are found to have linear effects. The selected model is a linear combination of mealpct, elpct and avginc. The command

> summarv(fit4)

leads to the following inferential summary of the coefficients:

	posterior mean	95% credible	interval
avginc	0.70911	0.50118	0.903100
elpct	-0.13935	-0.22267	-0.045745
meal nct.	-0.34517	-0.43269	-0.255400

It shows, for example, a significant elevation in mean mathematics scores for districts with higher average incomes.

Since these results are dependent on Markov chain Monte Carlo sampling, it is prudent to perform a check of the chains using:

> checkChains(fit4)

This leads to the chain visual summary shown in Figure 5. The chains are reasonably well-

predictor	trace	lag 1	acf
avginc	naliteterini		
elpct			Hananan
mealpct	Activities to July		Шин

Figure 5: The graphic produced by the command checkChains(fit4) for the gamselBayes() fit object fit4. The Figure 1 caption provides full details on checkChain() graphics.

behaved, but it may be worth considering larger warm-up and kept chain sizes. The following command leads to a new fit with a two-fold increase in the chain sizes compared to their default values:

```
> fit5 <- gamselBayes(y = y,Xlinear = Xlinear,Xgeneral = Xgeneral,
+ control = gamselBayes.control(nWarm = 2000,nKept = 2000))</pre>
```

The longer chains are visualised using:

> checkChains(fit5)

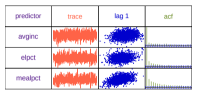


Figure 6: The graphic produced by the command checkChains(fit5) for the gamselBayes() fit object fit5. The Figure 1 caption provides full details on checkChain() graphics.

and the result is shown in Figure 6. The coefficient summary for this longer chain fit is obtained using:

> summary(fit.5)

	posterior mean	95%	credible	interval
avginc	0.69273		0.48851	0.886320
elpct	-0.14105		-0.22356	-0.045906
mealpct	-0.34925		-0.43759	-0.249040

The coefficient summary shows that the variables percentage of students qualifying for reducedprice lunches and percentage of English learners have statistically significant negative effect on mean mathematics score. The average income of the district has a significant positive effect.

3.2 Analysis with Some Logarithmically Transformed Predictors

Three of the continuous predictors used in the previous analysis are quite skewed. It may be worthwhile to see what happens if skewness-reducing logarithmic transformations are used. The following code achieves this:

log.avginc

The main change is the presence of a non-linear effect for the logarithm of the district average income. This effect can be viewed using:

> plot(fit6,xlab = "log(average income of school district)")

and is shown in Figure 7. The linear effects summary is:

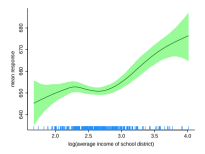


Figure 7: The plots produced from the command plot(fit6xlsb = "log(average income of school district)") for the first gamselBayes() fit, stored as fit6. The shaded regions correspond to pointwise 98% credible intervals. The rugs at the base of each plot show values of each predictor.

> summary(fit.6)

	posterior mean	95% credible	interval
log.elpct	-1.85110	-3.11310	0.00000
mealpct	-0.38873	-0.50444	-0.28386
upToYear8	-4.38100	-7.46970	0.00000

The gamselBayes chain check for fit6 involves:

> checkChains(fit6)

and leads to the chain visual summary shown in Figure 8.

This final gamselBayes() fit to the California schools data has the following aspects:

- 4 of the 9 candidate predictors are selected.
- . 3 of them, log(elpct+1), mealpct and upToYear8 have linear effects on the mean response
- 1 of them, log(avginc) impacts the response in a non-linear fashion, with two ramps and a plateau, as shown in Figure 7.

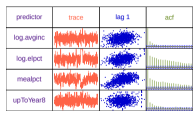


Figure 8: The graphic produced by the command checkChains(fit6) for the gamselBayes() fit object fit6. The Figure 1 caption provides full details on checkChain() graphics.

