# SC1\_Proj

### Alessio

## 13 January 2020

## Introduction

One of the most common types of cancer diagnosed in women is breast cancer. There are multiple tests that people are subjected to, but one of the most indicative ones is fine needle aspiration which involes extracting a sample of cells to be examined under a microscope. Multiple numerical metrics are computed from the obtained images. The aim is to use the extracted metrics to make accurate diagnoses.

The dataset consists of 569 images which have been processed as described and a total of 30 variables have been computed for each observation.

The aim of this report is to implement a number of classification algorithms, use them to obtain predictions, and compare their performances.

## Exploratory analysis

```
data <- read_csv("../data/data.csv")
colnames(data)[3:32] <- c(
    "radius_m", "texture_m", "perim_m", "area_m", "smooth_m", "compact_m",
    "concav_m", "concav_pt_m", "symmetry_m", "frac_dim_m", "radius_se",
    "texture_se", "perim_se", "area_se", "smooth_se", "compact_se",
    "concav_se", "concav_pt_se", "symmetry_se", "frac_dim_se", "radius_w",
    "texture_w", "perim_w", "area_w", "smooth_w", "compact_w", "concav_w",
    "concav_pt_w", "symmetry_w", "frac_dim_w"
)</pre>
```

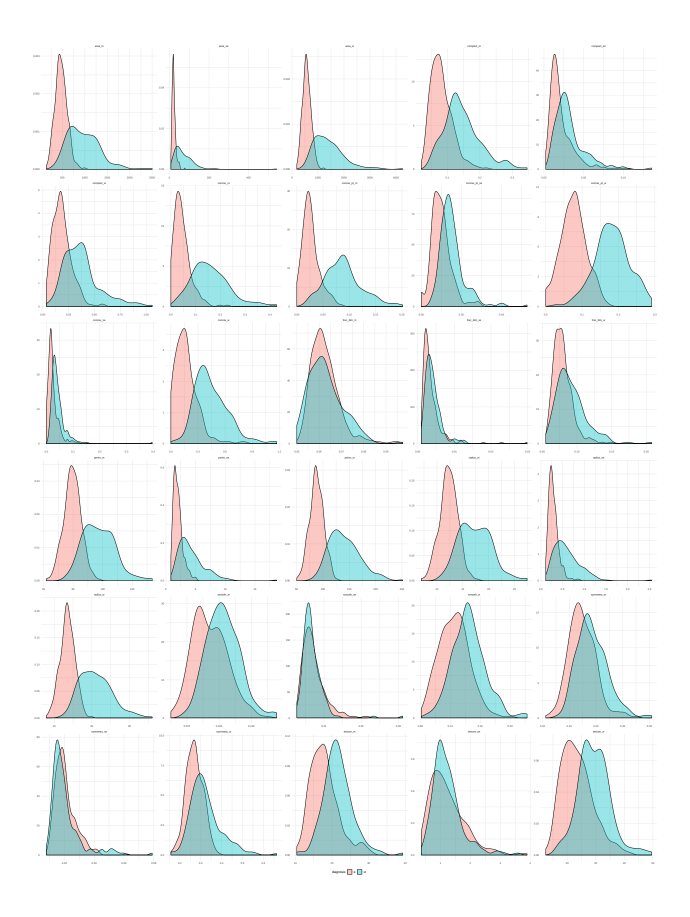
Check for missing values in every column.

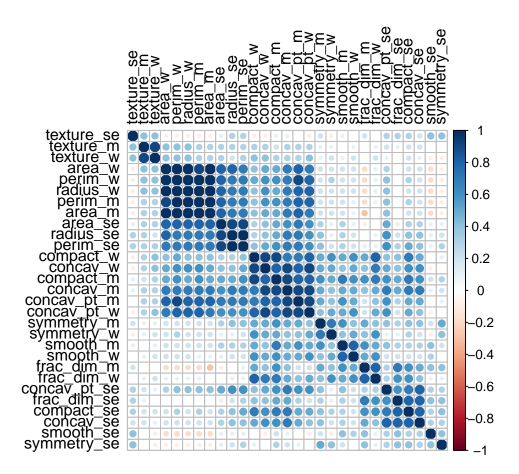
```
colSums(is.na(data))
```

```
##
              id
                     diagnosis
                                    radius_m
                                                  texture_m
                                                                   perim_m
                                                                                  area_m
##
                                            0
               0
                                                           0
                                                                                        0
##
       smooth m
                     compact m
                                     concav_m
                                                concav_pt_m
                                                               symmetry_m
                                                                              frac dim m
##
               0
                                            0
                                                           0
##
      radius_se
                    texture_se
                                    perim_se
                                                                smooth se
                                                                              compact_se
                                                    area_se
##
               0
##
                                               frac_dim_se
                                                                               texture_w
      concav_se concav_pt_se
                                 symmetry_se
                                                                 radius_w
##
               0
                              0
                                            0
##
        perim_w
                        area_w
                                     smooth_w
                                                  compact_w
                                                                  concav_w
                                                                             concav_pt_w
##
               0
                              0
                                            0
                                                                         0
##
                    frac_dim_w
                                          X33
     symmetry_w
```

