Bayesian learning Lab 2

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Linear and polynomial regression

The dataset TempLambohov.txt contains daily average temperatures (in degree Celcius) at Lambohov, Linköping over the course of the year 2019. The response variable is temp and the covariate is

$$time = \frac{the \ number \ of \ days \ since \ the \ beginning \ of \ the \ year}{365}$$

A Bayesian analysis of the following quadratic regression model is to be performed:

$$temp = \beta_0 + \beta_1.time + e, e \sim N(0, \sigma^2)$$

To answer this question a conjugate prior for the linear regression model will be used in which: The joint prior for β and σ^2 :

$$\beta | \sigma^2 \sim \mathcal{N}(\mu_0, \sigma^2 \Omega_0^{-1})$$

 $\sigma^2 \sim Inv - \chi^2(v_0, \sigma^2)$

While the posterior:

$$\beta|\sigma^{2}, y \sim \mathcal{N}(\mu_{n}, \sigma^{2}\Omega_{n}^{-1})$$

$$\sigma^{2} \sim Inv - \chi^{2}(v_{n}, \sigma^{2}_{n})$$
where:
$$\mu_{n} = (X'X\Omega_{0})^{-1} (X'X\hat{\beta} + \Omega_{0}\mu_{0})$$

$$\Omega_{n} = X'X + \Omega_{0}$$

$$v_{n} = v_{0} + n$$

$$v_{n}\sigma^{2} = v_{n}\sigma^{2} + (y'y + \mu'_{0}\Omega_{0}\mu_{0} - \mu'_{n}\Omega_{n}\mu_{n})$$

First we start by reading the files and then we Create the covariate_time vaiable as (the number of days sinvee the beginning of the year / 365)

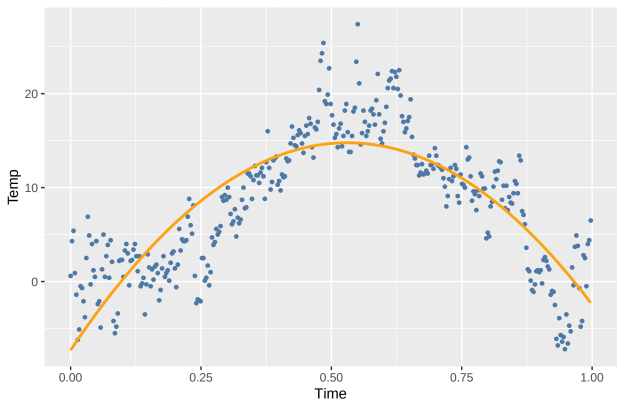
```
#reading the files
df<-read_xlsx("Linkoping2022.xlsx")
#Creating the covariate_time vaiable as
#(the number of days sinvce the beginning of the year / 365)
a<- df$datetime
#begunning of the year
b<- '2022-01-01'
a<-format.POSIXlt(strptime(a,'%Y-%m-%d'))
b<-format.POSIXlt(strptime(b,'%Y-%m-%d'))
#time diff from the
x<-as.vector(difftime(a,b,units='days'))
df$cov_tm<-x/365</pre>
```

\mathbf{A}

Use the conjugate prior for the linear regression model. The prior hyperparameters μ_0 , Ω_0 , v_0 , σ_0^2 shall be set to sensible values. Start with $\mu_0 = (-10, 100, -100)^T$, $\Omega_0 = 0.02.I_3$, $v_0 = 3$, $\sigma_0^2 = 2$. Check if this prior agrees with your prior opinions by simulating draws from the joint prior of all parameters and for every draw compute the regression curve. This gives a collection of regression curves; one for each draw from the prior. Does the collection of curves look reasonable? If not, change the prior hyperparameters until the collection of prior regression curves agrees with your prior beliefs about the regression curve. [Hint: R package mytnorm can be used and your $inv - \chi^2$ simulator of random draws from Lab 1.]

```
#initial parameters
mu_0<- c(-10,100.-100)
Omega_0<-0.02*diag(3)
v 0 <- 3
sigma2_0<-2
# First we fit the linear model by using lm function in R
degree <- 2 # Set the degree of the polynomial
x= df$cov_tm
y= df$temp
model <- lm(y ~ poly(x, degree, raw = TRUE))</pre>
df plt<- data.frame(x=x,y=y)</pre>
z=predict(model)
# Plot the data and regression line
plt <- ggplot(df_plt, aes(x = x, y = y)) +</pre>
  geom_point(color = "#4E79A7", size = 1)+
  geom_line(aes(y = z), color = "#FCA311", size=1 , linetype = 1)+
  labs(x = 'Time', y = 'Temp'
       ,title ='Polynomial Regression with Dgree 2')
## Warning: Using 'size' aesthetic for lines was deprecated in ggplot2 3.4.0.
## i Please use 'linewidth' instead.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
plt
```

Polynomial Regression with Dgree 2



summary(model)

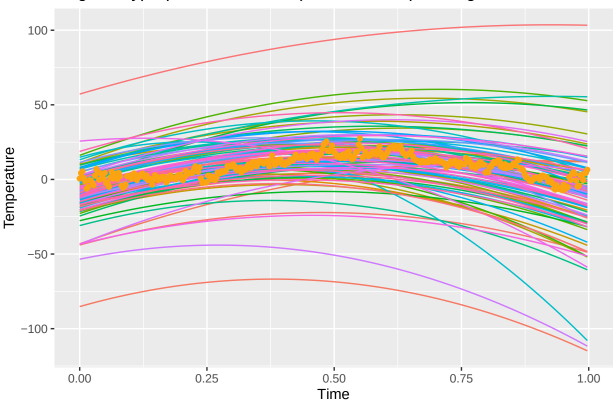
```
##
  lm(formula = y ~ poly(x, degree, raw = TRUE))
##
## Residuals:
        Min
                  1Q
                       Median
                                             Max
## -10.6557 -2.8525 -0.1874
                                2.5052
                                        12.6580
##
## Coefficients:
                                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                                 -7.3039
                                             0.6582
                                                     -11.10
                                                               <2e-16 ***
                                              3.0492
                                                       27.27
## poly(x, degree, raw = TRUE)1 83.1568
                                                               <2e-16 ***
                                                     -26.46
## poly(x, degree, raw = TRUE)2 -78.3093
                                             2.9600
                                                               <2e-16 ***
## ---
                     '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Residual standard error: 4.215 on 362 degrees of freedom
## Multiple R-squared: 0.6726, Adjusted R-squared: 0.6708
## F-statistic: 371.9 on 2 and 362 DF, p-value: < 2.2e-16
```

First we simulating draws from the joint prior of all parameters and for every draw compute the regression curve. This will gives us a collection of regression curves; one for each draw from the prior.

```
mu_0 < -c(-10,100,-100)
Omega_0<-0.02*diag(3)
v_0 <- 3
sigma2_0<-2
# We define our X and Y values
y <- df$temp
X <- cbind(1, df$cov tm, df$cov tm**2)</pre>
# We define the number of Itritaions we want
nIter<-100
# Afunction to find the values of betas and sigma2
lmreg<-function(y, X, mu_0, Omega_0, v_0, sigma2_0, nIter){</pre>
  # Compute posterior hyperparameters
  n = length(y) # Number of observations
  nCovs = dim(X)[2] # Number of covariates
  invOmega_n = solve(Omega_0)
  # The actual sampling
  sigma2Sample = rep(NA, nIter)
  betaSample = matrix(NA, nIter, nCovs)
  for (i in 1:nIter){
    # Simulate from p(sigma2 | y, X)
    sigma2 = rinvchisq(n=1, df=v_0, scale = sigma2_0)
    sigma2Sample[i] = sigma2
    # Simulate from p(beta | sigma2, y, X)
    beta_ = rmvnorm(n=1, mean = mu_0, sigma = sigma2*inv0mega_n)
    betaSample[i,] = beta_
  return(results = list(sigma2Sample = sigma2Sample, betaSample=betaSample))
}
# Finding Predicted values
pred<-lmreg(y, X, mu_0, Omega_0, v_0, sigma2_0, nIter)$betaSample
plot_data<- matrix(data=0,nrow = nrow(X), ncol = nIter)</pre>
for (i in 1:nIter){
  plot_data[,i]<-X%*%(pred)[i,]</pre>
#Plotting the curves
plot_data<- as.data.frame(cbind(time=df$cov_tm,plot_data))</pre>
plot_data <- plot_data %>%
  pivot_longer(
    cols = starts_with("V"), # Columns to pivot
    names_to = "Pred_Iter",
                                      # Name of the new column for variable names
    values_to = "Pred"
                                 # Name of the new column for values
```

```
plt <- ggplot(plot_data)+
  geom_line(aes(x = time, y = Pred, color = Pred_Iter)) +
  geom_point(data = df, aes(x = cov_tm, y = temp), color = "#FCA311") +
  ggtitle("Original Hyperparamters, mu = (-10,100,-100) and sigma = 2") + ylab("Temperature") + xlab("T
  theme(legend.position = "none")
plt</pre>
```

