Assignment_3

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#Assignment 3

##Part 1 - Simulating data

Use meta analysis reported in Parola et al (2020) to create informed simulated data - 100 pairs of schizophrenia and controls, each participant producing 10 repeated measures (10 trials with their speech recorded), for each recording produce 10 acoustic measures (6 from meta analysis and 4 with random noise)

 Do the same for a baseline data set including only 10 noise variables ##Sara ### seeting up variables

```
###Data simulation
n <- 100
trials <- 10

#Effect sizes definition = Informed effect mean and Skeptic effect mean
IEM <- c(-0.5,-1.26,-.74,1.89,0.25,1.3,0,0,0,0)
SEM <- rep(0,10)

#Defining individual vairability from populationd accross trials measurement error
ISD <- 1
TSD <- 0.5
E <- 0.2</pre>
```

Simulating the true effect size for each varibale for all pairs of participants

```
for (i in seq(10)){
  temp_informed <- tibble(</pre>
    ID=seq(n),
    TrueEffect = rnorm(n, IEM[i], ISD),
    Variable = paste0("V",i))
  temp_skeptic <- tibble(</pre>
    ID=seq(n),
    TrueEffect = rnorm(n, SEM[i], ISD),
    Variable = paste0("V",i))
  if(i==1){
    d_informed_true <- temp_informed</pre>
    d_skeptic_true <- temp_skeptic</pre>
  } else {
    d informed true <- rbind(d informed true, temp informed)
    d_skeptic_true <- rbind(d_skeptic_true, temp_skeptic)</pre>
  }
}
```

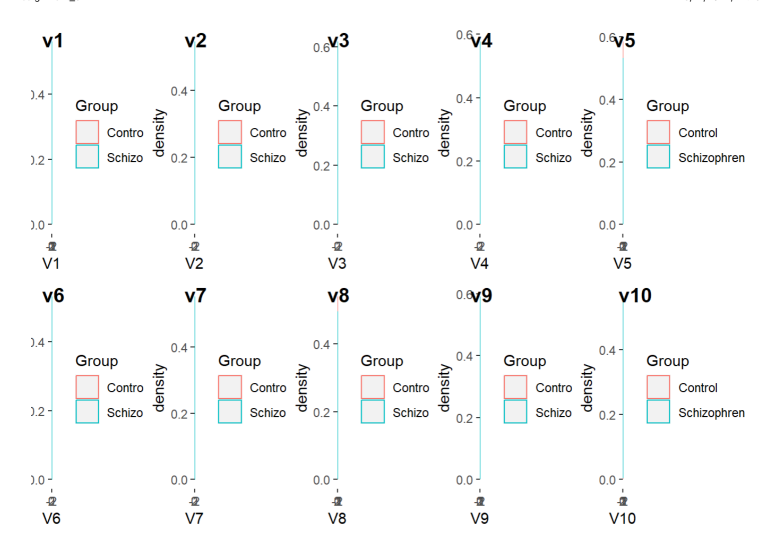
Creating one row per trial

Transforming the dataframe to a wide format based on the variable

```
d_informed_wide <- d_informed %>%
  mutate(TrueEffect=NULL) %>%
  pivot_wider(names_from = Variable,
              values_from = measurement)
d_skeptic_wide <- d_skeptic %>%
  mutate(TrueEffect=NULL) %>%
  pivot_wider(names_from = Variable,
              values from = measurement)
Schizo_ID <- d_informed_wide %>%
  filter(Group == 'Schizophrenia')
 control_ID <- d_informed_wide %>%
  filter(Group == 'Control')
control_ID[, 1] <- control_ID[,1] + 100</pre>
 d_informed_wide <- rbind(control_ID,Schizo_ID)</pre>
 Schizo ID skeptic <- d skeptic wide %>%
  filter(Group == 'Schizophrenia')
 control_ID_skeptic <- d_skeptic_wide %>%
  filter(Group == 'Control')
 control_ID_skeptic[,1] <- control_ID_skeptic[,1] + 100</pre>
 d_skeptic_wide <- rbind(control_ID_skeptic,Schizo_ID_skeptic)</pre>
```

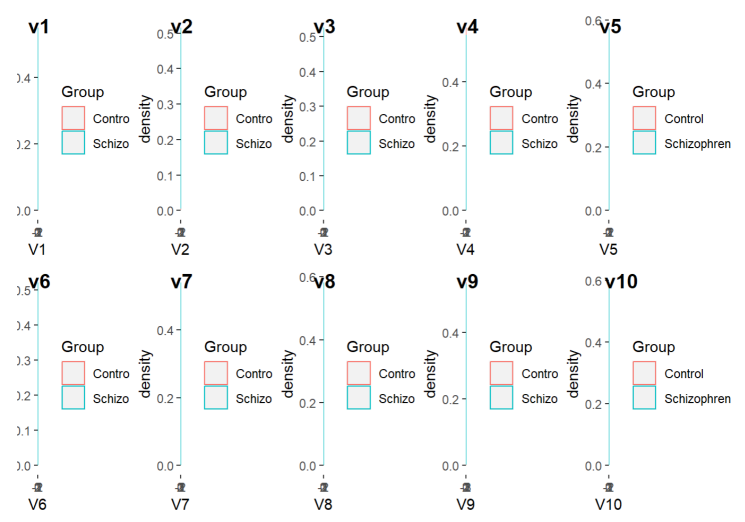
Visualizing the simulated informed data

```
plot1 <- d_informed_wide %>%
  ggplot(aes(x = V1, color = Group))+
  geom_density()
plot2 <- d informed wide %>%
  ggplot(aes(x = V2, color = Group))+
  geom density()
plot3 <- d_informed_wide %>%
  ggplot(aes(x = V3, color = Group))+
  geom_density()
plot4 <- d informed wide %>%
  ggplot(aes(x = V4, color = Group))+
  geom density()
plot5 <- d informed wide %>%
  ggplot(aes(x = V5, color = Group))+
  geom_density()
plot6 <- d informed wide %>%
  ggplot(aes(x = V6, color = Group))+
  geom_density()
plot7 <- d_informed_wide %>%
  ggplot(aes(x = V7, color = Group))+
  geom_density()
plot8 <- d informed wide %>%
  ggplot(aes(x = V8, color = Group))+
  geom density()
plot9 <- d_informed_wide %>%
  ggplot(aes(x = V9, color = Group))+
  geom_density()
plot10 <- d informed wide %>%
  ggplot(aes(x = V10, color = Group))+
  geom_density()
cowplot::plot_grid(plot1, plot2, plot3, plot4, plot5, plot6, plot7, plot8, plot9,
plot10,
          labels = c("v1", "v2", "v3", 'v4', 'v5', 'v6', 'v7', 'v8', 'v9', 'v10'),
          ncol = 5, nrow = 2)
```



Visualizing the simulated data for skeptical data frame

```
plot1_s <- d_skeptic_wide %>%
  ggplot(aes(x = V1, color = Group))+
  geom_density()
plot2 s <- d skeptic wide %>%
  ggplot(aes(x = V2, color = Group))+
  geom density()
plot3_s <- d_skeptic_wide %>%
  ggplot(aes(x = V3, color = Group))+
  geom_density()
plot4 s <- d skeptic wide %>%
  ggplot(aes(x = V4, color = Group))+
  geom density()
plot5_s <- d_skeptic_wide %>%
  ggplot(aes(x = V5, color = Group))+
  geom_density()
plot6 s <- d skeptic wide %>%
  ggplot(aes(x = V6, color = Group))+
  geom_density()
plot7_s <- d_skeptic_wide %>%
  ggplot(aes(x = V7, color = Group))+
  geom density()
plot8 s <- d skeptic wide %>%
  qqplot(aes(x = V8, color = Group))+
  geom density()
plot9_s <- d_skeptic_wide %>%
  ggplot(aes(x = V9, color = Group))+
  geom_density()
plot10 s <- d skeptic wide %>%
  ggplot(aes(x = V10, color = Group))+
  geom_density()
cowplot::plot_grid(plot1_s, plot2_s, plot3_s, plot4_s, plot5_s, plot6_s, plot7_s,
plot8_s, plot9_s, plot10_s,
          labels = c("v1", "v2", "v3", 'v4', 'v5', 'v6', 'v7', 'v8', 'v9', 'v10'),
          ncol = 5, nrow = 2)
```



##Patrik ##Part 2 - Machine larning pipeline on simulated data Build a machine leaning pipeline (separately on the 2 datasets) - create a data budget (e.g., balanced training and test sets) pree-process the data (e.g., scaling the features) - fit and assess a classification algorithm on the training data (e.g., bayesian multilevel logistic regression) - assess performance on the test set - discuss whether performance and feature importance is as expected