4 Analysis of Boston Mortgage Applications Data

The data frame BostonMortgages within the HRW package (Harezlak, Ruppert & Wand, 2021) contains data on 13 variables for 2,380 mortgage applications in Boston, U.S.A., during the years 1989–1999. The data are loaded usine:

```
> library(HRW) : data(BostonMortgages)
```

Descriptions of the variables are provided by:

> help(BostonMortgages)

A key variable is BostonMortgages["deny"], and codes whether or not the mortgage application was denied. A numerical version of this variable, for which I codes the application being denied and 0 codes the application being approved, is used as a response variable in this illustration. The code for setting up the response data vector is:

```
> v <- as.numeric(BostonMortgages[."denv"] == "ves")
```

Next we set up the data frame of vectors which enter the model linearly. The first phase involves conversion of five factor variables with yes/no coding to binary forms:

```
> Xlinear <- BostonMortgages[,c("pbcr","dmi","self","single","black")]
> for (i in 1:ncol(Xlinear))
```

```
+ Xlinear[,j] <- as.numeric(Xlinear[,j] == "yes")</p>
```

The indicator of whether or not the mortgage application pertains to a condominium is added on using:

> Xlinear\$condominium <- BostonMortgages[,"condominium"]

The BostonNortgages data frame includes a 6-level ordinal variable that represents credit risk. The following code creates and adds to Xlinear five indicator variables for each of the credit scores that differ from 1:

```
> Xlinear$CCSeq1 <- as.numeric(BostonMortgages$ccs==1)
> Xlinear$CCSeq2 <- as.numeric(BostonMortgages$ccs==3)
> Xlinear$CCSeq3 <- as.numeric(BostonMortgages$ccs==3)
> Xlinear$CCSeq4 <- as.numeric(BostonMortgages$ccs==5)
- Xlinear$CCSeq5 <- as.numeric(BostonMortgages$ccs==5)
```

The next set of commands a similar to the credit score processing, but are for a 4-level ordinal mortgage credit score variable:

```
> Xlinear$MCSeq1 <- as.numeric(BostonMortgages$mcs==1)
> Xlinear$MCSeq2 <- as.numeric(BostonMortgages$mcs==2)
> Xlinear$MCSeq3 <- as.numeric(BostonMortgages$mcs==3)</pre>
```

The variable BostonMortgages[,"uria"] is quantitative and represents the unemployment rate in the applicant's industry. However, it only has 10 unique values and is not conducive to spline-based estimation of non-linear effects. Therefore, it is included in the linear effects-only data frame using:

> Xlinear\$uria <- BostonMortgages[."uria"]

The three remaining predictors are quantitive ratio variables with hundreds of unique values. These are definitely worth considering for possible non-linear effects and make up the Xgeneral data frame:

> Xgeneral <- BostonMortgages[.c("dir", "hir", "lyr")]

In the call to ${\tt gamselBayes}$ () it is important to specify ${\tt family} = {\tt "binomial"}$ due to the binary nature of y for this example:

```
> fit7 <- gamselBayes(y = y,Xlinear = Xlinear,Xgeneral = Xgeneral,
+ family = "binomial")</pre>
```

The last command leads to the fit object being stored in fit7. Then:

```
> effectTypes(fit7)
```

leads to

```
Predictors selected as having linear effects:

CCSeq1 CCSeq2 black dmi pbcr

Predictors selected as having non-linear effects:
```

dir lvr

This output shows that

- · 5 of the candidate predictors are selected as having linear effects,
- · 2 of the candidate predictors are selected as having non-linear effects.

To assess the effects of the variables chosen as having linear effects, we issue:

> summarv(fit7)

which leads to the following inferential summary of the coefficients:

	posterior mean	95%	credible	interval
CCSeq1	-0.68844		-0.92380	-0.46025
CCSeq2	-0.32369		-0.60340	0.00000
black	0.35240		0.15877	0.53789
dmi	2.64750		2.09810	3.35000
pbcr	0.72527		0.49313	0.97320

It shows, for example, that 3 of the selected linear effect predictors have a positive impact on the probability of mortgage denial: indicators of the applicant being self-employed, single or African-American.

The two predictors having non-linear effects in fit7 are ratio of the debt payments to the total income and ratio of the loan size to the assessed value of the property. Figure 9 shows these effects.