Original Hyperparamters, mu = (-10,100,-100) and sigma = 2



It can be shown that the list of curves that we end up with does not seems to be so reasonable, in this case we can change in the prior hyperparamters until the collection of prior regression curves agrees with our prior beliefs, we choose to change μ_0 values to be (-7.3039, 83.1568, -78.3093) and we also changed the value of σ_0^2 to be .05.

```
mu_0_new<- c(-7.3039, 83.1568, -78.3093)
sigma2_0_new<-.05

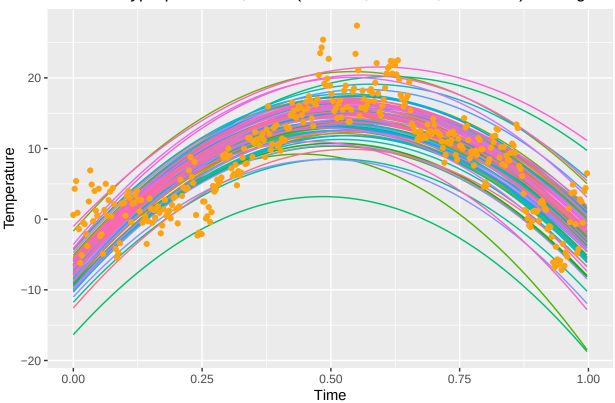
# Finding Predicted values
pred2<-lmreg(y, X, mu_0_new, Omega_0, v_0, sigma2_0_new, nIter)$betaSample

plot_data2<- matrix(data=0,nrow = nrow(X), ncol = nIter)
for (i in 1:nIter){
   plot_data2[,i]<-X%*%(pred2)[i,]
}

#Plotting the curves</pre>
```

```
plot_data2<- as.data.frame(cbind(time=df$cov_tm,plot_data2))</pre>
plot_data2 <- plot_data2 %>%
  pivot_longer(
    cols = starts_with("V"),
                              # Columns to pivot
    names_to = "Pred_Iter",
                                     # Name of the new column for variable names
    values to = "Pred"
                                 # Name of the new column for values
  )
plt <- ggplot(plot_data2)+</pre>
  geom_line(aes(x = time, y = Pred, color = Pred_Iter)) +
  geom_point(data = df, aes(x = cov_tm, y = temp), color = "#FCA311") +
  ggtitle("Modified Hyperparamters, mu = (-7.3039, 83.1568, -78.3093) and sigma = .05") + ylab("Tempera
  theme(legend.position = "none")
plt
```

Modified Hyperparamters, mu = (-7.3039, 83.1568, -78.3093) and sigma =



The resulted regression curves seemes to be more reasonable than the original one.

\mathbf{B}

Write a function that simulate draws from the joint posterior distribution of $\beta_0, \beta_1, \beta_2, \sigma^2$. i- Plot a histogram for each marginal posterior of the parameters. ii- Make a scatter plot of the temperature data and overlay a curve for the posterior median of the regression function $f(time) = E[temp|time] = \beta_0 + \beta_1.time + \beta_2.time^2$, i.e. the median of f(time) is computed for every value of time. In addition, overlay curves for the 95%

equal tail posterior probability intervals of f(time), i.e. the 2.5 and 97.5 posterior percentiles of f(time) is computed for every value of time. Does the posterior probability intervals contain most of the data points? Should they?

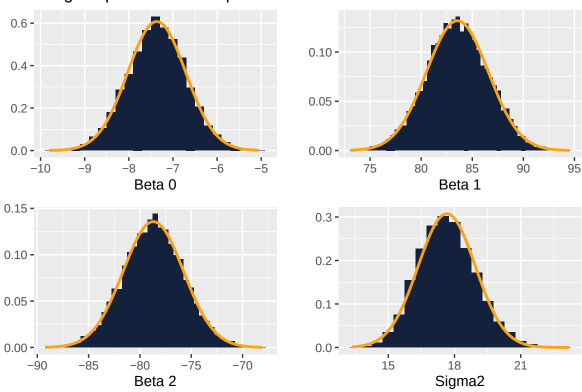
As we want to estimate the uncertainty in the model parameters. Simulating from the joint posterior allows us to obtain a set of plausible values for all the parameters in the model, taking into account the observed data and prior information.

1-Plot a histogram for each marginal posterior of the parameters.

```
simfun<- function(y, X, mu_0, Omega_0, v_0, sigma2_0, simIter){</pre>
  n = length(y) # Number of observations
  nCovs = dim(X)[2] # Number of covariates
  XX = t(X)%*%X
  betaHat <- solve(XX,t(X)%*%y)</pre>
  Omega_n = XX + Omega_0
  mu_n = solve(Omega_n,XX%*%betaHat+Omega_0%*%mu_0)
  v_n = v_0 + n
  sigma2_n = as.numeric((v_0*sigma2_0 + (t(y)%*%y + t(mu_0)%*%0mega_0%*%mu_0 - t(mu_n)%*%0mega_n%*%mu_i)
  invOmega n = solve(Omega n)
  # The actual sampling
  sigma2Sample = rep(NA, simIter)
  betaSample = matrix(NA, simIter, nCovs)
  for (i in 1:simIter){
    # Simulate from p(sigma2 | y, X)
    sigma2 = rinvchisq(n=1, df=v_n, scale = sigma2_n)
    sigma2Sample[i] = sigma2
    # Simulate from p(beta | sigma2, y, X)
    beta_ = rmvnorm(n=1, mean = mu_n, sigma = sigma2*inv0mega_n)
    betaSample[i,] = beta_
 }
 return(results = list(sigma2Sample = sigma2Sample, betaSample=betaSample))
simIter<-10000
res<-simfun(y, X, mu_0, Omega_0, v_0, sigma2_0, simIter)
beta_sim<-as.data.frame(res$betaSample)</pre>
sigma_simz<-data.frame(V1=res$sigma2Sample)</pre>
plt1 <- ggplot(beta_sim,aes(x = V1)) +geom_histogram(aes(y=..density..),</pre>
                                                   linetype=1,
                                                  fill='#14213D', binwidth = 0.2)+
  labs(x='Beta 0',y=' ',title ='Marginal posterior of the parameters')+
  stat_function(fun = dnorm, args = list(mean = mean(beta_sim$V1),
                                          sd = sd(beta_sim$V1)),
                color = "#FCA311", size = 1)
plt2 <- ggplot(beta_sim,aes(x = V2)) +geom_histogram(aes(y=..density..),</pre>
                                                   linetype=1,
                                                   fill='#14213D', binwidth = 0.4)+
 labs(x='Beta 1',y=' ')+
```

```
stat_function(fun = dnorm, args = list(mean = mean(beta_sim$V2),
                                         sd = sd(beta_sim$V2)),
                color = "#FCA311", size = 1)
plt3 <- ggplot(beta_sim,aes(x = V3)) +geom_histogram(aes(y=..density..),</pre>
                                                  linetype=1,
                                                  fill = '#14213D', binwidth = 0.5) +
  labs(x='Beta 2',y=' ',)+
  stat_function(fun = dnorm, args = list(mean = mean(beta_sim$V3),
                                         sd = sd(beta_sim$V3)),
                color = "#FCA311", size = 1)
plt4 <- ggplot(sigma_simz,aes(x = V1)) +geom_histogram(aes(y=..density..),</pre>
                                                      linetype=1,
                                                      fill='#14213D', binwidth = 0.5)+
  labs(x='Sigma2',y=' ',)+
  stat_function(fun = dnorm, args = list(mean = mean(sigma_simz$V1),
                                         sd = sd(sigma_simz$V1)),
                color = "#FCA311", size = 1)
plt1+plt2+plt3+plt4
## Warning: The dot-dot notation ('..density..') was deprecated in ggplot2 3.4.0.
## i Please use 'after_stat(density)' instead.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```

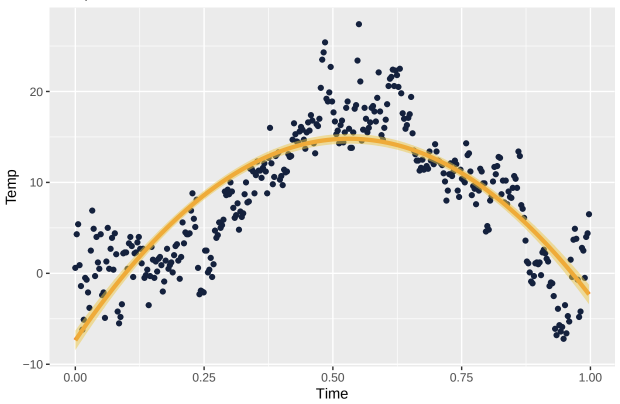
Marginal posterior of the parame



2-Does the posterior probability intervals contain most of the data points? Should they?