data %<>% mutate\_at(vars(diagnosis), factor)

```
train <- data %>% sample_frac(0.8)
test <- anti_join(data, train, by = "id")</pre>
# need ids for later
id_train <- train$id</pre>
id_test <- test$id</pre>
data %<>%
  dplyr::select(-c(id, X33))
train %<>%
  dplyr::select(-c(id, X33))
test %<>%
  dplyr::select(-c(id, X33))
sum(is.na(data))
## [1] 0
training_data <- train[2:dim(train)[2]]</pre>
training_classes <- train[1]</pre>
test_data <- test[2:dim(test)[2]]</pre>
test_classes <- test[1]</pre>
0.6
                                                  0.6
0.4 -
                                                  0.4
0.2
                                                  0.2
                                                  0.0
0.0
               В
                                  M
                                                                 В
                                                                                     M
```



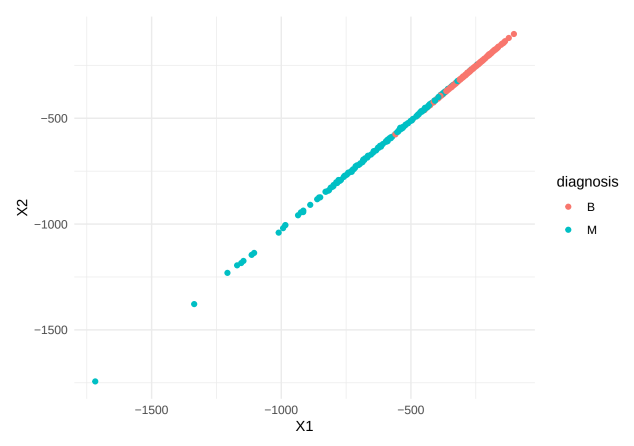


## Dimensionality Reduction and Feature Selection

## PCA

Code

```
normalise_z <- function(X) {</pre>
  mean_cols <- colMeans(X)</pre>
  sd_cols <- apply(X, 2, sd)</pre>
  mean_normalised_X <- t(apply(X, 1, function(x) {</pre>
    x - mean_cols
  }))
  normalised_X <- t(apply(mean_normalised_X, 1, function(x) {</pre>
    x / sd_cols
  }))
  return(normalised_X)
}
pca <- function(X, number_components_keep) {</pre>
  normalised_X <- normalise_z(X)</pre>
  corr_mat <- t(normalised_X) %*% normalised_X</pre>
  eigenvectors <- eigen(corr_mat, symmetric = TRUE)$vectors</pre>
  reduced_data <- X %*% eigenvectors[, 1:number_components_keep]</pre>
  relevant_eigs <- eigenvectors[, 1:number_components_keep]</pre>
  returnds <- list(reduced_data, relevant_eigs)</pre>
  names(returnds) <- c("reduced_data", "reduction_matrix")</pre>
  return(returnds)
Apply to dataset
pca_result <- pca(as.matrix(training_data), 2)</pre>
pca_reduced_training_data <- data.frame(cbind(pca_result$reduced_data, training_classes))</pre>
ggplot(data = pca_reduced_training_data, aes(x = X1, y = X2)) +
      geom_point(aes(colour = diagnosis))
```