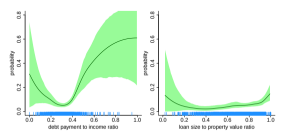


Figure 9: The plots produced from the command plot(fit7.xiim = rbind(c(0.1),c(0.1)),c(0.1)),xim = rbind(c(0.0.8),c(0.0.8)), xiab = c("debt payment to income ratio", "loan size to property value ratio") for the first genselbayes () fit, stored as fit7. The curves are vertically centred estimates of the effects of ratio of the debt payments to the total income (left panel) and ratio of the loan size to the assessed value of the property (right panel) on the probability of mortgage application denial. The shaded regions correspond to pointwise 95% credible intervals. The rugs at the base of each ylot show values of each prodictor.

Lastly, we carry out a check of the chains on which the fitting and inference of fit7 is based. Since there are 11 selected predictors in fit7, the checkChains() graphics are divided into two graphics with a maximum of 6 predictors per graphic.

> checkChains(fit7)

The result is shown in Figure 10. All chains are seen to be reasonably well-behaved and the approximate Bayesian inference is sound.

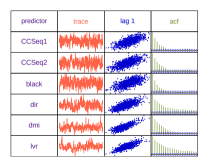
5 Analysis of Data on House Sales in Sydney, Australia

This example depends upon the R package HRW, which accompanies the Harezlak, Ruppert & Wand (2018) book. After making sure that HRW is installed, load the data using:

```
> library(HRW) ; data(SydneyRealEstate)
```

The command:

> help(SydneyRealEstate)



predictor	trace	lag 1	acf
pbcr			<u> </u>

Figure 10: The graphic produced by the command checkChains(fit7) for the gamselBayes() fit object fit7. The Figure 1 caption provides full details on checkChain() graphics.

leads to a detailed description of the SydneyRealEstate data frame. It has 39 variables corresponding to 37,676 house sales in Sydney, Australia, during the year 2001.

The following code determines the indices of important parts of the SydneyRealEstate data frame:

```
> indResponse <- 1; indsContinEarly <- c(2:4,7)
> indSaleDate <- 5; indSaleQtr <- 6
> indPostCode <- 8; indCrimeRate <- 10
> indAirNoise <- 24: indsContLater <- c(11:23.25:29.33:39)
```

Next, we obtain indicator variables for the sale quarter variable and presence of aircraft noise:

```
> saleQtrEq2 <- as.numeric(SydneyRealEstate[,indSaleQtr] == 2)
> saleQtrEq3 <- as.numeric(SydneyRealEstate[,indSaleQtr] == 3)
> saleQtrEq4 <- as.numeric(SydneyRealEstate[,indSaleQtr] == 4)
> aircraftNoise <- as.numeric(SydneyRealEstate[,indAirNoise] > 0)
```

Most of the other potential predictor variables are well-behaved. However, running the commands:

```
> par(mfrow = c(1,1))
> hist(SydneyRealEstate[,"crimeRate"],breaks = 100,col = "plum")
```

shows that SydneyRealEstate[,"crimeRate"], which is a measure of the crime rate of each house's suburb, is highly skewed and with a small fraction of the houses having a large outlying value (the histogram is not shown here). Application of the the $\log(x+1)$ transformation makes the data more amenable to the $\operatorname{\mathsf{gamsel}}()$ methodology. From now on we work with this transformation of $\operatorname{\mathsf{Sydepsel}}(\operatorname{\mathsf{state}})$ regimeRate¹]

Three other potential predictors, SydneyRealEstate[.c("neph","PM10","SO2")], are quantitative but have low numbers of unique values. To avoid problems with spline fitting, these predictors are considered as having only a linear effect, and not permitted to have possible non-linear effects.

We are now ready to set up the the response vector and predictor data frames:

Given the high sample size of 37,676 and the fact that the numbers of potential predictors in Xlinear and Xgeneral are, respectively, 7 and 30 we use the faster mean field variational Baves approach.

```
> fit8 <- gamselBayes(y = y,Xlinear = Xlinear,Xgeneral = Xgeneral,
+ method = "MFVB")</pre>
```

```
The effect types of the selected predictors are tabulated using:

> effectTypes(fit8)

are as follows:

Predictors selected as having linear effects:

distToFreeway distToMedical distToTunnel
neph saleQtrEq2 saleQtrEq3 saleQtrEq4

Predictors selected as having non-linear effects:

NO2 distToAmbulance distToBusStop
distToCoastline distToFactory distToFerry
distToGPU distToHighway distToHospital
distToMainRoad distToUniversity
distToMaselaeRoad discreainerRatio income
```

latitude log(crimeRate + 1) longitude lotSize

ozone

For the 7 selected linear effects the command:

```
> summary(fit8)
```

leads to

```
posterior mean 95% credible interval
distToFreeway
                                0.0085231 0.0110140
                 0.0097661
dietToModical
                -0 0046228
                               -0.0071508 -0.0020682
distToTunnel
                 0.0067200
                                0.0056759 0.0077919
nenh
                -0 2794900
                               -0.3610100 -0.1986600
saleOtrFo2
                 0.0203810
                                0.0105800 0.0300970
saleOtrFd3
                 0.0655930
                                0.0565690 0.0744180
                 0 1323800
saleOtrEq4
                                0.1233400 0.1417800
```

This output shows, for example, that houses sold later in the calendar year, such as the 4th quarter period of October–December, tend to have higher sales prices.

The commands:

lead to the plot of the estimated non-linear effects, shown in Figure 11. In all, there are 22 estimated non-linear effects shown in Figure 11. The first panel shows that the effect of lot size is mainly monotonically increasing, but with some ramps and a plateau. For distance to coastline there is a sharp rise for houses that are within 1 kilometre of the coast.

6 Analysis of Car Auction Data

This example also depends upon the ${\sf R}$ package ${\sf HRW}.$ Assuming that ${\sf HRW}$ is installed, load the data via:

```
> library(HRW) ; data(carAuction)
```

Then issue:

```
> help(carAuction)
```

to obtain a detailed description of the carAuction data frame. This data frame has 51 variables corresponding to 72, 983 cars purchased at auctions by automobile dealerships in the United States of America and originates from the competition titled "Don't Get Kicked!" that ran on the kaggle platform (www.kaggle.com) during 2011-2012.

In this illustration we hold back data on 2,983 of the cars for testing. The remaining 70,000 will be used as a training set for fitting a probit additive model via gamselBayes():

```
> namesNonPred <- c("Refid", "IsBadBuy")
> namesGeneral <- c("dodmRead", "price", "costAtPurch", "warrantyCost")
> namesLinear <- setdiff(names(carAuction), union(namesNonPred, namesGeneral))
```

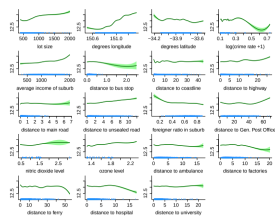


Figure 11: The graphic produced by the command plot(fit8,xlab = xlabDetailed) for the gamsolBayes() fit object fit8. The curves are vertically centred estimates of the effects of each predictor selected as having a non-linear effect. The shaded regions correspond to pointwise 95% credible intervals. The rugs at the base of each plot show values of each predictor. Due to the very larve sample size, random subsets of size 1,000 are used in the rugs.

- > nTotal <- nrow(carAuction)
- > nTest <- 2983 ; nTrain <- nTotal nTest
- > testInds <- sample(1:nTotal,nTest,replace = FALSE)
- > trainInds <- setdiff(1:nTotal.testInds)

Now set up the response data vector and predictor data frames:

- > yTrain <- carAuction\$IsBadBuy[trainInds]
- > XlinearTrain <- carAuction[trainInds,namesLinear]
- > XgeneralTrain <- carAuction[trainInds,namesGeneral]

The number of candidate predictors in XinearTrain is 45, whilst the number of candidate predictors in XgeneralTrain is 4. In the call to gamselBayes() it is important to specify family = "binomial" due to yTrain being binary:

```
> fit9 <- gamselBayes(y = yTrain,Xlinear = XlinearTrain,
```

+ Xgeneral = XgeneralTrain,family = "binomial")

The last command leads to the fit object being stored in fit9. Then:

> effectTypes(fit9)

leads to the following estimated effects types:

```
Predictors selected as having linear effects:

ageAtSale makeEqChevrolet makeEqChrysler
odomRead purchIn2010 purchInFlorida
purchInNorthCarolina purchInTexas
sizeEqMedium sizeEqSUV transEqManual
trimEqBas wheelEqAlloy wheelEqCovers

Predictors selected as having non-linear effects:
```

costAtPurch warrantyCost

We see from this output that, out of the 49 candidate parameters, 11 are selected as having a linear effect, and 2 are selected as having a non-linear effect.