```
# Calculating the median value point
median=as.matrix(apply(beta_sim, 2, median))
# we fint the regression model\ P(time) = beta_0 + beta_1 * time + beta_2 * time 2
predicted_response <- X%*% median</pre>
#storing the median values
posterior_median <- apply(predicted_response, 1, median)</pre>
#Finding the predicted interval
prd_int <- data.frame(nrow = nrow(X), nrow = 2)</pre>
colnames(prd_int) <- c("CI_lower", "CI_upper")</pre>
preds<- as.matrix(beta_sim)%*%t(X)</pre>
for(i in 1:nrow(X)){
  data_t <- preds[,i]</pre>
  #Here we have 95% CI using the function quantile
  prd_int[i,] <- quantile(data_t, probs = c(0.05,0.95))</pre>
plt_df=data.frame(x=df$cov_tm,y=df$temp,med=posterior_median)
plt_df= cbind(plt_df,prd_int)
# Calculate posterior median of the predicted response
plt <- ggplot(plt_df, aes(x = x, y = y)) +</pre>
```

The posterior median Curve and 95% CI



The answer is no as the credible intervals do not necessarily contain most of the data points. They are not intended to do so. Credible intervals are used to quantify the uncertainty or variability in the estimated parameter beta in our case. They provide a range of plausible values for the parameter based on the observed data and the model assumptions.

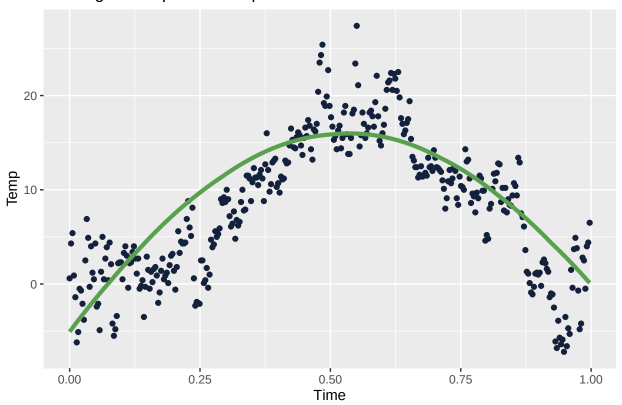
\mathbf{C}

It is of interest to locate the time with the highest expected temperature (i.e. the time where f(time) is maximal). Let's call this value \bar{x} . Use the simulated draws in (b) to simulate from the posterior distribution of \bar{x} . [Hint: the regression curve is a quadratic polynomial. Given each posterior draw of $\beta_0, \beta_1, \beta_2$, you can find a simple formula for \bar{x} .]

To simulate from the posterior distribution of the highest expected temperature \bar{X} we can use the all possible prediction values that we generated in the task before, then by taking the max value we get the point point wise highest expected temperature over time:

```
# Storing the data in one data frame
#Initite the storing vector
het<-c()
#Startingh the for loop
for (i in 1:nrow(X)) {
 het[i] <-max(preds[,i])</pre>
# binding the data into te ploting data frame
plt_df= cbind(plt_df,het)
#Ploting the data
plt <- ggplot(plt_df, aes(x = x, y = y)) +</pre>
  geom_point(color = "#14213D", size = 1.5)+
geom_line(aes(y = het), color = "#59A14F", linetype = 1,size=1.5)+
labs(x = 'Time', y = 'Temp'
,title ='The Highest Expected Temperature'
,color = "Line Legend") +
scale_color_manual(values = c("#14213D","#59A14F")
, labels = c("1","2","3","4"))+
theme(legend.position="bottom")
plt
```

The Highest Expected Temperature



\mathbf{D}

Say now that you want to estimate a polynomial regression of order 8, but you suspect that higher order terms may not be needed, and you worry about overfitting the data. Suggest a suitable prior that mitigates this potential problem. You do not need to compute the posterior. Just write down your prior. [Hint: the task is to specify μ_0 and Ω_0 in a suitable way.]

To mitigate the problem of over fitting when estimating a polynomial resgression of order 10 without being worry of the over fitting, one can use Smoothness/Shrinkage/Regularization of the prior by introducing λ a penalization parameter (See lec 5 Slide 9 Lasso) in this method we have:

$$\beta_j | \sigma^2 \sim N(o, \frac{\sigma^2}{\lambda})$$

Here we have a large values of λ gives smoother fit. More shrinkage. where:

$$\frac{\mu_0}{\Omega_0} = 0$$

$$\Omega_0 = \lambda I$$

Which equivalent to Penalized Likelihood:

$$-2log \ p(\beta|\sigma^2, y, x) \propto (y - X\beta)'(y - X\beta) + \lambda\beta'\beta$$

Thus, the Posterior mean/mode gives ridge regressoin estimator:

$$\tilde{\beta} = (XtX + \lambda I)^{-1}Xty$$

$$if \ XtX = I$$

$$Then$$

$$\tilde{\beta} = \frac{1}{(1+\lambda)}\hat{\beta}$$

We might also be interested to determine the lambda, which could be by performing a cross validation on the test data pf using the Bayesian inference, where to us a prior for λ . we have this hierarchical setup:

$$y|\beta, \sigma^{2}, x \sim N(X\beta, \sigma^{2}I_{n})$$
$$\beta|\sigma^{2}, \lambda \sim N(0, \sigma^{2}\lambda^{-1}I_{m})$$
$$\sigma^{2} \sim Inv - \chi^{2}(v_{o}, \sigma_{o}^{2})$$
$$\lambda \sim Inv - \chi^{2}(\eta_{0}, \lambda_{0})$$
$$so, \ \mu_{o} = 0, \ \Omega = \lambda I_{m}$$

and we have the joint posterior of β , σ^2 , and λ is:

$$\beta|\sigma^{2}, \lambda, y \sim N(\mu_{n}, \sigma^{2}\Omega_{n}^{-1})$$

$$\sigma^{2}|\lambda, y \sim Inv - \chi^{2}(v_{n}, \sigma_{n}^{2})$$

$$p(\lambda|y) \propto \sqrt{\frac{|\Omega_{0}|}{|X^{T}X + \Omega_{0}|}} (\frac{v_{n}\sigma_{n}^{2}}{2})^{\frac{-v_{n}}{2}}.p(\lambda)$$

$$where, \ \Omega_{0} = \lambda I_{m} \ and \ p(\lambda) \ is \ the \ prior \ for \ \lambda \ and :$$

$$\mu_{n} = (X^{T}X + \Omega_{0})^{-1}X^{T}y$$

$$\Omega_{n} = X^{T}X + \Omega_{0}$$

$$v_{n} = v_{0} + n$$

$$v_{n}\sigma_{n}^{2} = v_{0}\sigma_{0}^{2} + y^{T}y - \mu_{0}^{T}\Omega_{n}\mu_{n}$$

2. Posterior approximation for classification with logistic regression

The dataset WomenAtWork.dat contains n = 168 observations on the following eight

variables related to women:

Variable Variable	Data type	Meaning	Role
Work	Binary	Whether or not the woman works	Response y
Constant	1	Constant to the intercept	Feature
HusbandInc	Numeric	Husband's income	Feature
EducYears	Counts	Years of education	Feature
ExpYears	Counts	Years of experience	Feature
Age	Counts	Age	Feature
NSmallChild	Counts	Number of child ≤ 6 years in household	Feature
NBigChild	Counts	Number of child > 6 years in household	Feature

(a) Consider the logistic regression model:

$$\Pr(y = 1 | \mathbf{x}, \beta) = \frac{\exp(\mathbf{x}^T \beta)}{1 + \exp(\mathbf{x}^T \beta)},$$

where y equals 1 if the woman works and 0 if she does not. \mathbf{x} is a 7-dimensional vector containing the seven features (including a 1 to model the intercept). The goal is to approximate the posterior distribution of the parameter vector β with a multivariate normal distribution

$$\beta|\mathbf{y}, \mathbf{x} \sim N\left(\tilde{\beta}, J_{\mathbf{y}}^{-1}(\tilde{\beta})\right),$$

where $\tilde{\beta}$ is the posterior mode and $J(\tilde{\beta}) = -\frac{\partial^2 \ln p(\beta|\mathbf{y})}{\partial \beta \partial \beta^T}|_{\beta = \tilde{\beta}}$ is the negative of the observed Hessian evaluated at the posterior mode. Note that $\frac{\partial^2 \ln p(\beta|\mathbf{y})}{\partial \beta \partial \beta^T}$ is a 7×7 matrix with second derivatives on the diagonal and cross-derivatives

Figure 1: Alt Text

 $\frac{\partial^2 \ln p(\beta|\mathbf{y})}{\partial \beta_i \partial \beta_j}$ on the off-diagonal. You can compute this derivative by hand, but we will let the computer do it numerically for you. Calculate both $\tilde{\beta}$ and $J(\tilde{\beta})$ by using the optim function in R. [Hint: You may use code snippets from my demo of logistic regression in Lecture 6.] Use the prior $\beta \sim \mathcal{N}(0, \tau^2 I)$, where $\tau = 5$.

Present the numerical values of $\tilde{\beta}$ and $J_{\mathbf{y}}^{-1}(\tilde{\beta})$ for the WomenAtWork data. Compute an approximate 95% equal tail posterior probability interval for the regression coefficient to the variable NSmallChild. Would you say that this feature is of importance for the probability that a woman works?

[Hint: You can verify that your estimation results are reasonable by comparing the posterior means to the maximum likelihood estimates, given by: glmModel <- glm(Work ~ 0 + ., data = WomenAtWork, family = binomial).]

- (b) Use your normal approximation to the posterior from (a). Write a function that simulate draws from the posterior predictive distribution of $\Pr(y=1|\mathbf{x})$, where the values of \mathbf{x} corresponds to a 43-year-old woman, with two children (7 and 10 years old), 12 years of education, 8 years of experience, and a husband with an income of 20. Plot the posterior predictive distribution of $\Pr(y=1|\mathbf{x})$ for this woman.
 - [Hints: The R package mvtnorm will be useful. Remember that $\Pr(y = 1|\mathbf{x})$ can be calculated for each posterior draw of β .]
- (c) Now, consider 11 women which all have the same features as the woman in (b). Rewrite your function and plot the posterior predictive distribution for the number of women, out of these 11, that are working. [Hint: Simulate from the binomial distribution, which is the distribution for a sum of Bernoulli random variables.]

Figure 2: Alt Text

Posterior approximation for classification with logistic regression

\mathbf{A}

Firstly we want to calculate the value of $\tilde{\beta}$ the posterior mode and the negative of the observed 7x7 hessian evaluated at the posterior mode $J(\tilde{\beta}) = -\frac{\partial^2 \ln p(\beta|y)}{\partial \beta \partial \beta^T}|_{\beta = \tilde{\beta}}$. Using code snippets from my demo of logistic regression in Lecture 6. First we want to calculate the vale of $\tilde{\beta}$ and $J(\tilde{\beta}) = -\frac{\partial^2 \ln p(\beta|y)}{\partial \beta \partial \beta^T}|_{\beta = \tilde{\beta}}$ by using the optim, Note that we have $\tau = 2$ and $\beta \sim N(0, \frac{1}{\lambda}I)$.