Data budget (splitting) and pree-processing (scaling)

```
set.seed(260)

d_informed_wide <- d_informed_wide %>%
   mutate(pair_ID=ID) %>%
   mutate(pair_ID=ifelse(ID>100, ID-100, ID))

d_skeptic_wide <- d_skeptic_wide %>%
   mutate(pair_ID=ID) %>%
   mutate(pair_ID=ifelse(ID>100, ID-100, ID))

split_inf <- initial_split(d_informed_wide, prop = 4/5)

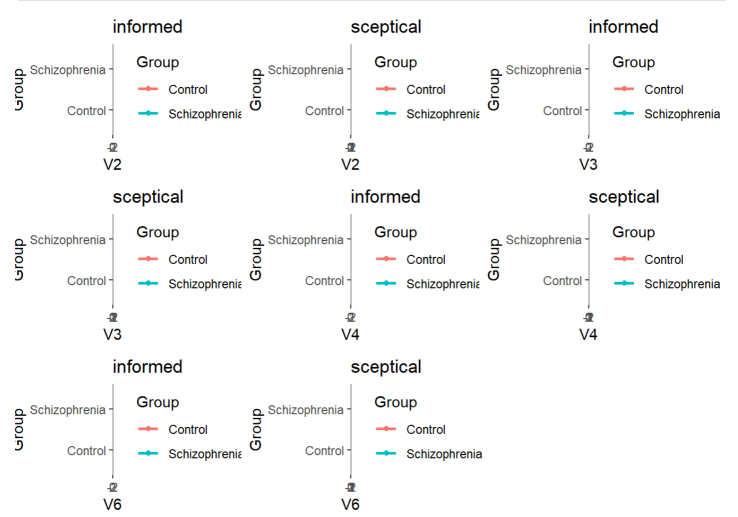
train_informed <- training(split_inf)
test_informed <- testing(split_inf)</pre>
```

```
split_skep <- initial_split(d_skeptic_wide, prop = 4/5)</pre>
train_skeptic <- training(split_skep)</pre>
test skeptic <- testing(split skep)</pre>
train_informed$ID <- as.factor(train_informed$ID)</pre>
train_skeptic$ID <- as.factor(train_skeptic$ID)</pre>
test informed$ID <- as.factor(test informed$ID)</pre>
test_skeptic$ID <- as.factor(test_skeptic$ID)</pre>
train_informed$pair_ID <- as.factor(train_informed$pair_ID)</pre>
train_skeptic$pair_ID <- as.factor(train_skeptic$pair_ID)</pre>
test informed$pair ID <- as.factor(test informed$pair ID)
test_skeptic$pair_ID <- as.factor(test_skeptic$pair_ID)</pre>
train informed$Trial <- as.factor(train informed$Trial)</pre>
train_skeptic$Trial <- as.factor(train_skeptic$Trial)</pre>
test informed$Trial <- as.factor(test informed$Trial)</pre>
test_skeptic$Trial <- as.factor(test_skeptic$Trial)</pre>
rec_informed <- train_informed %>%
  recipe(Group~.) %>%
  update role(ID, new role = 'ID') %>%
  step_scale(all_numeric()) %>%
  step_center(all_numeric()) %>%
  prep(training=train informed, retain=TRUE)
rec_skeptic <- train_skeptic %>%
  recipe(Group~.) %>%
  update_role(ID, new_role = 'ID') %>%
  step scale(all numeric()) %>%
  step center(all numeric()) %>%
  prep(training=train_informed, retain=TRUE)
train_informed_s <- juice(rec_informed)</pre>
test_informed_s <- bake(rec_informed, new_data = test_informed)</pre>
train_skeptic_s <- juice(rec_skeptic)</pre>
test_skeptic_s <- bake(rec_skeptic, new_data = test_skeptic)</pre>
```

Visual inspection of the data

```
plot_i2 <- ggplot(train_informed, aes(V2, Group, colour=Group))+</pre>
  geom point()+
  geom smooth(method = "glm", se=FALSE)+
  theme_bw()+
  ggtitle("informed")
plot s2 <- ggplot(train skeptic, aes(V2, Group, colour=Group))+</pre>
  geom point()+
  geom_smooth(method = "glm", se=FALSE)+
  theme_bw()+
  ggtitle("sceptical")
plot_i3 <- ggplot(train_informed, aes(V3, Group, colour=Group))+</pre>
  geom_point()+
  geom_smooth(method = "glm", se=FALSE)+
  theme bw()+
  ggtitle("informed")
plot_s3 <- ggplot(train_skeptic, aes(V3, Group, colour=Group))+</pre>
  geom point()+
  geom smooth(method = "glm", se=FALSE)+
  theme bw()+
  ggtitle("sceptical")
plot_i4 <- ggplot(train_informed, aes(V4, Group, colour=Group))+</pre>
  geom_point()+
  geom_smooth(method = "glm", se=FALSE)+
  theme_bw()+
  ggtitle("informed")
plot s4 <- ggplot(train skeptic, aes(V4, Group, colour=Group))+</pre>
  geom_point()+
  geom_smooth(method = "glm", se=FALSE)+
  theme bw()+
  ggtitle("sceptical")
plot_i6 <- ggplot(train_informed, aes(V6, Group, colour=Group))+</pre>
  geom point()+
  geom_smooth(method = "glm", se=FALSE)+
  theme bw()+
  ggtitle("informed")
plot s6 <- ggplot(train skeptic, aes(V6, Group, colour=Group))+</pre>
  geom point()+
  geom_smooth(method = "glm", se=FALSE)+
  theme_bw()+
  ggtitle("sceptical")
plot_grid(plot_i2, plot_s2, plot_i3, plot_s3, plot_i4, plot_s4, plot_i6, plot_s6)
```

```
## `geom_smooth()` using formula = 'y ~ x'
```



fit and asses a classification algorithm on training data (Baysian)

```
##Setting up the model
PR_f0 <-bf(Group~1+V1+V2+V3+V4+V5+V6+V7+V8+V9+V10)

PR_f1 <-bf(Group~1+V1+V2+V3+V4+V5+V6+V7+V8+V9+V10+(1|ID))
get_prior(PR_f0, train_informed_s, family = bernoulli)</pre>
```

```
##
                     prior
                                class coef group resp dpar nlpar 1b ub
                                                                                  source
##
                    (flat)
                                    b
                                                                                 default
##
                    (flat)
                                    b
                                         V1
                                                                            (vectorized)
##
                    (flat)
                                    b
                                        V10
                                                                            (vectorized)
##
                    (flat)
                                    b
                                         V2
                                                                            (vectorized)
##
                    (flat)
                                    b
                                         V3
                                                                            (vectorized)
##
                    (flat)
                                    b
                                         V4
                                                                            (vectorized)
##
                    (flat)
                                    b
                                         V5
                                                                            (vectorized)
##
                    (flat)
                                         V6
                                                                            (vectorized)
                                    b
##
                    (flat)
                                    b
                                         V7
                                                                            (vectorized)
##
                    (flat)
                                    b
                                         V8
                                                                            (vectorized)
##
                    (flat)
                                    b
                                         V9
                                                                            (vectorized)
##
                                                                                 default
    student_t(3, 0, 2.5) Intercept
```

```
get_prior(PR_f1, train_informed_s, family = bernoulli)
```

```
##
                     prior
                                class
                                            coef group resp dpar nlpar lb ub
##
                    (flat)
                                     b
##
                    (flat)
                                     b
                                              V1
##
                    (flat)
                                     b
                                              V10
##
                    (flat)
                                              V2
                                     b
##
                    (flat)
                                     b
                                               V3
##
                    (flat)
                                     b
                                              V4
                                               V5
##
                    (flat)
                                     b
##
                                               V6
                    (flat)
                                     b
##
                                              V7
                    (flat)
                                     b
##
                    (flat)
                                     b
                                               V8
##
                    (flat)
                                              V9
    student_t(3, 0, 2.5) Intercept
##
##
    student t(3, 0, 2.5)
                                                                            0
    student_t(3, 0, 2.5)
##
                                                                            0
                                   sd
                                                     ID
                                                                            0
##
    student_t(3, 0, 2.5)
                                   sd Intercept
                                                     TD
##
           source
##
          default
##
    (vectorized)
##
    (vectorized)
##
    (vectorized)
##
    (vectorized)
##
    (vectorized)
##
    (vectorized)
    (vectorized)
##
##
    (vectorized)
##
    (vectorized)
##
    (vectorized)
##
          default
##
          default
##
    (vectorized)
##
    (vectorized)
```

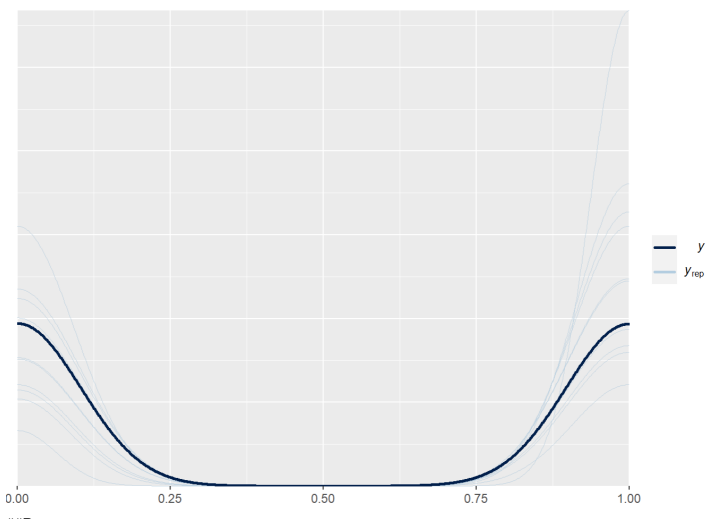
Setting priors

```
PR_p0 <- c(
    prior(normal(0, 1), class=Intercept),
    prior(normal(0, 0.3), class=b)
)

PR_p1 <- c(
    prior(normal(0, 1), class=Intercept),
    prior(normal(0, 0.3), class=sd),
    prior(normal(0, 0.3), class=b)
)</pre>
```