A Bayesian inferential summary of the linear effect coefficients is achieved using:

```
> summary(fit9)
```

and leads to

```
posterior mean 95% credible interval
ageAtSale
                    9.5264e-02 8.4529e-02 1.0658e-01
                   -1.1605e-01 -1.6643e-01 -5.3060e-02
makeEqChevrolet
makeEqChrvsler
                    9.1270e-02 0.0000e+00 1.5121e-01
                    3.4235e-06 2.2505e-06 4.4529e-06
odomBoad
purchIn2010
                    9.7396e-02 6.7546e-02 1.2879e-01
purchInFlorida
                   -1.1759e-01 -1.6779e-01 -7.2159e-02
purchInNorthCarolina -1.0757e-01 -1.5943e-01 -6.1273e-02
purchInTexas
                    8.9598e-02 5.0214e-02 1.2514e-01
                   -6.6150e-02 -9.7408e-02 -3.2660e-02
sizeEaMedium
                    1.8834e-01 1.4006e-01 2.3328e-01
sizeEqSUV
transEqManual
                   -1.6450e-01 -2.3466e-01 -9.5746e-02
                    7.0185e-02 2.7402e-02 1.1022e-01
trimEqBas
wheelEqAlloy
                   -1.5214e+00 -1.5665e+00 -1.4754e+00
wheelEqCovers
                    -1.5905e+00 -1.6388e+00 -1.5410e+00
```

Some of the effects match intuition, such as older cars having a higher probability of being a bad buy at auction. Less intuitive is the fact that the same is true for cars that were purchased in North Carolina.

The command:

leads to the figure shown in Figure 12, which shows the effects of

the acquisition cost paid for the car at the time of purchase in U.S. dollars

and

the warranty cost in U.S. dollars

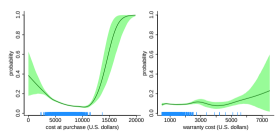


Figure 12: The plots produced from the command plot(fit9.xim = rbind(c(0.2000),rep(NA.2)), xlab = c("cost at purchase (U.S. dollars)", "warranty cost (U.S. dollars)")) for the gammelBayes() fit, stored as fit9. The shaded regions correspond to pointwise 95% credible intervals. The rugs at the base of each plot show values of each predictor. Due to the very large sample size, random subsets of size 1.000 are used in the rugs.

The former has an interesting U-shaped effect.

Next we assess how well cars in the test data set are classified according to the fit in fit9, starting with setting up test versions of the v. Xlinear and Xgeneral inputs:

```
> yTest <- carAuction$IsBadBuy[testInds]
> XlinearTest <- carAuction[testInds,namesLinear]
> XgeneralTest <- carAuction[testInds.namesGeneral]</pre>
```

The vector of predictions on the probit (Standard Normal quantile function) scale is obtained using:

> probVecTest <- predict(fit9.newdata = list(XlinearTest.XgeneralTest))

The following code then obtains and prints the confusion matrix:

Lastly, we obtain an estimate of the misclassification rate via:

classified bad buy

57

76

```
> estMisClassRate <- 100*sum(yTestHat != yTest)/nTest
> cat("The estimated misclassification rate is ",
+ signif(estMisClassRate,4), "%.\n",sep = "")
```

The estimated misclassification rate is 11 3%

The classifier based on fit9 classifies about 88.5% of cars correctly in terms of their bad buy versus good buy status.