```
# First we want to calculate the vale of beta and the Jacopiang
#inv of beta by using three optim function and the code from the lec notes
# Note that we have tau = 2 and prior beta follows N(0, tau^2I)
### Select Logistic or Probit regression and install packages ###
Probit <- 0
### Prior and data inputs ###
Covs <- c(2:8) # Select which covariates/features to include
standardize <- F # If TRUE, covariates/features are standardized
                  #to mean O and variance 1
lambda <- 4 # scaling factor for the prior of beta in our case tau = 2
# Loading out data set
wat<-read.table("WomenAtWork.dat",header = T) # read data from file
Nobs <- dim(wat)[1] # number of observations
y \leftarrow wat[1] \# y=1 \ if \ the \ women \ is \ working, \ otherwise \ y=0.
X <- as.matrix(wat[,Covs]) # Covs matrix 7*7</pre>
Xnames <- colnames(X)</pre>
# Standraizing the covs matrix
if (standardize){
 Index <- 2:(length(Covs)-1)</pre>
 X[,Index] <- scale(X[,Index])</pre>
Npar \leftarrow dim(X)[2]
# This is to add y variable as binary response and adding
#intercept, for now it's not needed
# for (ii in 1:Nobs){
  if (wat$quality[ii] > 5){
     y[ii] \leftarrow 1
#
#
# }
# wat <- data.frame(intercept=rep(1,Nobs),wat) # add intercept
# Setting up the prior
mu <- as.matrix(rep(0,Npar)) # Prior mean vector</pre>
Sigma <- (1/lambda)*diag(Npar) # Prior covariance matrix
# Functions that returns the log posterior for the logistic and probit regression.
# First input argument of this function must be the parameters we optimize on,
# i.e. the regression coefficients beta.
```

```
LogPostLogistic <- function(betas,y,X,mu,Sigma){</pre>
  linPred <- X%*%betas;</pre>
  logLik <- sum( linPred*y - log(1 + exp(linPred)) );</pre>
  #if (abs(logLik) == Inf) logLik = -20000; # Likelihood is
  #not finite, stear the optimizer away from here!
  logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
 return(logLik + logPrior)
}
LogPostProbit <- function(betas,y,X,mu,Sigma){</pre>
  linPred <- X%*%betas;</pre>
  SmallVal <- .Machine$double.xmin</pre>
  logLik <- sum(y*log(pnorm(linPred)+SmallVal) +</pre>
                   (1-y)*log(1-pnorm(linPred)+SmallVal) )
 logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
  return(logLik + logPrior)
# Select the initial values for beta
initVal <- matrix(0,Npar,1)</pre>
if (Probit==1){
  logPost = LogPostProbit;
} else{
  logPost = LogPostLogistic;
# The argument control is a list of options to
#the optimizer optim, where fnscale=-1 means that we minimize
# the negative log posterior. Hence, we maximize the log posterior.
OptimRes <- optim(initVal,logPost,gr=NULL,y,X,mu,
                   Sigma,method=c("BFGS"),control=list(fnscale=-1),hessian=TRUE)
# Printing the results to the screen
names(OptimRes$par) <- Xnames # Naming the coefficient by covariates</pre>
# Computing approximate standard deviations.
approxPostStd <- sqrt(diag(solve(-OptimRes$hessian)))</pre>
names(approxPostStd) <- Xnames # Naming the coefficient by covariates</pre>
print('The posterior mode is:')
## [1] "The posterior mode is:"
print(OptimRes$par)
##
                 [,1]
## [1,] -0.056386120
## [2,] -0.031827146
## [3,] 0.122159465
## [4,] 0.117428313
## [5,] -0.034825257
## [6,] -0.843613303
```

```
## [7,] -0.004864958
## attr(,"names")
## [1] "Constant"
                     "HusbandInc"
                                  "EducYears"
                                                 "ExpYears"
                                                               "Age"
## [6] "NSmallChild" "NBigChild"
print('The Hessian Matrix:')
## [1] "The Hessian Matrix:"
print(OptimRes$hessian)
##
                            [,2]
                                         [,3]
                                                      [,4]
                                                                             [,6]
                [,1]
                                                                  [,5]
## [1,]
         -30.680116
                       -529.3142
                                  -327.50822
                                               -236.69284 -1090.0231
                                                                        -6.677466
## [2,]
        -529.314175 -13341.9063
                                 -6766.54274
                                              -4843.93682 -21997.1466 -120.512938
## [3,]
        -327.508217
                     -6766.5427
                                 -4173.35502
                                              -2920.20417 -13322.2994
                                                                       -89.131991
## [4,]
        -236.692843 -4843.9368 -2920.20417 -3210.07312 -10128.0426 -52.468406
## [5,] -1090.023088 -21997.1466 -13322.29943 -10128.04260 -46241.4185 -222.328139
## [6,]
                      -120.5129
          -6.677466
                                   -89.13199
                                                -52.46841
                                                            -222.3281 -12.483127
## [7,]
         -35.842534
                      -680.3382
                                 -427.53675
                                               -263.56474 -1376.1948
                                                                        -7.970026
##
                [,7]
## [1,]
         -35.842534
        -680.338176
## [2,]
## [3,]
        -427.536753
## [4,]
        -263.564737
## [5,] -1376.194769
## [6,]
          -7.970026
## [7,]
         -95.588839
print('The approximate posterior standard deviation is:')
## [1] "The approximate posterior standard deviation is:"
print(approxPostStd)
      Constant HusbandInc
##
                            EducYears
                                         ExpYears
                                                          Age NSmallChild
##
   0.48111702
               0.02091553
                           ##
    NBigChild
##
  0.14263857
Now we compare with the results from the regression model: glmModel < -glm(Work \sim 0 + ., data =
WomenAtWork, family = binomial
glmModel<- glm(Work ~ 0 + ., data = wat, family = binomial)</pre>
summary(glmModel)
##
## Call:
## glm(formula = Work ~ 0 + ., family = binomial, data = wat)
## Coefficients:
```

```
##
               Estimate Std. Error z value Pr(>|z|)
                0.02263
                           1.93083
                                     0.012 0.990649
## Constant
               -0.03796
## HusbandInc
                           0.02229
                                    -1.703 0.088573 .
## EducYears
                0.18447
                           0.10007
                                     1.844 0.065253
## ExpYears
                0.12132
                           0.03353
                                     3.618 0.000297 ***
                                    -1.462 0.143686
               -0.04858
                           0.03323
## Age
                                    -3.064 0.002187 **
## NSmallChild -1.56485
                           0.51078
## NBigChild
               -0.02526
                           0.17716
                                    -0.143 0.886618
## ---
                  0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Signif. codes:
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 182.99
                                      degrees of freedom
##
                              on 132
## Residual deviance: 146.73 on 125 degrees of freedom
## AIC: 160.73
##
## Number of Fisher Scoring iterations: 4
```

The Coefficients of model results about shows relationship between each predictor variable and the log-odds of the outcome variable. variables like HusbandInc, Age, NSmallChild and NBigChild have a negative impact on the y variable in which the direction of the relationship, while the rest have a positive impact, our results from the teammate tell the same direction of the model estimates (i.e variables with negative impact HusbandInc, Age, NSmallChild and NBigChild). However the magnitude of the coefficient which indicates the strength of the relationship is a bit different.

Now we Compute an approximate 95% equal tail posterior probability interval for the regression coefficient to the variable NSmallChild