- Correlation Feature SelectionLDA

## Classification

add list of methods+measure

### SVM

From the application of PCA to the dataset we can see that, after reducing to 2 dimensions, the data appears to be almost linearly separable. Given this, an appropriate method of classifying the data would be to apply a soft-margin SVM to the reduced dimension data. Soft-margin SVMs solve the problem of classifying non-separable data by permitting a certain number of points to be incorrectly classifed however the number and the amount they violate the constraints by must be as small as possible. After manipulating the reformulated optimisation problem we end up with the optimisation problem

$$\min_{\lambda} \frac{\overline{\lambda}XX^T\overline{\lambda^T}}{4} + \lambda^T \mathbf{1}$$
such that  $0 \le \lambda_i \le C$ 
and  $\sum_{i=1}^{n} \lambda_i y_i = 0$ 

where

$$X = \begin{pmatrix} x_1^T \\ \vdots \\ x_n^T \end{pmatrix} \text{ and } \overline{\lambda} = [\lambda_1 \cdot y_i, ..., \lambda_n \cdot y_n] \text{ and } \mathbf{1} = [1, ..., 1] \in \mathbb{R}^{\times}$$

For some predefined C. This is a quadratic programming problem with linear constraints which can be solved using the R package quadprogwith the function solve.QP. From its documentation, this function can solve (for b) problems in the form  $\min_b(-d^Tb + \frac{1}{2}b^TDb)$  with the constraints that  $A^Tb \ge b_0$ . By transforming the above problem into this format we can implement soft-margin SVM using the following code

```
train_soft_svm <- function(X, y, C) {
  num_observation <- nrow(X)
  dim_num <- ncol(X)

Dmat2 <- diag(y) * X %*% t(X) %*% diag(y)
  diag(Dmat2) <- diag(Dmat2) + 1e-6
  dv2 <- rep(1, num_observation)

A2 <- rbind( y, diag(num_observation))
  A2 <- rbind(A2, -1*diag(num_observation))

bv2 <- c(c(0), rep(0, num_observation), rep(-C, num_observation))

model <- solve.QP(Dmat2, dv2, t(A2), bv2, meq = 1)
}</pre>
```

In order to recover w and b from  $\lambda$  we use the relationship

$$w = \sum_{i=0}^{n-1} \lambda_i x_i^T y_i$$

and

$$b = \operatorname{mean}(\sum_{i=0}^{k} y_i - w \cdot x_i) \cdot \forall i \cdot 0 < \lambda_i < C$$

Which can be made as functions in R as so:

```
calculate_b <- function(w, X, y, a, C) {
    ks <- sapply(a, function(x) {
        return(x > 0 && x < C)
    })
    indices <- which(ks)
    sum_bs <- 0
    for (i in indices) {
        sum_bs <- sum_bs + (y[i] - w %*% X[i, ])
    }
    return(sum_bs / length(indices))
}

recover_w <- function(a, y, X) {
    colSums(diag(a) %*% diag(y) %*% X)
}</pre>
```

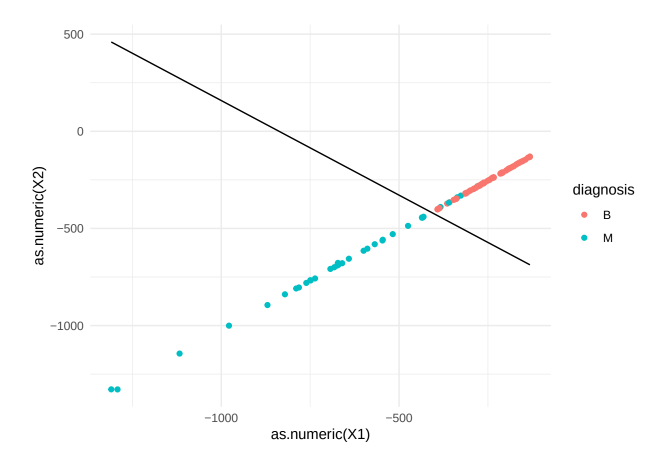
From the parameters we can recover the equation of the line corresopnding the decision boundary which can later be used for plotting

```
soft_margin_svm_plotter <- function(w, b) {
  plotter <- function(x) {
    return(1 / w[2] * -(b + (w[1] * as.numeric(x))))
  }
  return(plotter)
}</pre>
```

#### TODO: prediction function

We are now able to define a function which first embeds the vector in the reduced space and then predicts the class based upon the prediction function produced by the parameters found by the SVM. Below is code that will do the following based upon a package written for this assignment found here