7 Fitting and Plotting Options

Whenever gamselBayes() is applied to a particular data set there are numerous options concerning, for example, the size of the spline basis and hyperparameter choices. In this section we illustrate the tweaking of such values. Similar comments apply to plotting a gamselBayes() object

Throughout this section, we work with the California schools data set that was described and analysed in Section 3. The following commands from that section load the required packages and set up the input data objects:

7.1 Sparsity Imposition Options

Let β be a generic regression coefficient attached to one of the linear predictors. In models used by gamselBayes(), $\beta = \gamma_{\beta} \tilde{\beta}$ where γ_{β} is binary and $\tilde{\beta}$ is continuous. Therefore

$$P(\beta = 0|y) = P(\gamma_{\beta} = 0|y),$$

and the posterior distribution of γ_{β} can be used to decide between hypotheses $H_0: \beta=0$ and $H_1: \beta\neq0$. A natural rule is to accept H_0 if and only if $P(\gamma_{\beta}=0|\mathbf{y})>\frac{1}{2}$. However, with parsimony in mind, $\mathbf{gansolBayes}()$ also supports less stringent rules. Rather than thresholding $P(\gamma_{\beta}=0|\mathbf{y})$ at $\frac{1}{2}$, consider a family of rules indexed by a threshold parameter $\tau\in[0,1)$. For the case of τ being zero or very small it is useful in practice to have an "effective zero" parameter, $\varepsilon_{off,soro}$, which is set to a small positive number such as 0.00001. Note that $\tau=0$ corresponds to maximum a posteriori estimation of β . After fixing τ and $\varepsilon_{off,soro}$, our strategy for deciding between an effect being zero or linear is

the effect is zero if $P(\gamma_{\beta} = 0|y) > \max(\tau, \varepsilon_{\text{eff,zero}})$, otherwise the effect is linear.

Analogous rules are used for deciding between an effect being linear or non-linear.

Lower values of τ lead to sparser fits, so in the gamselBayes() function we use the identifier lowerMakesSparser for the τ parameter. The default value of lowerMakesSparser is 0.1 when method="MCMC" and 0 when method="MFVB". The following command stipulates that lowerMakesSparser be set to the higher value of 0.5:

```
> fit10 <- gamselBayes(y = y,Xlinear = Xlinear,Xgeneral = Xgeneral,
+ lowerMakesSparser = 0.5)</pre>
```

For this choice of thresholding, the command: > effectTynes(fit10) shows that the estimated effect types become: Predictors selected as having linear effects: calwort log.elpct mealpct upToYear8 Predictor selected as having a non-linear effect. log.avginc which is less sparse than the default fit. The command-> summarv(fit10) then leads to the following inferential summary for the linear coefficients: posterior mean 95% credible interval -0.12442 -0.26213 0.00000 -2.02740 -3.20800 -0.96832 -0.37274 -0.46437 -0.28234 calunct log.elpct mealnct upToYear8 -4 51630 -7 33090 0 00000 The function gamselBayesUpdate() allows one to change the value of lowerMakesSparser without having to re-do the fitting phase. If, for example, a change of from τ from 0.5 to 0.2 is of interest then the command: > fit11 <- gamselBayesUpdate(fit10,lowerMakesSparser = 0.2) quickly provides the new fit object with the newer thresholding. Maximum a posteriori model section corresponds to $\tau = 0$, as used by the following fit: > fit12 <- gamselBayesUpdate(fit10.lowerMakesSparser = 0) We then have: > effectTypes(fit12) Predictor selected as having a linear effect: mealpct Predictor selected as having a non-linear effect: log.avginc and

posterior mean 95% credible interval

-0.38203 -0.47367 -0.28358

> summary(fit12)

mealpct

7.2 Control Options

The control argument of the gamselBayes() function can be used to specify various aspects of the Bayesian model and the strategy for achieving approximate inference. Suppose, for example, that the following departures from the default settines are desired:

- the number of interior knots in the spline basis functions is 15.
- the spline basis includes a linear function.
- the prior Normal distribution standard deviation of the linear coefficients parameters is σ₂ = 150.
- the prior Normal distribution standard deviation of the spline basis coefficients parameters is σ_v = 2,000.
- the prior Beta distribution shape parameters of the of the spline basis coefficients spikeand-slab auxiliary indicator variables probability parameter are A_n = 9 and B_n = 2.
- the number of warm-up Markov chain Monte Carlo samples is 3,000.
- the number of kept Markov chain Monte Carlo samples is 10,000.
- the thinning factor of the kept Markov chain Monte Carlo samples is 2.

```
Then the following call to gamselBayes() achieves this:
```