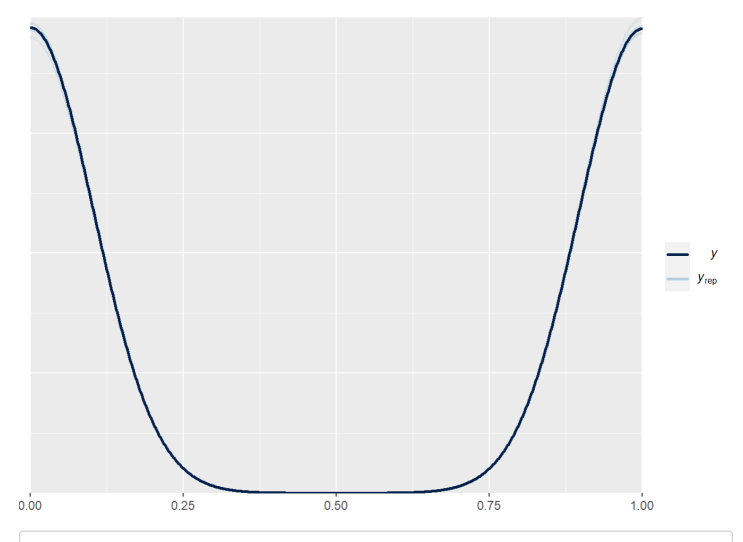
Fitting the first model on both the skeptical and informed data

```
##Model fit on priors
pr_m0_inf <- brm(</pre>
  PR_f0,
  data = train_informed_s,
  prior = PR_p0,
  family = bernoulli,
  refresh=0,
  sample prior = 'only',
  iter=6000,
  warmup = 2500,
  backend = "cmdstanr",
  threads = threading(2),
  chains = 4,
  cores = 4,
  control = list(
    adapt delta = 0.9,
    max\_treedepth = 20)
)
#pp_check(pr_m0_inf)
pr_m0_skep <- brm(</pre>
  PR_f0,
  data = train_skeptic_s,
  prior = PR_p0,
  family = bernoulli,
  refresh=0,
  sample prior = 'only',
  iter=6000,
  warmup = 2500,
  backend = "cmdstanr",
  threads = threading(2),
  chains = 4,
  cores = 4,
  control = list(
    adapt_delta = 0.9,
    max_treedepth = 20)
)
pp_check(pr_m0_skep)
```

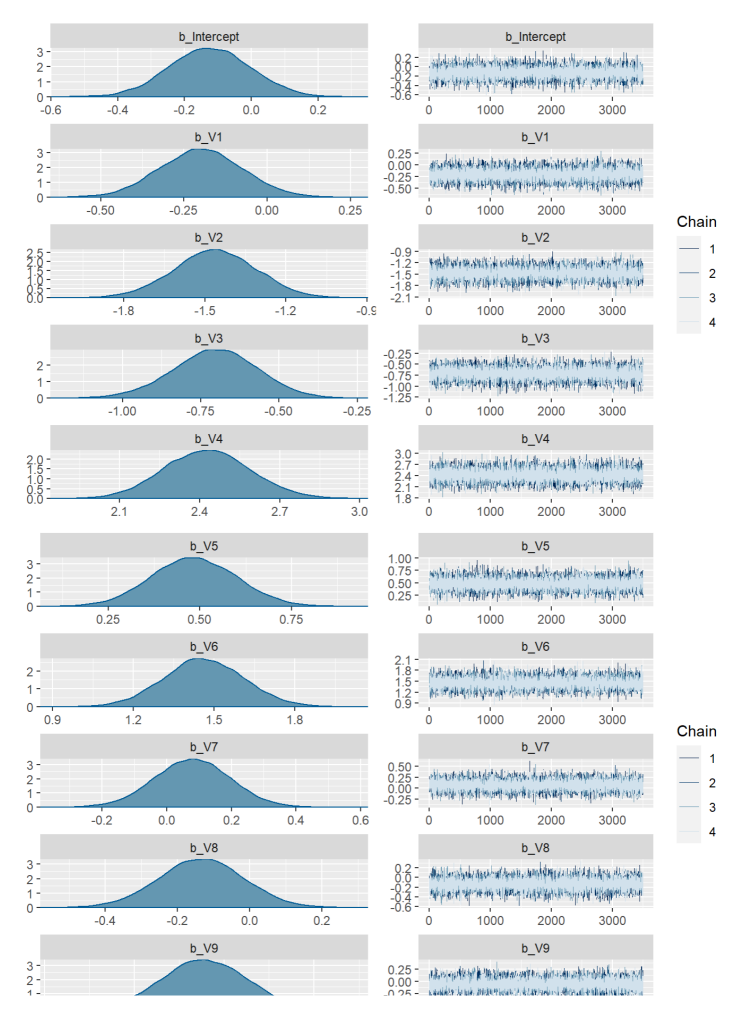


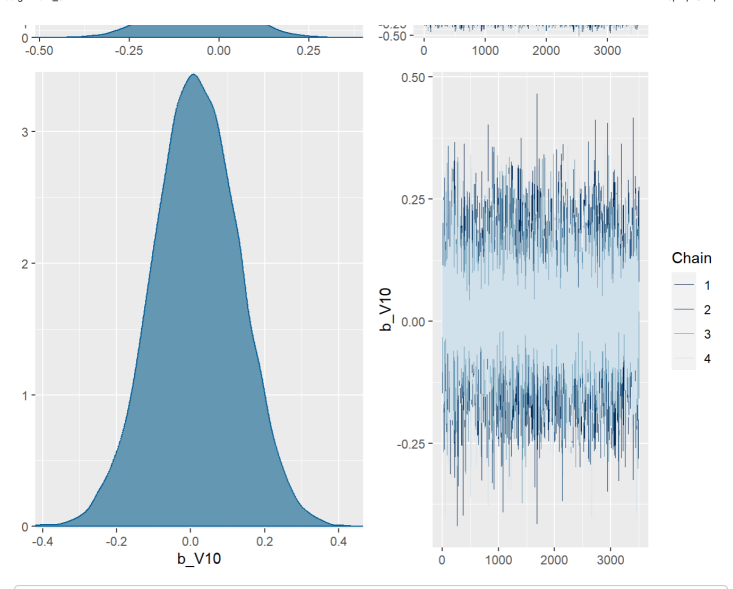
##Bryan

```
##Model fit on informed data
pr_m0_fit_inf <- brm(</pre>
  PR_f0,
  data = train_informed_s,
  prior = PR_p0,
  family = bernoulli,
  refresh=0,
  sample_prior = TRUE,
  iter=6000,
  warmup = 2500,
  backend = "cmdstanr",
  threads = threading(2),
  chains = 4,
  cores = 4,
  control = list(
    adapt_delta = 0.9,
    max_treedepth = 20)
)
pp_check(pr_m0_fit_inf)
```



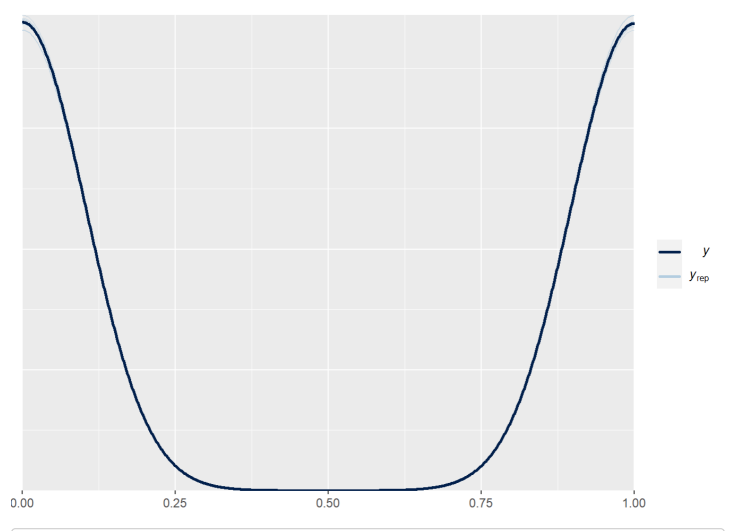
plot(pr_m0_fit_inf)