When running the trained SVM prediction functio on the test set we achieve 94.7368421 %. We can plot this code as below



## **Naive Bayes**

## Mathematical setting

Let y be the class label that we want to assign to an observation  $\mathbf{x} = (x_1, \dots x_d)$ , where  $x_1, \dots x_d$  are the features. The probability of an observation having label y is given by Bayes rule,

$$P(y|x_1,\dots,x_d) = \frac{P(x_1,\dots,x_d|y_k)P(y)}{P(x_1,\dots,x_d)}$$
$$\propto P(x_1,\dots,x_d|y_k)P(y).$$

The prior class probability P(y) can be easily obtained by the proportion of observation that are in the given class.

The main assumption is that every feature is conditionally independent given the class label y. The reason why this classifier is called naive is that very often this assumption is not actually realistic.

This assumption simplifies the posterior to

$$P(y|x_1,\dots,x_d) \propto P(y) \prod_{i=1}^d P(x_i|y).$$

There are various types of Naive Bayes classifiers based on the type of features. In our case, since we have continuous variables we assume that all features are normally distributed. Therefore, the conditional probabilities can be calculated as

$$P(x_i|y) = \frac{1}{\sqrt{2\pi\sigma_y^2}} exp\left(-\frac{(x_i - \mu_y)^2}{2\sigma_y^2}\right)$$

Finally, to assign the class to an observation we use the Maximum A Posteriori decision rule. For every observation, we pick the class the has the highest probability

$$y = \underset{y}{\operatorname{argmax}} P(y) \prod_{i=1}^{d} P(x_i|y).$$

## Implementation

Here are some code snippets just to illustrate how these theoretical aspects are implemented. The full code can be found in the package.

The observations are stored as rows in X and the corresponding class labels are entires in the column matrix y.

First we calculate the prior class probabilities based on the number of observations in each class.

```
n <- dim(X)[1]
d <- dim(X)[2]
classes <- sort(unique(y)[, 1])
k <- length(classes)

prior <- rep(0, k)
for (i in 1:k) {
   prior[i] <- sum(y == classes[i]) / n
}</pre>
```

Then we create an array of the mean and sd of the data split by classes and features.

```
summaries <- array(rep(1, d * k * 2), dim = c(k, d, 2))
for (i in 1:k) {
    X_k <- X[which(y == (i - 1)), ]
    summaries[i, , 1] <- apply(X_k, 2, mean)
    summaries[i, , 2] <- apply(X_k, 2, sd)
}</pre>
```

Finally, the predictions are obtained by taking the largest posterior class probability. Note that in order to avoid underflow, we take the maximum of the log posterior class probabilities.

```
probs <- matrix(rep(0, n * k), nrow = n)
for (obs in 1:n) {
    for (class in 1:k) {
        class_prob <- log(prior[class])
        for (feat in 1:d) {
            mu <- summaries[class, feat, 1]
            sd <- summaries[class, feat, 2]
            cond <- dnorm(x_new[obs, feat], mu, sd, log = TRUE)
            class_prob <- class_prob + cond
        }
        probs[obs, class] <- class_prob
    }
}
pred <- apply(probs, 1, which.max)</pre>
```

### Fit model to dataset

```
levels(training_classes$diagnosis) <- c(0, 1)
training_classes %<>% as.matrix
mode(training_classes) <- "numeric"

levels(test_classes$diagnosis) <- c(0, 1)
test_classes %<>% as.matrix
mode(test_classes) <- "numeric"</pre>
```

Fit the Naive Bayes model to the data, calculate predictions and check the accuracy using.

```
model_naive <- naive_bayes(training_data, training_classes)
pred_naive <- predict(model_naive, as.matrix(test_data))
# confusion_plot(test_classes,pred_naive)
calc_accuracy <- function(ytest, yhat) sum(drop(yhat) == drop(ytest)) / length(drop(ytest))
acc_naive <- calc_accuracy(test_classes, pred_naive)
acc_naive</pre>
```

```
## [1] 0.9385965
```

Fit the model to the PCA-reduced dataset.

```
model_naive_pca <- naive_bayes(pca_result$reduced_data, training_classes)</pre>
```

```
pred_naive_pca <- predict(model_naive_pca, pca_reduced_test_data)
acc_naive_pca <- calc_accuracy(test_classes, pred_naive_pca)
acc_naive</pre>
```