```
# We use the function rmvnorm to generate the
#variates using OptimRes$par as our mean and approxPostStd as sigma
postmode<-as.matrix(OptimRes$par[,1])
poststd<-solve(-OptimRes$hessian)
watvar<-data.frame(rmvnorm(n=10000,mean = postmode, sigma = poststd))
#For a 95% CI, you would typically calculate
#the lower and upper bounds at quantiles 0.025 and 0.975, respectively.
print('An approximate 95% equal tail posterior probability
    interval for the regression coeffcient to the variable NSmallChild is:')</pre>
```

[1] "An approximate 95% equal tail posterior probability\n interval for the regression coeffcient

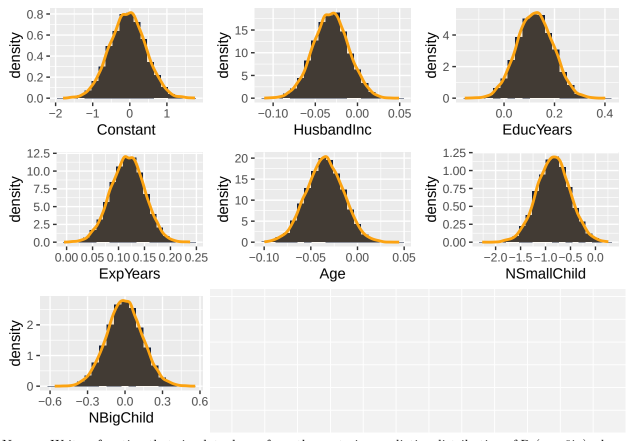
```
print(quantile(watvar[,6],c(0.025,.975)))
## 2.5% 97.5%
## -1.4780606 -0.2128472
```

The results of the CI tell us that the NSmallChild regression coefficient have a significant impact on the log-odds of the outcome variable as we can see the coefficient fall between the lower and the upper bound of the CI, it also important that the 0 is not included in the lower and the upper bounds for the CI which tells that the paramter has a significant impact, the direction of this impact in negative and have the highest magnitude in the model coefficients. We can also observe that 0 is not included in our interval which means the coefficient is important.

\mathbf{B}

Now we write a function that simulate draws from the posterior predictive distribution of $\Pr(y = 0|x)$, where the values of x corresponds to a 40-year-old woman, with two children (4 and 7 years old), 11 years of education, 7 years of experience, and a husband with an income of 18. Plot the posterior predictive distribution of $\Pr(y = 0|x)$ for this woman. First we estimate the values of β using the normal approximation to the posterior from (a) above, the below graph shows the distribution of the estimated βs :

```
betas<-rmvnorm(n=10000,mean = postmode, sigma = poststd)</pre>
# ploting the beta distribution
p data<- as.data.frame(betas)</pre>
colnames(p_data)<-colnames(wat[2:8])</pre>
names<-colnames(p_data)</pre>
p_fun<- function(coln){</pre>
          plt <- ggplot(p_data,aes_string(x = coln)) +</pre>
               geom_histogram(aes(y=..density..),linetype=1
                               fill='#14213D', bins = 20)+
               geom_density(alpha=.2,color="#FCA311",size=1,fill="#FCA311")
plt
}
plot(arrangeGrob(grobs = lapply(names, p_fun)))
## Warning: 'aes_string()' was deprecated in ggplot2 3.0.0.
## i Please use tidy evaluation idioms with 'aes()'.
## i See also 'vignette("ggplot2-in-packages")' for more information.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```



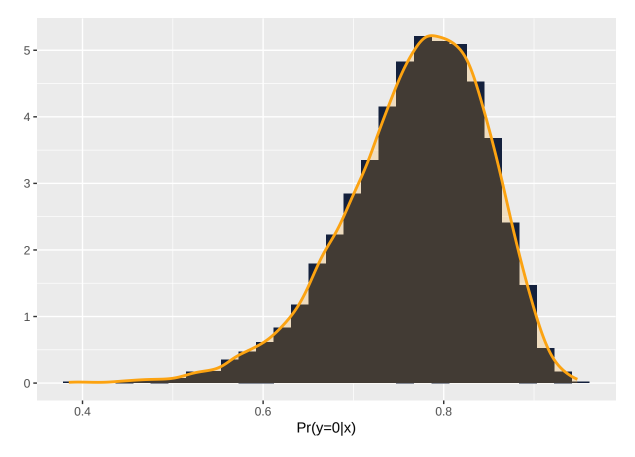
Now we Write a function that simulate draws from the posterior predictive distribution of Pr(y = 0jx), where the values of x

```
pred_prob<- function(ndraws,x_new){</pre>
         ### Select Logistic or Probit regression and install packages ###
        Probit <- 0
        ### Prior and data inputs ###
        Covs <- c(2:8) # Select which covariates/features to include
        standardize <- F # If TRUE, covariates/features
         #are standardized to mean 0 and variance 1
        lambda <- 2 # scaling factor for the prior of beta in our case tau = 2
        # Loading out data set
        wat<-read.table("WomenAtWork.dat",header = T) # read data from file</pre>
        Nobs <- dim(wat)[1] # number of observations
        y \leftarrow wat[1] \# y=1 \ if \ the \ women \ is \ working, \ otherwise \ y=0.
        X <- as.matrix(wat[,Covs]) # Covs matrix 7*7</pre>
        Xnames <- colnames(X)</pre>
         # Standraizing the covs matrix
        if (standardize){
           Index <- 2:(length(Covs)-1)</pre>
          X[,Index] <- scale(X[,Index])</pre>
        }
        Npar \leftarrow dim(X)[2]
        # Setting up the prior
        mu <- as.matrix(rep(0,Npar)) # Prior mean vector</pre>
        Sigma <- (lambda)^2 *diag(Npar) # Prior covariance matrix</pre>
        LogPostLogistic <- function(betas,y,X,mu,Sigma){</pre>
```

```
linPred <- X%*%betas;</pre>
  logLik <- sum( linPred*y - log(1 + exp(linPred)) );</pre>
   if (abs(logLik) == Inf){
               logLik = -20000
      }# Likelihood is not finite, stear the optimizer away from here!
  logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
  return(logLik + logPrior)
# Not in use we change the value to 0 at the beginning of the code
LogPostProbit <- function(betas,y,X,mu,Sigma){</pre>
  linPred <- X%*%betas;</pre>
  SmallVal <- .Machine$double.xmin</pre>
  logLik <- sum(y*log(pnorm(linPred)+SmallVal) +</pre>
                  (1-y)*log(1-pnorm(linPred)+SmallVal))
  logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
  return(logLik + logPrior)
# Select the initial values for beta
initVal <- matrix(0,Npar,1)</pre>
if (Probit==1){
 logPost = LogPostProbit;
} else{
  logPost = LogPostLogistic;
# The argument control is a list of options to the optimizer optim,
#where fnscale=-1 means that we minimize
# the negative log posterior. Hence, we maximize the log posterior.
OptimRes <- optim(initVal,logPost,gr=NULL,y,X,mu,Sigma,method=c("BFGS")
                  ,control=list(fnscale=-1),hessian=TRUE)
postmode<-as.matrix(OptimRes$par[,1])</pre>
poststd<- solve(-OptimRes$hessian)</pre>
x_new<-as.matrix(x_new,ncol=1)</pre>
betas<-rmvnorm(n=ndraws,mean = postmode, sigma = poststd)</pre>
# Finding the value y givan the new Xs
pr_y<-data.frame(x=betas%*%x_new)</pre>
# Finding the probabilities using the logistics function
pr_y$x_logit<-1-(1/(1+exp(-pr_y$x)))
# Ploting the dataset
plt <- ggplot(pr_y,aes(x = x_logit)) +geom_histogram(aes(y=..density..),</pre>
         linetype=1, fill='#14213D')+
          geom_density(alpha=.2,color="#FCA311",size=1,fill="#FCA311")+
         labs(x='Pr(y=0|x)',y=' ',)
plt
```

```
pred_prob(10000,c(1,18,11,7,40,1,1))
```

'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.