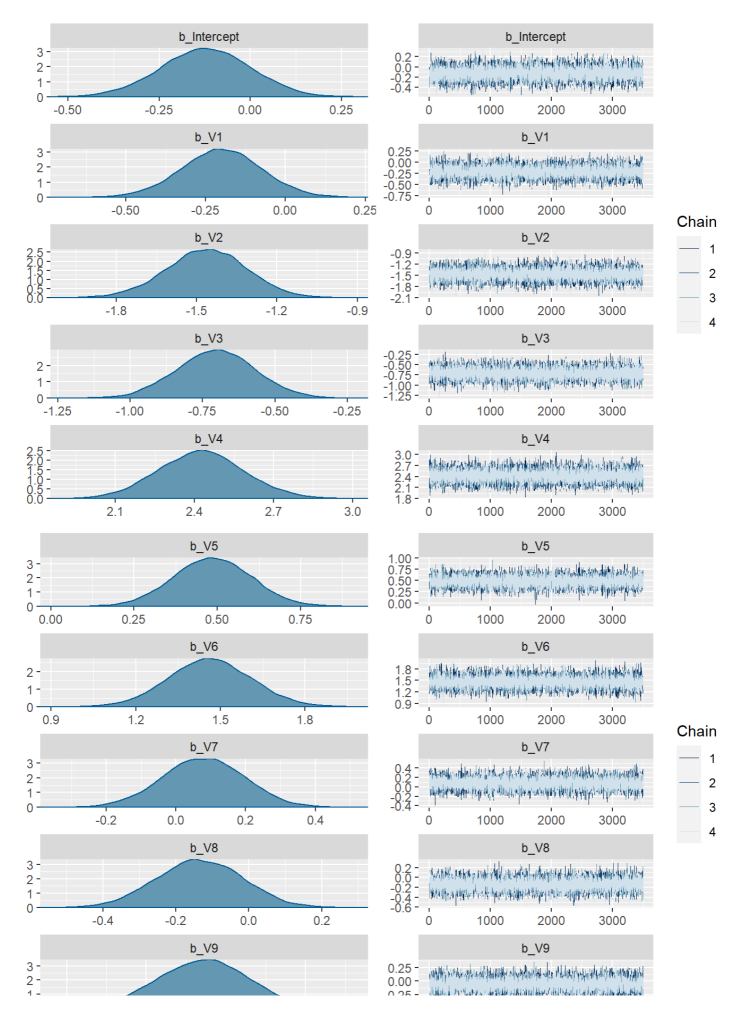


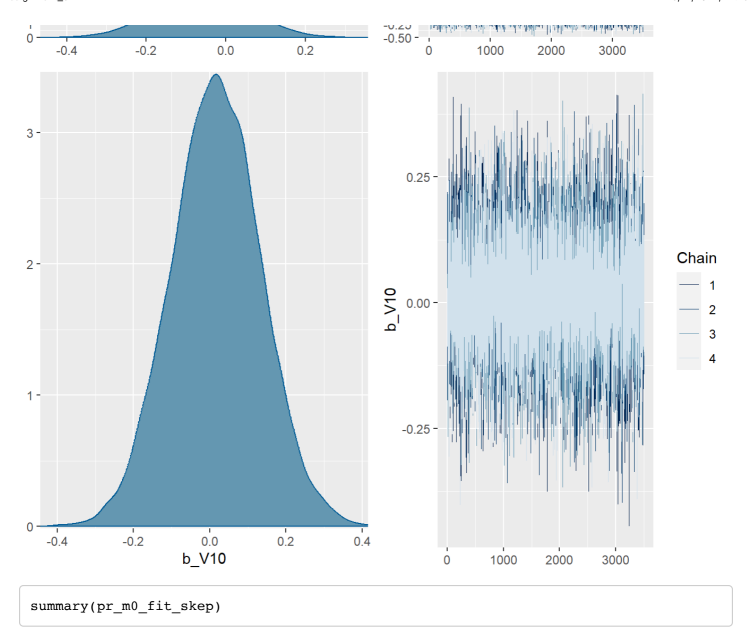
summary(pr_m0_fit_inf)

```
##Model fit on skeptical data
pr_m0_fit_skep <- brm(</pre>
  PR_f0,
  data = train_skeptic_s,
  prior = PR_p0,
  family = bernoulli,
  refresh=0,
  sample prior = TRUE,
  iter=6000,
  warmup = 2500,
  backend = "cmdstanr",
  threads = threading(2),
  chains = 4,
  cores = 4,
  control = list(
    adapt_delta = 0.9,
    max\_treedepth = 20)
)
pp_check(pr_m0_fit_skep)
```



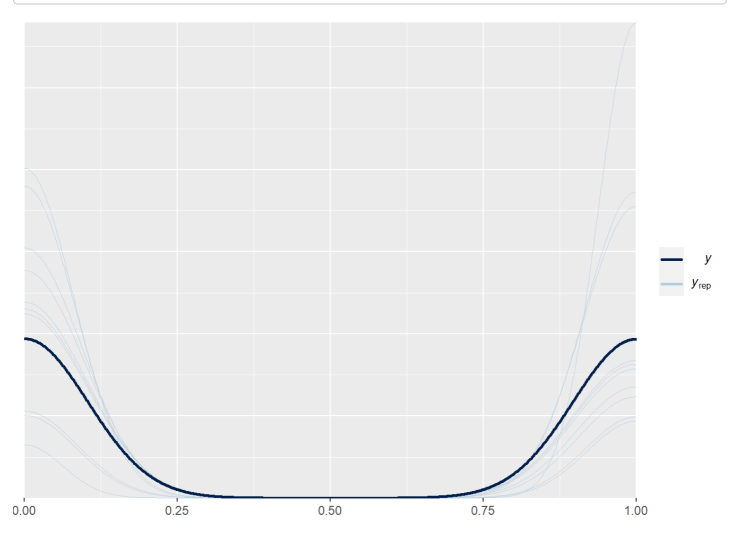
plot(pr_m0_fit_skep)



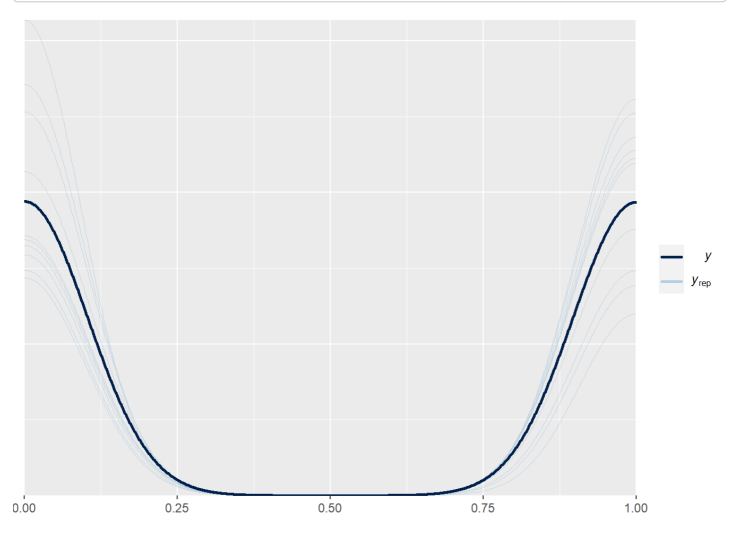


Fitting the second model on skeptical and informed data

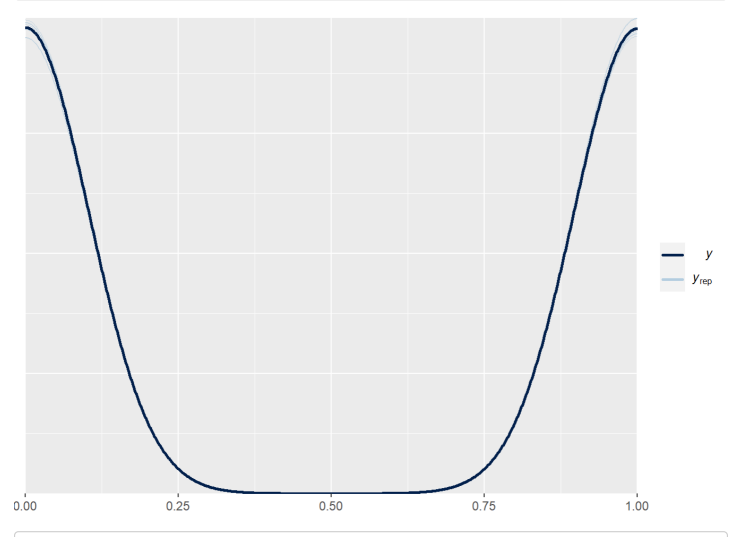
```
##Second model fit on priors
pr_m1_inf <- brm(</pre>
  PR_f1,
  data = train_informed_s,
  prior = PR p1,
  family = bernoulli,
  refresh=0,
  sample prior = 'only',
  iter=6000,
  warmup = 2500,
  backend = "cmdstanr",
  threads = threading(2),
  chains = 4,
  cores = 4,
  control = list(
    adapt_delta = 0.9,
    max\_treedepth = 20)
)
pp_check(pr_m1_inf)
```



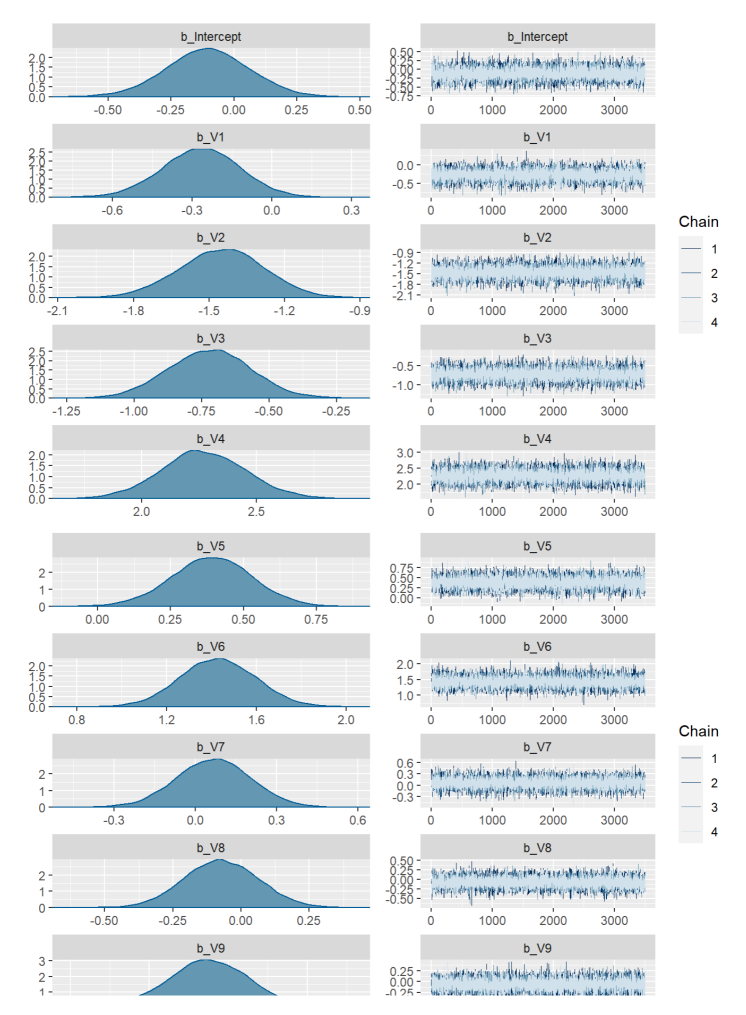
```
pr_m1_skep <- brm(</pre>
  PR_f1,
  data = train_skeptic_s,
  prior = PR_p1,
  family = bernoulli,
  refresh=0,
  sample_prior = 'only',
  iter=6000,
  warmup = 2500,
  backend = "cmdstanr",
  threads = threading(2),
  chains = 4,
  cores = 4,
  control = list(
    adapt_delta = 0.9,
    max\_treedepth = 20)
)
pp_check(pr_m1_skep)
```