For fit13, the estimated effect types are found by issuing

```
> effectTypes(fit13)
```

are:

```
Predictors selected as having linear effects:
```

log.elpct mealpct upToYear8

```
Predictor selected as having a non-linear effect:
```

```
log.avginc
```

and the linear coefficients summary from

```
> summary(fit13)
```

is

lo me

	posterior mean	95% credible	interval
g.elpct	-2.20060	-3.51570	-0.76229
alpct	-0.34451	-0.46268	-0.23367
ToYear8	-4.66180	-7.61850	0.00000

Access to the full set of control options is provided by:

```
> help(gamselBayes.control)
```

7.3 Plotting Options

Lastly, we demonstrate some of the options available for display of non-linear effects via the plot() function for gamselBayes() fit objects. These illustrations are for the fit object fit13 from the precious subsections.

The command:

```
> plot(fit13,estCol = "darkmagenta",varBandCol = "gold",rugCol = "seagreen",
+ xlab = "log(average income of school district)".cex.lab = 1.75)
```

modifies the colours of the function estimates, variability bands and predictor data rug-plot displays and leads to the result shown in Figure 13. Our plotting options are use of dashed

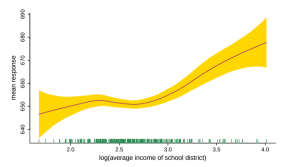


Figure 13: The plots produced from the command plot(fit13,estCol = "darkmagenta",varBandCol = "gold",rugCol = "seagreen", xlab = "log(average income of school district)",cex.lab = 1.75) for the gamselBayes() fit, stored as fit13. The shaded regions correspond to pointwise 95% credible intervals. The rue at the base of the plot show values of each predictor.

curves, rather than shaded polygons, to display variability bands and specification of the vertical frame limits using the vim argument. These are illustrated by the command:

```
> plot(fit13, shade = FALSE, estCol = "steelblue", varBandCol = "darkred",
+ rugCol = "limegreen", xlab = "log(average income of school district)",
+ cex.lab = 1.75)
```

and leads to Figure 14. Further details on plotting options can be obtained by issuing:

> help(plot.gamselBayes)

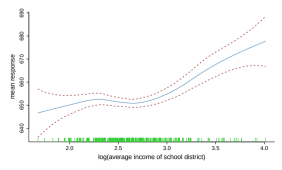


Figure 14: The plots produced from the command plot(fit13,shade = FALSE,estCol = "steel-blue",arBandCol = "darkred", rugCol = "limegreen",xlab = "log(average income of school district)", cc..lab = 1.75) for the gamselBayes() fit, stored as fit13. The shaded regions correspond to pointwise 95% credible intervals. The rug at the base of the vlot show values of each predictor.

8 Limitations and Trouble-Shooting

The gamselBayes package was written and is maintained, solely, by the authors of this vignette.

A reasonable amount of effort has been made to safeguard against breakdown for arbitrary
inputs. However, since gamselBayes is a two-person and non-industrial software package it
has limitations in this regard. In this final section we list some trouble-shooting tips that may
aid successful use of gamselBayes:

- Ensure that the input data is free of missing values and non-numerical objects. In other words, the input data in y, Xlinear and Xgeneral should contain only numbers.
- Predictors containing gross outliers can lead to problems with penalized spline fitting, so checks and pre-processing to avoid such an issue may be worthwhile.
- 3. If a predictor is strongly skewed then problems with penalized spline fitting are also more likely. Hence, depending on the strength of the skewness, pre-transformation of such predictors may be beneficial.
- 4. Each of the predictors in the Xgeneral input are treated as continuous variables. In theory, such variables have a large number of unique values. However, if its measurements have been discretized to the point that the number of unique value is smaller than around 15 25 then this can lead to problems with penalized spline fitting. One simple remedy is to move the predictor to the Xlinear input and live with its effect being restricted to zero or linear, but not non-linear.
- 5. In a similar vein to the last point, if the sample size is smaller than around 15 25 then all candidate predictors have the limitation described there and it may not be feasible to include them in the Xgeneral input.

Feedback concerning your experiences with gamselBayes may be sent to the package maintainer at the e-mail address provided by the command help(package = "gamselBayes").

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