consider 13 women which all have the same features as the woman in (b). Rewrite your function and plot the posterior predictive distribution for the number of women, out of these 13, that are not working.

```
pred_prob2<- function(ndraws,x_new){</pre>
      ### Select Logistic or Probit regression and install packages ###
      Probit <- 0
      ### Prior and data inputs ###
      Covs <- c(2:8) # Select which covariates/features to include
      standardize <- F # If TRUE, covariates/features are
                         #standardized to mean O and variance 1
      lambda <- 2 # scaling factor for the prior of beta in our case tau = 2
      # Loading out data set
      wat <- read.table ("Women At Work.dat", header = T) # read data from file
      Nobs <- dim(wat)[1] # number of observations
      y \leftarrow wat[1] \# y=1 \ if \ the \ women \ is \ working, \ otherwise \ y=0.
      X <- as.matrix(wat[,Covs]) # Covs matrix 7*7</pre>
      Xnames <- colnames(X)</pre>
      # Standraizing the covs matrix
      if (standardize){
        Index <- 2:(length(Covs)-1)</pre>
        X[,Index] <- scale(X[,Index])</pre>
      Npar \leftarrow dim(X)[2]
      # Setting up the prior
      mu <- as.matrix(rep(0,Npar)) # Prior mean vector</pre>
      Sigma <- (lambda)^2 *diag(Npar) # Prior covariance matrix
```

```
LogPostLogistic <- function(betas,y,X,mu,Sigma){</pre>
  linPred <- X%*%betas;</pre>
  logLik <- sum( linPred*y - log(1 + exp(linPred)) );</pre>
   if (abs(logLik) == Inf){
                logLik = -20000
  # Likelihood is not finite, stear the optimizer away from here!
  logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
  return(logLik + logPrior)
# Not in use we change the value to 0 at the beginning of the code
LogPostProbit <- function(betas, y, X, mu, Sigma){</pre>
  linPred <- X%*%betas;</pre>
  SmallVal <- .Machine$double.xmin</pre>
  logLik <- sum(y*log(pnorm(linPred)+SmallVal) +</pre>
                  (1-y)*log(1-pnorm(linPred)+SmallVal))
  logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
  return(logLik + logPrior)
}
# Select the initial values for beta
initVal <- matrix(0,Npar,1)</pre>
if (Probit==1){
 logPost = LogPostProbit;
} else{
  logPost = LogPostLogistic;
# The argument control is a list of options to the optimizer optim,
#where fnscale=-1 means that we minimize
# the negative log posterior. Hence, we maximize the log posterior.
OptimRes <- optim(initVal,logPost,gr=NULL,y,X,mu,Sigma,method=c("BFGS")</pre>
                  ,control=list(fnscale=-1),hessian=TRUE)
postmode<-as.matrix(OptimRes$par[,1])</pre>
poststd<- solve(-OptimRes$hessian)</pre>
x_new<-as.matrix(x_new,ncol=1)</pre>
betas<-rmvnorm(n=ndraws,mean = postmode, sigma = poststd)</pre>
# Finding the value y givan the new Xs
pr_y<-data.frame(x=betas%*%x_new)</pre>
# Finding the probabilities using the logistics function
pr_y$x_logit<-1/(1+exp(-pr_y$x))
#Adding the clasifier
pr_y$job_flag <- ifelse(pr_y$x_logit <= 0.5, 0, 1)</pre>
plt <- ggplot(pr_y, aes(x = x_logit, y = job_flag)) +</pre>
  geom_point(colour="#14213D") +
  # stat_smooth(method="glm",colour="#FCA311",
                alpha = 0.5, se=FALSE, fullrange=TRUE,
               method.args = list(family=binomial)) +
  xlab("Predictor") + xlim(c(0,1))+
  ylab("Probability of Outcome") +
  ggtitle("Logistic Regression function with 0.5 as decision boundary")+
```

pred_prob2(10000,c(1,18,11,7,40,1,1))

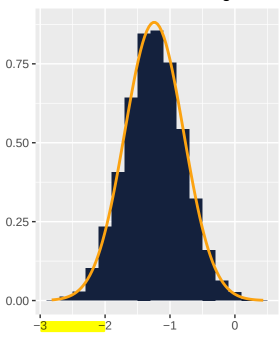


0.25

0.50

Predictor

Posterior predictive distribution for , out of these 13, that are not working





References:

0.00 -

0.00

1- Bertil Wegmann (2023). Bayesian Learning [Lecture notes]. 732A73, Department of Computer and Information Science,LiU University.