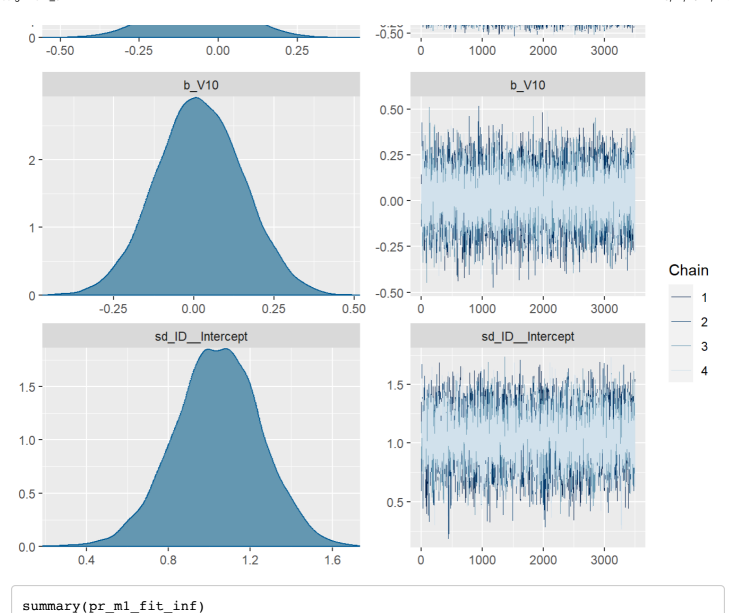


```
##second model fit on informed data
pr_m1_fit_inf <- brm(</pre>
  PR_f1,
  data = train_informed_s,
  prior = PR p1,
  family = bernoulli,
  refresh=0,
  sample prior = TRUE,
  iter=6000,
  warmup = 2500,
  backend = "cmdstanr",
  threads = threading(2),
  chains = 4,
  cores = 4,
  control = list(
    adapt_delta = 0.9,
    max\_treedepth = 20)
)
pp_check(pr_m1_fit_inf)
```

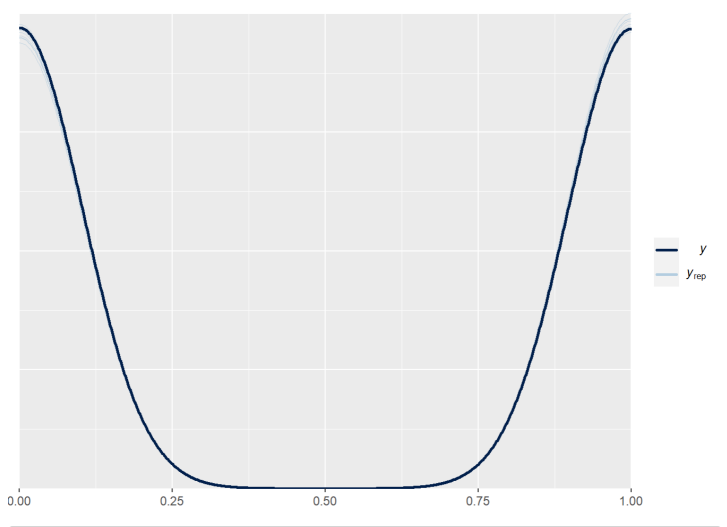


plot(pr_ml_fit_inf)

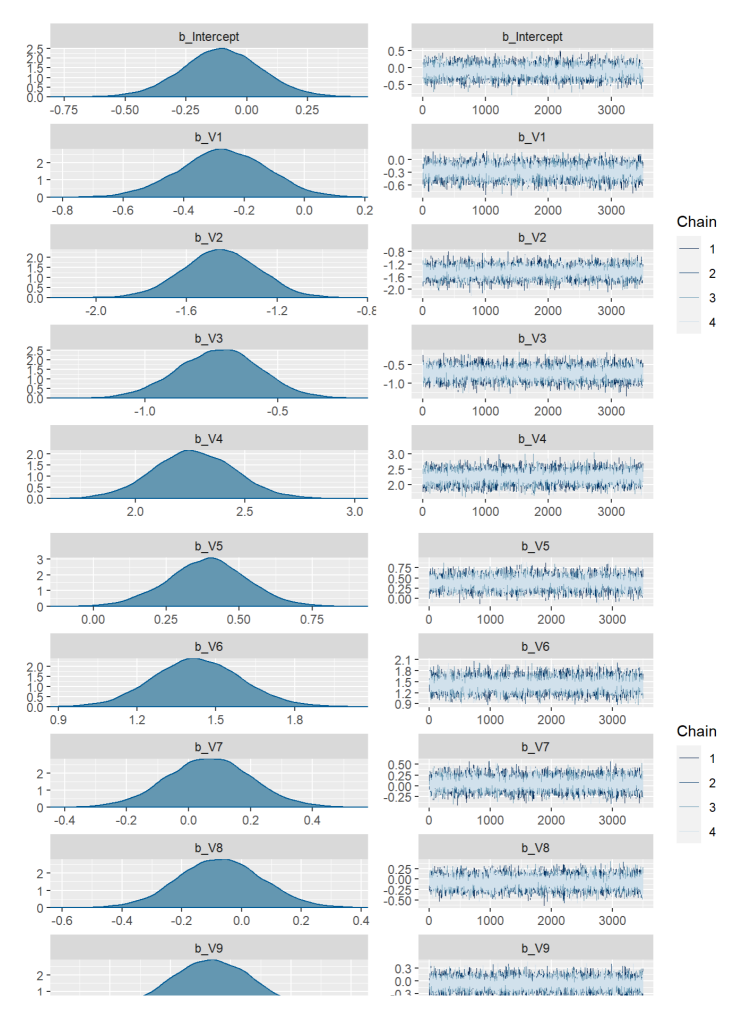


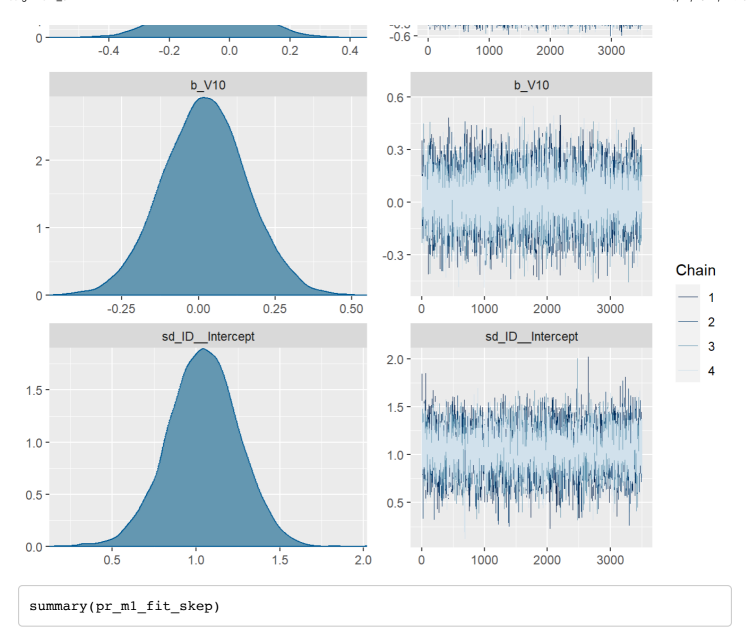


```
## second model fit on skeptical data
pr_m1_fit_skep <- brm(</pre>
  PR_f1,
  data = train_skeptic_s,
  prior = PR p1,
  family = bernoulli,
  refresh=0,
  sample prior = TRUE,
  iter=6000,
  warmup = 2500,
  backend = "cmdstanr",
  threads = threading(2),
  chains = 4,
  cores = 4,
  control = list(
    adapt_delta = 0.9,
    max\_treedepth = 20)
)
pp_check(pr_m1_fit_skep)
```



plot(pr_m1_fit_skep)





##Ditlev ### Asses performance on test data

train_informed_s\$PredictionsPerc0 <- predict(pr_m0_fit_inf)[, 1]
train_informed_s\$Prediction0[train_informed_s\$PredictionsPerc0 > 0.5] <- "Schizoph
renia"</pre>

Warning: Unknown or uninitialised column: `Prediction0`.

train_informed_s\$Prediction0[train_informed_s\$PredictionsPerc0 <= 0.5] <- "Control"</pre>

train_informed_s\$PredictionsPerc1 <- predict(pr_m1_fit_inf)[, 1]
train_informed_s\$Prediction1[train_informed_s\$PredictionsPerc1 > 0.5] <- "Schizoph
renia"</pre>

Warning: Unknown or uninitialised column: `Prediction1`. train informed s\$Prediction1[train informed s\$PredictionsPerc1 <= 0.5] <- "Control train_skeptic_s\$PredictionsPerc0 <- predict(pr_m0_fit_skep)[, 1]</pre> train skeptic s\$Prediction0[train skeptic s\$PredictionsPerc0 > 0.5] <- "Schizophre nia" ## Warning: Unknown or uninitialised column: `Prediction0`. train skeptic s\$Prediction0[train skeptic s\$PredictionsPerc0 <= 0.5] <- "Control" train skeptic s\$PredictionsPerc1 <- predict(pr m1 fit skep)[, 1]</pre> train_skeptic_s\$Prediction1[train_skeptic_s\$PredictionsPerc1 > 0.5] <- "Schizophre nia" ## Warning: Unknown or uninitialised column: `Prediction1`. train_skeptic_s\$Prediction1[train_skeptic_s\$PredictionsPerc1 <= 0.5] <- "Control" train informed s <- train informed s %>% mutate(Group = as.factor(Group), Prediction0 = as.factor(Prediction0), Prediction1 = as.factor(Prediction1))

```
train_skeptic_s$Prediction1[train_skeptic_s$PredictionsPerc1 <= 0.5] <- "Control"

train_informed_s <- train_informed_s %>%
    mutate(
    Group = as.factor(Group),
    Prediction0 = as.factor(Prediction0),
    Prediction1 = as.factor(Prediction1)
)

train_skeptic_s <- train_skeptic_s %>%
    mutate(
    Group = as.factor(Group),
    Prediction0 = as.factor(Prediction0),
    Prediction1 = as.factor(Prediction1)
)

test_informed_s$PredictionsPerc0 <- predict(pr_m0_fit_inf, newdata = test_informed_s, allow_new_levels = T)[, 1]
test_informed_s$Prediction0[test_informed_s$PredictionsPerc0 > 0.5] <- "Schizophre nia"</pre>
```