## [1] 0.9385965

## Logistic Regression

## **Mathematical Setting**

Let  $Y_i \mid \mathbf{x}_i \sim \text{Bernoulli}(p_i)$  with  $p_i = \sigma(\mathbf{x}_i^{\top} \boldsymbol{\beta})$  where  $\sigma(\cdot)$  is the **sigmoid function**. The joint log-likelihood is given by

$$\ln p(\mathbf{y} \mid \boldsymbol{\beta}) = \sum_{i=1}^{n} y_i \ln(p_i) + (1 - y_i) \ln(1 - p_i) = -\sum_{i=1}^{n} \ln \left( 1 + \exp((1 - 2y_i) \mathbf{x}_i^{\top} \boldsymbol{\beta}) \right)$$

### **Maximum Likelihood Estimation**

Maximizing the likelihood is equivalent to minimizing the negative log-likelihood. Minimizing the negative log likelihood is equivalent to solving the following optimization problem

$$\min_{\beta} \sum_{i=1}^{n} \ln \left( 1 + \exp((1 - 2y_i) \mathbf{x}_i^{\top} \boldsymbol{\beta}) \right)$$

## Maximum-A-Posteriori and Ridge Regularization

We can introduce an isotropic Gaussian prior on **all** the coefficients  $p(\beta) = N(\mathbf{0}, \sigma_{\beta}^2 I)$ . Maximizing the posterior  $p(\beta \mid \mathbf{y})$  is equivalent to minimizing the negative log posterior  $-\ln p(\beta \mid \mathbf{y})$  giving

$$\min_{\boldsymbol{\beta}} \sigma_{\boldsymbol{\beta}}^2 \sum_{i=1}^n \ln \left( 1 + \exp((1-2y_i)\mathbf{x}_i^\top \boldsymbol{\beta}) \right) + \frac{1}{2} \boldsymbol{\beta}^\top \boldsymbol{\beta}$$

#### Gradient Ascent for Maximum Likelihood Estimation

Updates take the form

$$\boldsymbol{\beta}_{k+1} \leftarrow \boldsymbol{\beta}_k + \gamma \boldsymbol{X}^\top (\mathbf{y} - \boldsymbol{\sigma}(\boldsymbol{X} \boldsymbol{\beta}_k))$$

#### Gradient Ascent for Maximum-A-Posteriori

The update takes the form

$$\boldsymbol{\beta}_{k+1} \leftarrow \boldsymbol{\beta}_k + \gamma_k \left[ \boldsymbol{\sigma}_{\boldsymbol{\beta}}^2 \boldsymbol{X}^\top (\mathbf{y} - \boldsymbol{\sigma}(\boldsymbol{X} \boldsymbol{\beta}_k)) - \boldsymbol{\beta}_k \right]$$

### Newton's Method for Maximum Likelihood Estimation

The iterations are as follows, where for stability one can add a learning rate  $\alpha$ , which is in practice often set to  $\alpha = 0.1$ .

$$\boldsymbol{\beta}_{k+1} \leftarrow \boldsymbol{\beta}_k + \alpha (\boldsymbol{X}^\top \boldsymbol{D} \boldsymbol{X})^{-1} \boldsymbol{X}^\top (\mathbf{y} - \sigma(\boldsymbol{X} \boldsymbol{\beta}_k))$$

In practice we would solve the corresponding system for d

$$(X^{\top}DX)\mathbf{d}_k = \alpha X^{\top}(\mathbf{y} - \sigma(X\boldsymbol{\beta}_k))$$

and then perform the update

$$\beta_{k+1} \leftarrow \beta_k + \mathbf{d}_k$$

### Newton's Method for Maximum-A-Posteriori

The update takes the form

$$\boldsymbol{\beta}_{k+1} \leftarrow \boldsymbol{\beta}_k + \alpha \left[ \boldsymbol{\sigma}_{\boldsymbol{\beta}}^2 \boldsymbol{X}^\top \boldsymbol{D} \boldsymbol{X} + \boldsymbol{I} \right]^{-1} \left( \boldsymbol{\sigma}_{\boldsymbol{\beta}}^2 \boldsymbol{X}^\top (\mathbf{y} - \boldsymbol{\sigma}(\boldsymbol{X} \boldsymbol{\beta}_k)) - \boldsymbol{\beta}_k \right)$$

## **Implementation**

First, we need to add a column of 1s to the design matrix so that we can fit the bias coefficient.

```
Xtrain <- as.matrix(cbind(1, training_data))
Xtest <- as.matrix(cbind(1, test_data))</pre>
```

#### Maximum a Posteriori

We implement BFGS, Newton's Method and Gradient Ascent.