1.00

0.75

Code Appendix

```
knitr::opts_chunk$set(echo = TRUE)
set.seed(123)
library(ggplot2)
library(mvtnorm)
library(readxl)
library(LaplacesDemon)
library(patchwork)
library(gridExtra)
library(matrixStats)
library(tidyverse)
#reading the files
df<-read_xlsx("Linkoping2022.xlsx")</pre>
#Creating the covariate_time vaiable as
#(the number of days sinvce the beginning of the year / 365)
a<- df$datetime</pre>
#begunning of the year
b<- '2022-01-01'
a<-format.POSIX1t(strptime(a,'%Y-\m-\mathcal{d}'))
b<-format.POSIX1t(strptime(b,'%Y-\m-\mathcal{d}'))
#time diff from the
x<-as.vector(difftime(a,b,units='days'))</pre>
df$cov_tm<-x/365
#initial parameters
mu_0 < -c(-10, 100. -100)
Omega_0 < -0.02 * diag(3)
v_0 <- 3
sigma2_0<-2
# First we fit the linear model by using lm function in R
degree <- 2 # Set the degree of the polynomial
x= df$cov tm
y= df$temp
model <- lm(y ~ poly(x, degree, raw = TRUE))</pre>
df_plt<- data.frame(x=x,y=y)</pre>
z=predict(model)
# Plot the data and regression line
plt <- ggplot(df_plt, aes(x = x, y = y)) +</pre>
  geom_point(color = "#4E79A7", size = 1)+
  geom_line(aes(y = z), color = "#FCA311", size=1 , linetype = 1)+
  labs(x = 'Time', y = 'Temp'
        ,title ='Polynomial Regression with Dgree 2')
plt
summary(model)
mu_0 < -c(-10, 100, -100)
Omega 0 < -0.02 * diag(3)
v_0 <- 3
sigma2_0<-2
```

```
# We define our X and Y values
y <- df$temp
X <- cbind(1, df$cov tm, df$cov tm**2)</pre>
# We define the number of Itritaions we want
nIter<-100
# Afunction to find the values of betas and sigma2
lmreg<-function(y, X, mu_0, Omega_0, v_0, sigma2_0, nIter){</pre>
  # Compute posterior hyperparameters
  n = length(y) # Number of observations
  nCovs = dim(X)[2] # Number of covariates
  invOmega_n = solve(Omega_0)
  # The actual sampling
  sigma2Sample = rep(NA, nIter)
  betaSample = matrix(NA, nIter, nCovs)
  for (i in 1:nIter){
    # Simulate from p(sigma2 | y, X)
    sigma2 = rinvchisq(n=1, df=v_0, scale = sigma2_0)
    sigma2Sample[i] = sigma2
    # Simulate from p(beta | sigma2, y, X)
    beta_ = rmvnorm(n=1, mean = mu_0, sigma = sigma2*inv0mega_n)
    betaSample[i,] = beta_
  }
  return(results = list(sigma2Sample = sigma2Sample, betaSample=betaSample))
# Finding Predicted values
pred<-lmreg(y, X, mu_0, Omega_0, v_0, sigma2_0, nIter)$betaSample
plot_data<- matrix(data=0,nrow = nrow(X), ncol = nIter)</pre>
for (i in 1:nIter){
  plot_data[,i]<-X%*%(pred)[i,]</pre>
#Plotting the curves
plot_data<- as.data.frame(cbind(time=df$cov_tm,plot_data))</pre>
plot_data <- plot_data %>%
  pivot_longer(
   cols = starts_with("V"), # Columns to pivot
   names_to = "Pred_Iter",
                                    # Name of the new column for variable names
    values_to = "Pred"
                                 # Name of the new column for values
  )
plt <- ggplot(plot_data)+</pre>
  geom_line(aes(x = time, y = Pred, color = Pred_Iter)) +
  geom_point(data = df, aes(x = cov_tm, y = temp), color = "#FCA311") +
```

```
ggtitle("Original Hyperparamters, mu = (-10,100,-100) and sigma = 2") + ylab("Temperature") + xlab("T
  theme(legend.position = "none")
plt
mu_0_new < -c(-7.3039, 83.1568, -78.3093)
sigma2_0_new < -.05
# Finding Predicted values
pred2<-lmreg(y, X, mu_0_new, Omega_0, v_0, sigma2_0_new, nIter)$betaSample
plot_data2<- matrix(data=0,nrow = nrow(X), ncol = nIter)</pre>
for (i in 1:nIter){
  plot_data2[,i]<-X%*%(pred2)[i,]
#Plotting the curves
plot_data2<- as.data.frame(cbind(time=df$cov_tm,plot_data2))</pre>
plot_data2 <- plot_data2 %>%
  pivot_longer(
    cols = starts_with("V"), # Columns to pivot
                               # Name of the new column for variable names
   names_to = "Pred_Iter",
    values_to = "Pred"
                                # Name of the new column for values
plt <- ggplot(plot_data2)+</pre>
  geom_line(aes(x = time, y = Pred, color = Pred_Iter)) +
  geom_point(data = df, aes(x = cov_tm, y = temp), color = "#FCA311") +
  ggtitle("Modified Hyperparamters, mu = (-7.3039, 83.1568, -78.3093) and sigma = .05") + ylab("Tempera
 theme(legend.position = "none")
plt
simfun<- function(y, X, mu_0, Omega_0, v_0, sigma2_0, simIter){</pre>
  n = length(y) # Number of observations
  nCovs = dim(X)[2] # Number of covariates
  XX = t(X)%*%X
  betaHat <- solve(XX,t(X)%*%y)</pre>
  Omega_n = XX + Omega_0
  mu_n = solve(Omega_n,XX%*%betaHat+Omega_0%*%mu_0)
  v_n = v_0 + n
  sigma2_n = as.numeric((v_0*sigma2_0 + (t(y)%*%y + t(mu_0)%*%0mega_0%*%mu_0 - t(mu_n)%*%0mega_n%*%mu_i)
  invOmega_n = solve(Omega_n)
  # The actual sampling
  sigma2Sample = rep(NA, simIter)
  betaSample = matrix(NA, simIter, nCovs)
  for (i in 1:simIter){
    # Simulate from p(sigma2 | y, X)
    sigma2 = rinvchisq(n=1, df=v_n, scale = sigma2_n)
    sigma2Sample[i] = sigma2
```

```
# Simulate from p(beta | sigma2, y, X)
    beta_ = rmvnorm(n=1, mean = mu_n, sigma = sigma2*inv0mega_n)
    betaSample[i,] = beta_
  }
 return(results = list(sigma2Sample = sigma2Sample, betaSample=betaSample))
}
simIter<-10000
res<-simfun(y, X, mu_0, Omega_0, v_0, sigma2_0, simIter)
beta_sim<-as.data.frame(res$betaSample)</pre>
sigma_simz<-data.frame(V1=res$sigma2Sample)</pre>
plt1 <- ggplot(beta_sim,aes(x = V1)) +geom_histogram(aes(y=..density..),</pre>
                                                   linetype=1,
                                                   fill='#14213D', binwidth = 0.2)+
  labs(x='Beta 0',y=' ',title ='Marginal posterior of the parameters')+
  stat_function(fun = dnorm, args = list(mean = mean(beta_sim$V1),
                                           sd = sd(beta_sim$V1)),
                color = "#FCA311", size = 1)
plt2 <- ggplot(beta_sim,aes(x = V2)) +geom_histogram(aes(y=..density..),</pre>
                                                   linetype=1,
                                                   fill = '#14213D', binwidth = 0.4) +
  labs(x='Beta 1',y=' ')+
  stat_function(fun = dnorm, args = list(mean = mean(beta_sim$V2),
                                          sd = sd(beta sim V2)),
                color = "#FCA311", size = 1)
plt3 <- ggplot(beta_sim,aes(x = V3)) +geom_histogram(aes(y=..density..),</pre>
                                                   linetype=1,
                                                   fill='#14213D', binwidth = 0.5)+
  labs(x='Beta 2',y=' ',)+
  stat_function(fun = dnorm, args = list(mean = mean(beta_sim$V3),
                                          sd = sd(beta_sim$V3)),
                color = "#FCA311", size = 1)
plt4 <- ggplot(sigma_simz,aes(x = V1)) +geom_histogram(aes(y=..density..),</pre>
                                                       linetype=1.
                                                       fill = '#14213D', binwidth = 0.5) +
  labs(x='Sigma2',y=' ',)+
  stat_function(fun = dnorm, args = list(mean = mean(sigma_simz$V1),
                                          sd = sd(sigma_simz$V1)),
                color = "#FCA311", size = 1)
plt1+plt2+plt3+plt4
# Calculating the median value point
median=as.matrix(apply(beta_sim, 2, median))
# we fint the regression model P(time)=beta_0+beta_1*time+beta_2*time2
predicted_response <- X%*% median</pre>
#storing the median values
posterior_median <- apply(predicted_response, 1, median)</pre>
```

```
#Finding the predicted interval
prd_int <- data.frame(nrow = nrow(X), nrow = 2)</pre>
colnames(prd_int) <- c("CI_lower", "CI_upper")</pre>
preds<- as.matrix(beta sim)%*%t(X)</pre>
for(i in 1:nrow(X)){
  data_t <- preds[,i]</pre>
  #Here we have 95% CI using the function quantile
 prd int[i,] \leftarrow quantile(data t, probs = c(0.05,0.95))
plt_df=data.frame(x=df$cov_tm,y=df$temp,med=posterior_median)
plt_df= cbind(plt_df,prd_int)
# Calculate posterior median of the predicted response
plt \leftarrow ggplot(plt_df, aes(x = x, y = y)) +
  geom_point(color = "#14213D", size = 1.5)+
  geom_line(aes(y = med), color = "#F28E2B", linetype = 1,size=1.5)+
  geom_ribbon(aes(ymin = CI_lower, ymax = CI_upper)
               , alpha = 0.5,fill = "#EDC948")+
  labs(x = 'Time', y = 'Temp'
       ,title ='The posterior median Curve and 95% CI')
plt
# Storing the data in one data frame
#Initite the storing vector
het<-c()
#Startingh the for loop
for (i in 1:nrow(X)) {
 het[i]<-max(preds[,i])</pre>
# binding the data into te ploting data frame
plt_df= cbind(plt_df,het)
#Ploting the data
plt \leftarrow ggplot(plt_df, aes(x = x, y = y)) +
  geom_point(color = "#14213D", size = 1.5)+
geom_line(aes(y = het), color = "#59A14F", linetype = 1,size=1.5)+
labs(x = 'Time', y = 'Temp'
,title ='The Highest Expected Temperature'
,color = "Line Legend") +
scale_color_manual(values = c("#14213D","#59A14F")
, labels = c("1","2","3","4"))+
theme(legend.position="bottom")
plt
# First we want to calculate the vale of beta and the Jacopiang
#inv of beta by using thee optim function and the code from the lec notes
# Note that we have tau = 2 and prior beta follows N(0,tau^2I)
```

```
### Select Logistic or Probit regression and install packages ###
Probit <- 0
### Prior and data inputs ###
Covs <- c(2:8) # Select which covariates/features to include
standardize <- F # If TRUE, covariates/features are standardized
                  #to mean O and variance 1
lambda <- 4 # scaling factor for the prior of beta in our case tau = 2
# Loading out data set
wat<-read.table("WomenAtWork.dat",header = T) # read data from file</pre>
Nobs <- dim(wat)[1] # number of observations
y \leftarrow wat[1] \# y=1 \ if \ the \ women \ is \ working, \ otherwise \ y=0.
X <- as.matrix(wat[,Covs]) # Covs matrix 7*7</pre>
Xnames <- colnames(X)</pre>
# Standraizing the covs matrix
if (standardize){
  Index <- 2:(length(Covs)-1)</pre>
 X[,Index] <- scale(X[,Index])</pre>
Npar \leftarrow dim(X)[2]
# This is to add y variable as binary response and adding
#intercept, for now it's not needed
# for (ii in 1:Nobs){
  if (wat quality[ii] > 5) {
     y[ii] <- 1
#
# }
# wat <- data.frame(intercept=rep(1,Nobs),wat) # add intercept
# Setting up the prior
mu <- as.matrix(rep(0,Npar)) # Prior mean vector</pre>