Warning: Unknown or uninitialised column: `Prediction0`.

```
test_informed_s$Prediction0[test_informed_s$PredictionsPerc0 <= 0.5] <- "Control"

test_informed_s$PredictionsPerc1 <- predict(pr_ml_fit_inf, newdata = test_informed
_s, allow_new_levels = T)[, 1]
test_informed_s$Prediction1[test_informed_s$PredictionsPerc1 > 0.5] <- "Schizophre
nia"</pre>
```

Warning: Unknown or uninitialised column: `Prediction1`.

```
test_informed_s$Prediction1[test_informed_s$PredictionsPerc1 <= 0.5] <- "Control"

test_skeptic_s$PredictionsPerc0 <- predict(pr_m0_fit_skep, newdata = test_informed
_s, allow_new_levels = T)[, 1]
test_skeptic_s$Prediction0[test_skeptic_s$PredictionsPerc0 > 0.5] <- "Schizophrenia"</pre>
```

Warning: Unknown or uninitialised column: `Prediction0`.

```
test_skeptic_s$Prediction0[test_skeptic_s$PredictionsPerc0 <= 0.5] <- "Control"

test_skeptic_s$PredictionsPerc1 <- predict(pr_m1_fit_skep, newdata = test_informed
_s, allow_new_levels = T)[, 1]
test_skeptic_s$Prediction1[test_skeptic_s$PredictionsPerc1 > 0.5] <- "Schizophrenia"</pre>
```

Warning: Unknown or uninitialised column: `Prediction1`.

```
test_skeptic_s$Prediction1[test_skeptic_s$PredictionsPerc1 <= 0.5] <- "Control"

test_informed_s <- test_informed_s %>%
    mutate(
        Group = as.factor(Group),
        Prediction0 = as.factor(Prediction0),
        Prediction1 = as.factor(Prediction1)
)

test_skeptic_s <- test_skeptic_s %>%
    mutate(
        Group = as.factor(Group),
        Prediction0 = as.factor(Prediction0),
        Prediction1 = as.factor(Prediction1)
)
```

confusionMatrix(train_skeptic_s\$Group, train_skeptic_s\$Prediction0)

```
## Confusion Matrix and Statistics
##
##
                  Reference
## Prediction
                   Control Schizophrenia
##
     Control
                       778
                                       23
     Schizophrenia
                        15
                                      784
##
##
##
                  Accuracy : 0.9762
                    95% CI: (0.9675, 0.9831)
##
##
       No Information Rate: 0.5044
       P-Value [Acc > NIR] : <2e-16
##
##
##
                     Kappa: 0.9525
##
##
    Mcnemar's Test P-Value: 0.2561
##
##
               Sensitivity: 0.9811
##
               Specificity: 0.9715
            Pos Pred Value: 0.9713
##
##
            Neg Pred Value: 0.9812
##
                Prevalence: 0.4956
            Detection Rate: 0.4863
##
      Detection Prevalence: 0.5006
##
##
         Balanced Accuracy: 0.9763
##
          'Positive' Class : Control
##
##
```

```
confusionMatrix(test_skeptic_s$Group, test_skeptic_s$Prediction0)
```

```
## Confusion Matrix and Statistics
##
##
                  Reference
## Prediction
                   Control Schizophrenia
##
     Control
                       198
     Schizophrenia
                         5
##
                                      188
##
##
                  Accuracy: 0.965
                    95% CI: (0.942, 0.9807)
##
##
       No Information Rate: 0.5075
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa : 0.93
##
    Mcnemar's Test P-Value: 0.4227
##
##
##
               Sensitivity: 0.9754
##
               Specificity: 0.9543
##
            Pos Pred Value: 0.9565
##
            Neg Pred Value: 0.9741
##
                Prevalence: 0.5075
            Detection Rate: 0.4950
##
##
      Detection Prevalence: 0.5175
##
         Balanced Accuracy: 0.9648
##
          'Positive' Class : Control
##
##
```

```
confusionMatrix(train_skeptic_s$Group, train_skeptic_s$Prediction1)
```

```
## Confusion Matrix and Statistics
##
##
                  Reference
## Prediction
                   Control Schizophrenia
##
     Control
                       797
     Schizophrenia
                         2
                                      797
##
##
##
                  Accuracy : 0.9962
                    95% CI: (0.9919, 0.9986)
##
##
       No Information Rate: 0.5006
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa: 0.9925
##
    Mcnemar's Test P-Value: 0.6831
##
##
##
               Sensitivity: 0.9975
##
               Specificity: 0.9950
            Pos Pred Value: 0.9950
##
##
            Neg Pred Value: 0.9975
                Prevalence: 0.4994
##
            Detection Rate: 0.4981
##
##
      Detection Prevalence: 0.5006
##
         Balanced Accuracy: 0.9963
##
          'Positive' Class : Control
##
##
```

```
confusionMatrix(test_skeptic_s$Group, test_skeptic_s$Prediction1)
```

```
## Confusion Matrix and Statistics
##
##
                  Reference
## Prediction
                   Control Schizophrenia
##
     Control
                       199
     Schizophrenia
                         2
                                      191
##
##
##
                  Accuracy: 0.975
                    95% CI: (0.9545, 0.9879)
##
##
       No Information Rate: 0.5025
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa : 0.95
##
    Mcnemar's Test P-Value: 0.1138
##
##
##
               Sensitivity: 0.9900
##
               Specificity: 0.9598
##
            Pos Pred Value: 0.9614
##
            Neg Pred Value: 0.9896
##
                Prevalence: 0.5025
            Detection Rate: 0.4975
##
##
      Detection Prevalence: 0.5175
##
         Balanced Accuracy: 0.9749
##
          'Positive' Class : Control
##
##
```

```
confusionMatrix(train_informed_s$Group, train_informed_s$Prediction0)
```

```
## Confusion Matrix and Statistics
##
##
                  Reference
## Prediction
                   Control Schizophrenia
##
     Control
                       777
                                       24
     Schizophrenia
                        15
                                      784
##
##
##
                  Accuracy : 0.9756
                    95% CI: (0.9668, 0.9826)
##
##
       No Information Rate: 0.505
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa : 0.9513
##
    Mcnemar's Test P-Value: 0.2002
##
##
##
               Sensitivity: 0.9811
##
               Specificity: 0.9703
##
            Pos Pred Value: 0.9700
##
            Neg Pred Value: 0.9812
##
                Prevalence: 0.4950
            Detection Rate: 0.4856
##
##
      Detection Prevalence: 0.5006
##
         Balanced Accuracy: 0.9757
##
##
          'Positive' Class : Control
##
```

```
confusionMatrix(test_informed_s$Group, test_informed_s$Prediction0)
```

```
## Confusion Matrix and Statistics
##
##
                  Reference
## Prediction
                   Control Schizophrenia
##
     Control
                       198
                                        1
     Schizophrenia
                         5
                                      196
##
##
##
                  Accuracy: 0.985
                    95% CI: (0.9676, 0.9945)
##
##
       No Information Rate: 0.5075
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa : 0.97
##
    Mcnemar's Test P-Value: 0.2207
##
##
##
               Sensitivity: 0.9754
##
               Specificity: 0.9949
##
            Pos Pred Value: 0.9950
##
            Neg Pred Value: 0.9751
##
                Prevalence: 0.5075
            Detection Rate: 0.4950
##
##
      Detection Prevalence: 0.4975
##
         Balanced Accuracy: 0.9851
##
          'Positive' Class : Control
##
##
```

```
confusionMatrix(train_informed_s$Group, train_informed_s$Prediction1)
```

```
## Confusion Matrix and Statistics
##
##
                  Reference
## Prediction
                   Control Schizophrenia
##
     Control
                       797
     Schizophrenia
                         2
                                      797
##
##
##
                  Accuracy : 0.9962
                    95% CI: (0.9919, 0.9986)
##
##
       No Information Rate: 0.5006
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa: 0.9925
##
    Mcnemar's Test P-Value: 0.6831
##
##
##
               Sensitivity: 0.9975
##
               Specificity: 0.9950
##
            Pos Pred Value: 0.9950
##
            Neg Pred Value: 0.9975
##
                Prevalence: 0.4994
            Detection Rate: 0.4981
##
##
      Detection Prevalence: 0.5006
##
         Balanced Accuracy: 0.9963
##
          'Positive' Class : Control
##
##
```

```
confusionMatrix(test_informed_s$Group, test_informed_s$Prediction1)
```

```
## Confusion Matrix and Statistics
##
                  Reference
##
## Prediction
                   Control Schizophrenia
##
     Control
                       199
##
     Schizophrenia
                         2
                                      199
##
##
                  Accuracy: 0.995
##
                    95% CI: (0.9821, 0.9994)
       No Information Rate: 0.5025
##
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa : 0.99
##
    Mcnemar's Test P-Value: 0.4795
##
##
               Sensitivity: 0.9900
##
##
               Specificity: 1.0000
##
            Pos Pred Value: 1.0000
##
            Neg Pred Value: 0.9900
                Prevalence: 0.5025
##
            Detection Rate: 0.4975
##
##
      Detection Prevalence: 0.4975
##
         Balanced Accuracy: 0.9950
##
##
          'Positive' Class : Control
##
```