### **Maximum Likelihood Estimation**

Similarly, we also implement the same algorithms for Maximum Likelihood Estimation.

### Comparing Cost Functions and Optimization Methods

Below we can see the accuracy of the different optimization methods (BFGS, Newton's method and Gradient Ascent) on the different cost functions (MAP or MLE).

```
# put everything together into a nice table
results_matrix <- matrix(c(
   acc_mle_bfgs, acc_mle_nm, acc_mle_ga,
   acc_map_bfgs, acc_map_nm, acc_map_ga
),</pre>
```

```
dimnames = list(c("BFGS", "NM", "GA"), c("MLE", "MAP")),
nrow = 3, ncol = 2
)
results <- data.frame(results_matrix)
kable(results)</pre>
```

	MLE	MAP
BFGS	0.9824561	0.9473684
NM	0.9473684	0.9473684
GA	0.9298246	0.4035088

#### Random-Walk Metropolis-Hastings Implementation on Reduced Data

Next, we can define a function that performs Random Walk Metropolis Hastings with a normal proposal distribution. To make it more efficient we can work with the log posterior and change the decision rule. In addition, we can pre-calculate all the normal and uniform samples and just access them later.

```
rwmh_multivariate_log <- function(start, niter, logtarget, vcov, thinning, burnin){</pre>
    # Set current z to the initial point and calculate its log target to save computations
                   # It's a column vector
    z <- start
    pz <- logtarget(start)</pre>
    # create vector deciding iterations where we record the samples
    store <- seq(from=(1+burnin), to=niter, by=thinning)</pre>
    #n_samples <- (niter - burnin) %/% thinning</pre>
    # Generate matrix containing samples. Initialize with the starting value
    samples <- matrix(0, nrow=length(store), ncol=nrow(start))</pre>
    samples[1, ] <- start</pre>
    # Generate uniform random numbers in advance, to save computation. Log them.
    log u <- log(runif(niter))</pre>
    # Proposal is a multivariate standard normal distribution. Generate samples and
    # later on use linearity property of Gaussian distribution
    vcov <- diag(nrow(start)) %*% vcov</pre>
    normal_shift <- mvrnorm(n=niter, mu=c(0,0,0), Sigma=vcov)</pre>
    for (i in 2:niter){
        # Sample a candidate
        candidate <- z + normal shift[i, ]</pre>
        # calculate log target of candidate and store it in case it gets accepted
        p_candidate <- logtarget(candidate)</pre>
        # use decision rule explained in blog posts
        if (log_u[i] <= p_candidate - pz){</pre>
            # Accept!
            z <- candidate
            pz <- p_candidate
        # Finally add the sample to our matrix of samples
        if (i %in% store) samples[which(store==i), ] <- z</pre>
    }
    # Finally add the sample to our matrix of samples
    if (i %in% store) samples[which(store == i), ] <- z</pre>
  return(samples)
```

Now we can add a column of 1s to the reduced dataset created by PCA so that we can run Logistic Regression

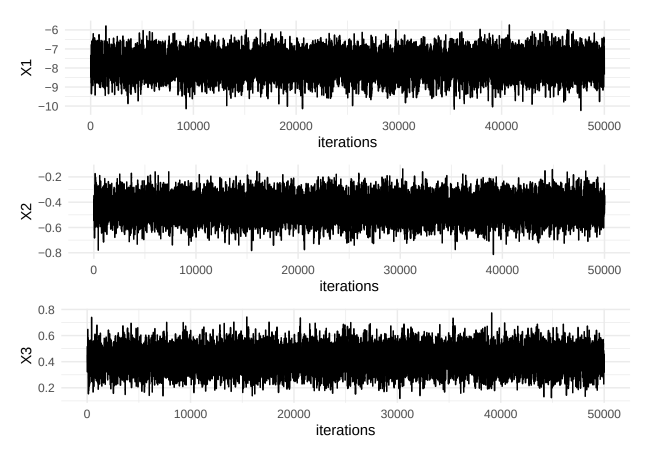
on the reduced data.

```
Xrwmh <- cbind(1, as.matrix(pca_reduced_training_data[, c(1, 2)]))</pre>
```