Sigma <- (1/lambda)*diag(Npar) # Prior covariance matrix
# Functions that returns the log posterior for the logistic and probit regression.
# First input argument of this function must be the parameters we optimize on,
# i.e. the regression coefficients beta.
LogPostLogistic <- function(betas,y,X,mu,Sigma){</pre>
 linPred <- X%*%betas;</pre>
  logLik <- sum( linPred*y - log(1 + exp(linPred)) );</pre>
  #if (abs(logLik) == Inf) logLik = -20000; # Likelihood is
  #not finite, stear the optimizer away from here!
 logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
 return(logLik + logPrior)
LogPostProbit <- function(betas,y,X,mu,Sigma){</pre>
  linPred <- X%*%betas;</pre>
```

```
SmallVal <- .Machine$double.xmin</pre>
  logLik <- sum(y*log(pnorm(linPred)+SmallVal) +</pre>
                   (1-y)*log(1-pnorm(linPred)+SmallVal) )
  logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
  return(logLik + logPrior)
# Select the initial values for beta
initVal <- matrix(0,Npar,1)</pre>
if (Probit==1){
  logPost = LogPostProbit;
} else{
  logPost = LogPostLogistic;
# The argument control is a list of options to
#the optimizer optim, where fnscale=-1 means that we minimize
# the negative log posterior. Hence, we maximize the log posterior.
OptimRes <- optim(initVal,logPost,gr=NULL,y,X,mu,</pre>
                   Sigma, method=c("BFGS"), control=list(fnscale=-1), hessian=TRUE)
# Printing the results to the screen
names(OptimRes$par) <- Xnames # Naming the coefficient by covariates</pre>
# Computing approximate standard deviations.
approxPostStd <- sqrt(diag(solve(-OptimRes$hessian)))</pre>
names(approxPostStd) <- Xnames # Naming the coefficient by covariates</pre>
print('The posterior mode is:')
print(OptimRes$par)
print('The Hessian Matrix:')
print(OptimRes$hessian)
print('The approximate posterior standard deviation is:')
print(approxPostStd)
glmModel<- glm(Work ~ 0 + ., data = wat, family = binomial)</pre>
summary(glmModel)
# We use the function rmunorm to generate the
\#variates\ using\ OptimRes\$par\ as\ our\ mean\ and\ approx PostStd\ as\ sigma
postmode<-as.matrix(OptimRes$par[,1])</pre>
poststd<-solve(-OptimRes$hessian)</pre>
watvar<-data.frame(rmvnorm(n=10000,mean = postmode, sigma = poststd))</pre>
#For a 95% CI, you would typically calculate
#the lower and upper bounds at quantiles 0.025 and 0.975, respectively.
print('An approximate 95% equal tail posterior probability
      interval for the regression coeffcient to the variable NSmallChild is:')
print(quantile(watvar[,6],c(0.025,.975)))
betas<-rmvnorm(n=10000,mean = postmode, sigma = poststd)
# ploting the beta distribution
p_data<- as.data.frame(betas)</pre>
colnames(p_data)<-colnames(wat[2:8])</pre>
names<-colnames(p_data)</pre>
p_fun<- function(coln){</pre>
```

```
plt <- ggplot(p_data,aes_string(x = coln)) +</pre>
              geom_histogram(aes(y=..density..),linetype=1
                             fill='#14213D', bins = 20)+
              geom_density(alpha=.2,color="#FCA311",size=1,fill="#FCA311")
plt
plot(arrangeGrob(grobs = lapply(names, p fun)))
pred_prob<- function(ndraws,x_new){</pre>
        ### Select Logistic or Probit regression and install packages ###
       Probit <- 0
        ### Prior and data inputs ###
       Covs <- c(2:8) # Select which covariates/features to include
        standardize <- F # If TRUE, covariates/features
        #are standardized to mean 0 and variance 1
       lambda <- 2 # scaling factor for the prior of beta in our case tau = 2
        # Loading out data set
       wat<-read.table("WomenAtWork.dat",header = T) # read data from file</pre>
       Nobs <- dim(wat)[1] # number of observations
       y \leftarrow wat[1] \# y=1 \ if \ the \ women \ is \ working, \ otherwise \ y=0.
        X <- as.matrix(wat[,Covs]) # Covs matrix 7*7</pre>
       Xnames <- colnames(X)</pre>
        # Standraizing the covs matrix
        if (standardize){
          Index <- 2:(length(Covs)-1)</pre>
          X[,Index] <- scale(X[,Index])</pre>
       Npar \leftarrow dim(X)[2]
        # Setting up the prior
       mu <- as.matrix(rep(0,Npar)) # Prior mean vector</pre>
        Sigma <- (lambda)^2 *diag(Npar) # Prior covariance matrix</pre>
        LogPostLogistic <- function(betas,y,X,mu,Sigma){</pre>
          linPred <- X%*%betas;</pre>
          logLik <- sum( linPred*y - log(1 + exp(linPred)) );</pre>
           if (abs(logLik) == Inf){
                        logLik = -20000
              }# Likelihood is not finite, stear the optimizer away from here!
          logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
          return(logLik + logPrior)
        # Not in use we change the value to 0 at the beginning of the code
        LogPostProbit <- function(betas,y,X,mu,Sigma){</pre>
          linPred <- X%*%betas;</pre>
          SmallVal <- .Machine$double.xmin</pre>
          logLik <- sum(y*log(pnorm(linPred)+SmallVal) +</pre>
                          (1-y)*log(1-pnorm(linPred)+SmallVal))
          logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
          return(logLik + logPrior)
        # Select the initial values for beta
        initVal <- matrix(0,Npar,1)</pre>
```

```
if (Probit==1){
          logPost = LogPostProbit;
        } else{
          logPost = LogPostLogistic;
        # The argument control is a list of options to the optimizer optim,
        \#where\ fnscale = -1\ means\ that\ we\ minimize
        # the negative log posterior. Hence, we maximize the log posterior.
        OptimRes <- optim(initVal,logPost,gr=NULL,y,X,mu,Sigma,method=c("BFGS")
                            ,control=list(fnscale=-1),hessian=TRUE)
        postmode<-as.matrix(OptimRes$par[,1])</pre>
        poststd<- solve(-OptimRes$hessian)</pre>
        x_new<-as.matrix(x_new,ncol=1)</pre>
        betas<-rmvnorm(n=ndraws,mean = postmode, sigma = poststd)</pre>
        # Finding the value y givan the new Xs
        pr_y<-data.frame(x=betas%*%x_new)</pre>
        # Finding the probabilities using the logistics function
        pr_y$x_logit<-1-(1/(1+exp(-pr_y$x)))
        # Ploting the dataset
        plt <- ggplot(pr_y,aes(x = x_logit)) +geom_histogram(aes(y=..density..),</pre>
                   linetype=1, fill='#14213D')+
                   geom_density(alpha=.2,color="#FCA311",size=1,fill="#FCA311")+
                   labs(x='Pr(y=0|x)',y='',)
        plt
  }
pred_prob(10000,c(1,18,11,7,40,1,1))
  pred_prob2<- function(ndraws,x_new){</pre>
        ### Select Logistic or Probit regression and install packages ###
        Probit <- 0
        ### Prior and data inputs ###
        Covs <- c(2:8) # Select which covariates/features to include
        standardize <- F # If TRUE, covariates/features are
                           #standardized to mean O and variance 1
        lambda <- 2 # scaling factor for the prior of beta in our case tau = 2
        # Loading out data set
        wat <- read.table ("Women At Work.dat", header = T) # read data from file
        Nobs <- dim(wat)[1] # number of observations
        y \leftarrow wat[1] \# y=1 \ if \ the \ women \ is \ working, \ otherwise \ y=0.
        X <- as.matrix(wat[,Covs]) # Covs matrix 7*7</pre>
        Xnames <- colnames(X)</pre>
        # Standraizing the covs matrix
        if (standardize){
          Index <- 2:(length(Covs)-1)</pre>
          X[,Index] <- scale(X[,Index])</pre>
        }
        Npar \leftarrow dim(X)[2]
        # Setting up the prior
        mu <- as.matrix(rep(0,Npar)) # Prior mean vector</pre>
        Sigma <- (lambda)^2 *diag(Npar) # Prior covariance matrix
        LogPostLogistic <- function(betas,y,X,mu,Sigma){</pre>
          linPred <- X%*%betas;</pre>
```

```
logLik <- sum( linPred*y - log(1 + exp(linPred)) );</pre>
      if (abs(logLik) == Inf){
                   logLik = -20000
      # Likelihood is not finite, stear the optimizer away from here!
     logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
     return(logLik + logPrior)
    # Not in use we change the value to 0 at the beginning of the code
    LogPostProbit <- function(betas,y,X,mu,Sigma){</pre>
     linPred <- X%*%betas:</pre>
     SmallVal <- .Machine$double.xmin</pre>
     logLik <- sum(y*log(pnorm(linPred)+SmallVal) +</pre>
                     (1-y)*log(1-pnorm(linPred)+SmallVal))
     logPrior <- dmvnorm(betas, mu, Sigma, log=TRUE);</pre>
     return(logLik + logPrior)
    # Select the initial values for beta
   initVal <- matrix(0,Npar,1)</pre>
   if (Probit==1){
     logPost = LogPostProbit;
   } else{
     logPost = LogPostLogistic;
    # The argument control is a list of options to the optimizer optim,
   #where fnscale=-1 means that we minimize
    # the negative log posterior. Hence, we maximize the log posterior.
   OptimRes <- optim(initVal,logPost,gr=NULL,y,X,mu,Sigma,method=c("BFGS")
                     ,control=list(fnscale=-1),hessian=TRUE)
   postmode<-as.matrix(OptimRes$par[,1])</pre>
   poststd<- solve(-OptimRes$hessian)</pre>
   x_new<-as.matrix(x_new,ncol=1)</pre>
   betas<-rmvnorm(n=ndraws,mean = postmode, sigma = poststd)</pre>
    # Finding the value y givan the new Xs
   pr_y<-data.frame(x=betas%*%x_new)</pre>
    # Finding the probabilities using the logistics function
   pr_y$x_logit<-1/(1+exp(-pr_y$x))
   #Adding the clasifier
   pr_y$job_flag <- ifelse(pr_y$x_logit <= 0.5, 0, 1)</pre>
   plt <- ggplot(pr_y, aes(x = x_logit, y = job_flag)) +</pre>
     geom_point(colour="#14213D") +
     # stat_smooth(method="glm",colour="#FCA311",
                   alpha = 0.5, se=FALSE, fullrange=TRUE,
                   method.arqs = list(family=binomial)) +
     xlab("Predictor") + xlim(c(0,1))+
     ylab("Probability of Outcome") +
      ggtitle("Logistic Regression function with 0.5 as decision boundary")+
geom_vline(aes(xintercept = 0.5), color = "#14213D", size=1, alpha = 0.1) +
geom_hline(aes(yintercept = 0.5), color = "#14213D", size=1, alpha = 0.1)
```