Discuss whether performance and feature importance is as expected

##Part 3 - Applying the machine learning pipeline to empirical data

```
real_data <- read_csv("assignment_3_data.csv")
```

```
## Rows: 1889 Columns: 398
## — Column specification
-
## Delimiter: ","
## chr (5): NewID, Diagnosis, Language, Gender, Trial
## dbl (393): PatID, Corpus, Duration_Praat, F0_Mean_Praat, F0_SD_Praat, Intens...
##
## i Use `spec()` to retrieve the full column specification for this data.
## i Specify the column types or set `show_col_types = FALSE` to quiet this messag e.
```

```
real_data <- real_data %>%
  select(-Language) %>%
  select(-Corpus) %>%
  select(-NewID)
```

Apply your machine learning pipeline to the empirical data

Warning: in simulated data you only have 10 features, now you have many more - Consider the impact a higher number of features will have on your ML inference and decide if you want to cut down the number of features before running the pipeline (alternatibely expand the pipeline to add feature selection)

data budgeting and pree-processing of the data

```
set.seed(304)
split_real_data <- initial_split(real_data, prop = 4/5, strata = Gender)
train_data <- training(split_real_data)
test_data <- testing(split_real_data)

train_data <- train_data %>%
    select(-Gender) %>%
    select(-Trial)

test_data <- test_data %>%
    select(-Gender) %>%
    select(-Trial)

train_data$PatID <- as.factor(train_data$PatID)

test_data$PatID <- as.factor(test_data$PatID)</pre>
```

```
recipe_real <- train_data %>%
  recipe(Diagnosis~.) %>%
  update_role(PatID, new_role = 'PatID') %>%
  step_scale(all_numeric()) %>%
  step_center(all_numeric()) %>%
  prep(training=train_data, retain=TRUE)

train_data_s <- juice(recipe_real)
test_data_s <- bake(recipe_real, new_data = test_data)</pre>
```

Principle component analysis

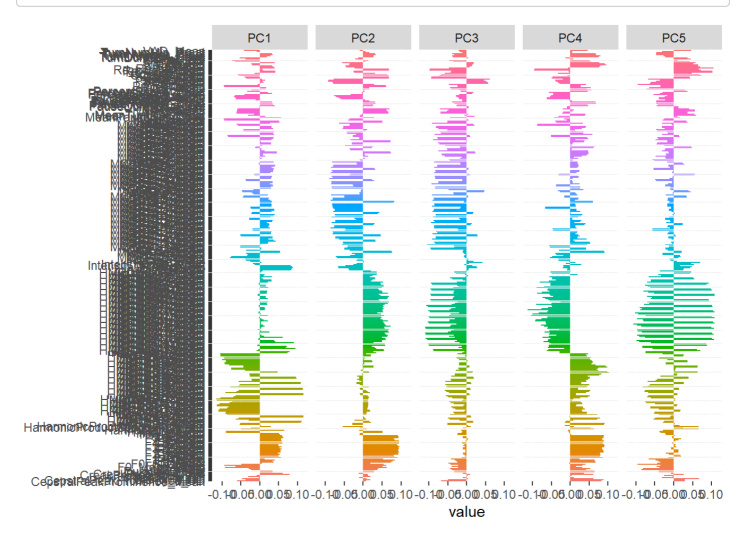
```
pca_recipe <- recipe(Diagnosis~., data = train_data_s) %>%
  update_role(PatID, new_role = "id") %>%
  step_pca(all_numeric(), id = "pca")%>%
  prep()

tidy_pca <- tidy(pca_recipe, 1)

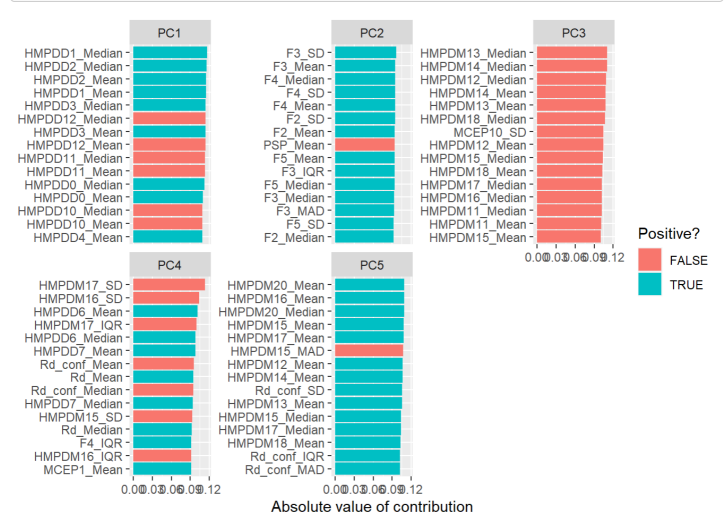
bake_pca <- bake(pca_recipe, train_data_s)

pca_b_test <- bake(pca_recipe, test_data_s)

tidy_pca %>%
  filter(component %in% paste0("PC", 1:5)) %>%
  mutate(component = fct_inorder(component)) %>%
  ggplot(aes(value, terms, fill = terms)) +
  geom_col(show.legend = FALSE) +
  facet_wrap(~component, nrow = 1) +
  labs(y = NULL)
```



```
tidy_pca %>%
  filter(component %in% paste0("PC", 1:5)) %>%
  group_by(component) %>%
  top_n(15, abs(value)) %>%
  ungroup() %>%
  mutate(terms = reorder_within(terms, abs(value), component)) %>%
  ggplot(aes(abs(value), terms, fill = value > 0)) +
  geom_col() +
  facet_wrap(~component, scales = "free_y") +
  scale_y_reordered() +
  labs(
    x = "Absolute value of contribution",
    y = NULL, fill = "Positive?"
)
```



??reorder_within

starting httpd help server ... done

##Manuela ### feature selection

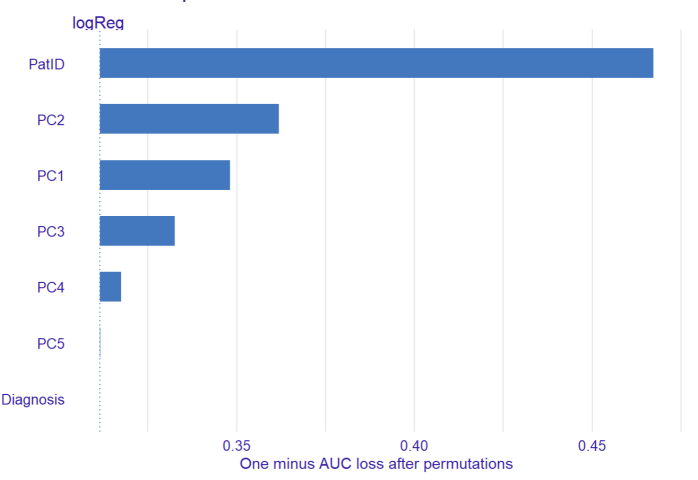
```
d_data <- bake_pca %>%
  mutate(PatID = NULL, Trail = NULL, NewID = NULL, Gender = NULL, Language = NULL,
Corpus = NULL)

LogisticRegression_train<- logistic_reg() %>%
  set_mode('classification') %>%
  set_engine('glm') %>%
  fit(Diagnosis ~ . , data = bake_pca)
```

```
explainer_lm <-
  explain_tidymodels(
    LogisticRegression_train,
    data = bake_pca,
    y = as.numeric(d_data$Diagnosis) -1,
    label = 'logReg',
    verbose = FALSE
)

explainer_lm %>% model_parts() %>% plot(show_boxplots = FALSE) + ggtitle('Feature importance', ''')
```

Feature importance



fit and assess a classification algorithm on the training data

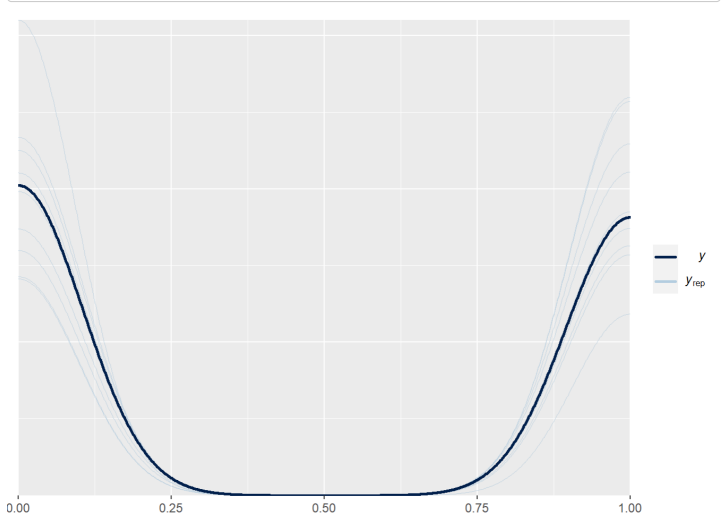
```
form1 <- bf(Diagnosis~1+PC2+PC1+PC3+PC4+(1|PatID))
get_prior(form1, bake_pca, family = bernoulli)</pre>
```

```
##
                    prior
                               class
                                            coef group resp dpar nlpar lb ub
##
                    (flat)
##
                    (flat)
                                    b
                                             PC1
##
                    (flat)
                                             PC2
##
                                             PC3
                    (flat)
                                    b
##
                    (flat)
                                             PC4
##
    student t(3, 0, 2.5) Intercept
##
    student_t(3, 0, 2.5)
                                                                          0
    student_t(3, 0, 2.5)
##
                                                 PatID
    student_t(3, 0, 2.5)
                                                                           0
##
                                   sd Intercept PatID
##
           source
##
         default
##
    (vectorized)
##
    (vectorized)
##
    (vectorized)
    (vectorized)
##
         default
##
##
          default
##
    (vectorized)
##
    (vectorized)
```

```
prior_f1 <- c(
  prior(normal(0, 1), class=Intercept),
  prior(normal(0, 1), class=sd),
  prior(normal(0, 0.3), class=b)
)</pre>
```