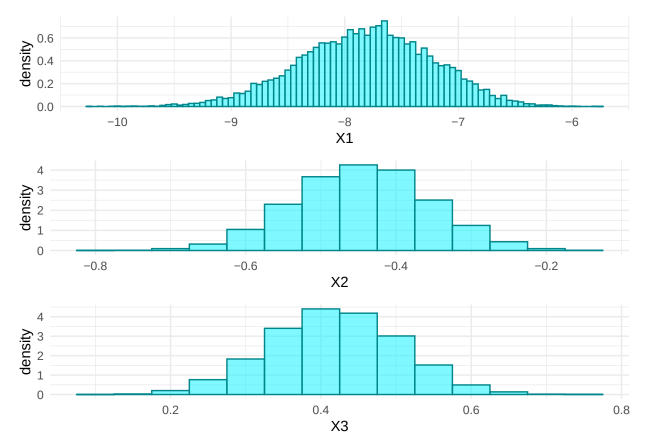
We run logistic regression using the BFGS optimization method on MAP. Since we need the Hessian matrix for sampling later on, we simply use the optim function on the unnormalized version of the posterior distribution for logistic regression.

We're now ready to start sampling starting at the MAP estimate and using, as a proposal, the normal distribution with variance-covariance matrix given by the inverse of the approximated hessian matrix coming from the optimization routine.

We can look at the trace plots and notice how the chain is mixing quickly, signalling a good exploratory behavior.

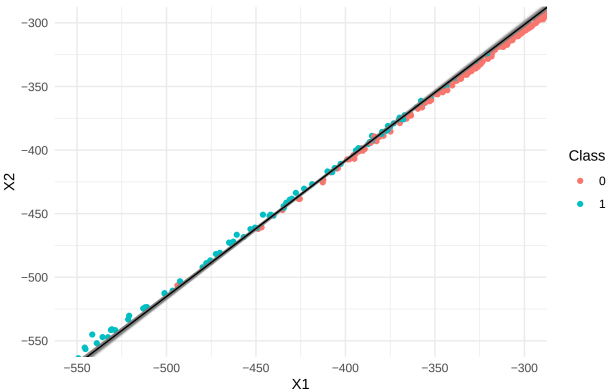


Plotting the histograms of the samples for each coordinate of the parameter vector shows that RWMH is indeed sampling correctly.



The next thing to do is to plot the decision boundary found by BFGS together with the sampled lines.





Finally, we can look at the performance of logistic regression on this reduced dataset. As a decision rule we just use a threshold.

```
Xtest_embedded <- cbind(1, as.matrix(embedded_test_data[, c(1, 2)]))
lr_embedded_preds <- round(1.0 / (1.0 + exp(-Xtest_embedded %*% start)))
calc_accuracy(test_classes, lr_embedded_preds)</pre>
```

## [1] 0.9649123

## Conclusion

Merge all predictions

```
id <- seq(length(id_test))</pre>
all_pred <- cbind(id, numeric_test_labels, pred_naive, pred_svm)
colnames(all_pred) <- c("id", "actual", "naive", "svm")</pre>
all_pred[all_pred == -1] <- 0
all_pred %<>% as.data.frame()
confusion_plot <- function(actual, predicted) {</pre>
  confusion_matrix <- as.data.frame(table(actual, predicted))</pre>
  g <- ggplot(confusion_matrix, aes(x = actual, y = predicted)) +</pre>
    geom_tile(aes(fill = Freq)) +
    geom_text(aes(label = sprintf("%1.0f", Freq)), color = "white", fontface = "bold") +
    labs(x = "Actual class", y = "Predicted class") +
    theme_minimal()
  return(g)
}
errors <- all_pred %>%
  mutate(
   naive = naive == actual,
    svm = svm == actual,
   logistic = yhat_map_bfgs == actual
  ) %>%
  dplyr::select(-"actual") %>%
  melt(id = "id")
```

Visualize which are the observations that the models missclassify.

```
ggplot(errors, aes(x = id, y = variable, fill = value)) +
  geom_raster() +
  theme_minimal()
```