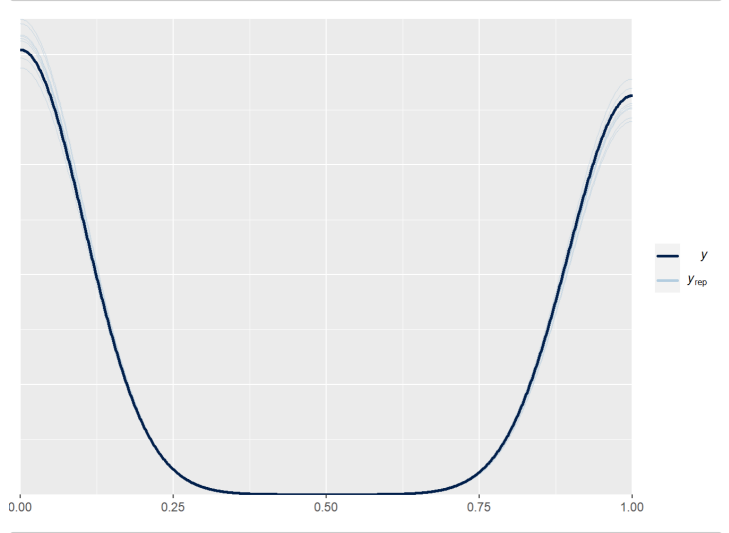
Fitting the model with only the priors

```
pr_m1 <- brm(</pre>
  form1,
  data = bake_pca,
  prior = prior_f1,
  family = bernoulli,
  refresh=0,
  sample_prior = 'only',
  iter=6000,
  warmup = 2500,
  backend = "cmdstanr",
  threads = threading(2),
  chains = 4,
  cores = 4,
  control = list(
    adapt_delta = 0.9,
    max\_treedepth = 20)
)
pp_check(pr_m1)
```

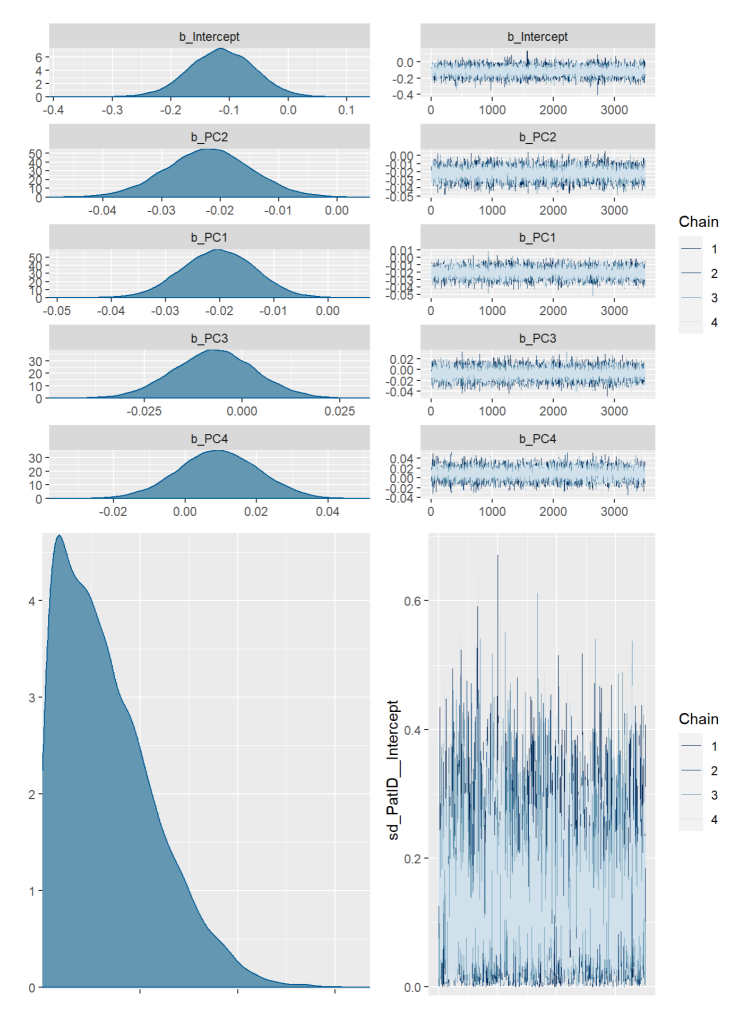


Fitting model on the data

```
pr_m1_fit <- brm(</pre>
  form1,
  data = bake_pca,
  prior = prior_f1,
  family = bernoulli,
  refresh=0,
  sample_prior = TRUE,
  iter=6000,
  warmup = 2500,
  backend = "cmdstanr",
  threads = threading(2),
  chains = 4,
  cores = 4,
  control = list(
    adapt_delta = 0.9,
    max\_treedepth = 20)
)
pp_check(pr_m1_fit)
```



plot(pr_m1_fit)



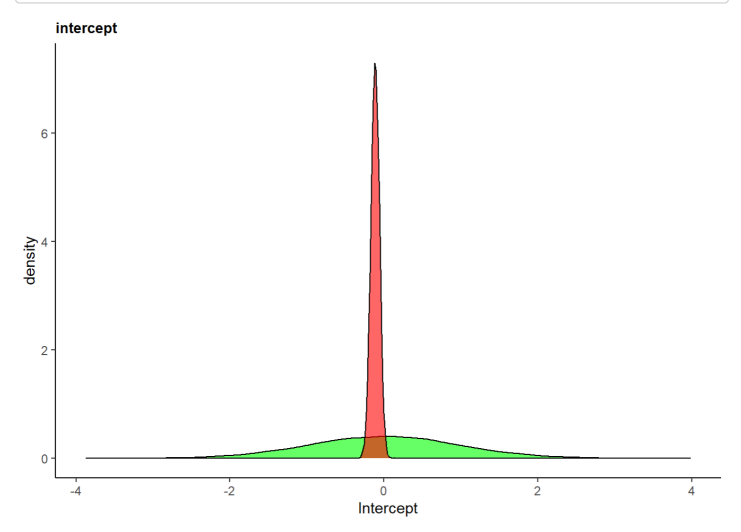


```
summary(pr_m1_fit)
```

Prior posterior update check

```
Posterior_f1 <- as_draws_df(pr_ml_fit)

ggplot(Posterior_f1) +
  geom_density(aes(prior_Intercept), fill="green", color="black",alpha=0.6) +
  geom_density(aes(b_Intercept), fill="red", color="black",alpha=0.6) +
  xlab('Intercept') +
  theme_classic()+
  ggtitle("intercept")+
  theme(plot.title = element_text(size = 10, face = "bold"))</pre>
```



assess performance on the test set

```
bake_pca$PredictionsPerc1 <- predict(pr_m1_fit)[, 1]
bake_pca$Prediction1[bake_pca$PredictionsPerc1 > 0.5] <- "SCZ"</pre>
```

```
## Warning: Unknown or uninitialised column: `Prediction1`.
```

```
bake_pca$Prediction1[bake_pca$PredictionsPerc1 <= 0.5] <- "CT"

bake_pca <- bake_pca %>%
    mutate(
        Diagnosis = as.factor(Diagnosis),
        Prediction1 = as.factor(Prediction1))

pca_b_test$PredictionsPerc1 <- predict(pr_m1_fit, newdata = pca_b_test, allow_new_levels = T)[, 1]

pca_b_test$Prediction1[pca_b_test$PredictionsPerc1 > 0.5] <- "SCZ"</pre>
```

```
## Warning: Unknown or uninitialised column: `Prediction1`.
```

```
pca_b_test$Prediction1[pca_b_test$PredictionsPerc1 <= 0.5] <- "CT"

pca_b_test <- pca_b_test %>%
   mutate(
    Diagnosis = as.factor(Diagnosis),
    Prediction1 = as.factor(Prediction1)
)
```

Accuracy

```
confusionMatrix(bake pca$Diagnosis, bake pca$Prediction1)
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction CT SCZ
##
          CT 545 251
          SCZ 444 270
##
##
##
                  Accuracy: 0.5397
                    95% CI: (0.5142, 0.5651)
##
##
       No Information Rate: 0.655
##
       P-Value [Acc > NIR] : 1
##
##
                     Kappa : 0.0637
##
    Mcnemar's Test P-Value: 3.265e-13
##
##
##
               Sensitivity: 0.5511
##
               Specificity: 0.5182
##
            Pos Pred Value: 0.6847
##
            Neg Pred Value: 0.3782
                Prevalence: 0.6550
##
            Detection Rate: 0.3609
##
##
      Detection Prevalence: 0.5272
##
         Balanced Accuracy: 0.5346
##
          'Positive' Class : CT
##
##
```

```
confusionMatrix(pca_b_test$Diagnosis, pca_b_test$Prediction1)
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction CT SCZ
##
          CT 115
                   78
          SCZ 104 82
##
##
                  Accuracy : 0.5198
##
##
                    95% CI: (0.4682, 0.5711)
##
       No Information Rate: 0.5778
##
       P-Value [Acc > NIR] : 0.99011
##
##
                     Kappa : 0.0368
##
    Mcnemar's Test P-Value: 0.06386
##
##
##
               Sensitivity: 0.5251
##
               Specificity: 0.5125
##
            Pos Pred Value: 0.5959
##
            Neg Pred Value: 0.4409
##
                Prevalence: 0.5778
            Detection Rate: 0.3034
##
##
      Detection Prevalence: 0.5092
##
         Balanced Accuracy: 0.5188
##
          'Positive' Class : CT
##
##
```

###discuss whether performance and feature importance is as expected