Classification with linear SVM

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Load Packages

If the libraries are not installed yet, you need to install them using, for example, the command: install.packages("ggplot2"). For the Hrate package this is different, since it comes from github. The devtools library needs to be installed, and then the install github() function is used.

```
library(dplyr)
library(e1071)
library(class)
library(gmodels)
library(ggpubr)
library(caret)
```

Load Data

Load data table with values per text file.

```
# load estimations from stringBase corpus
estimations.df <- read.csv("~/Github/NaLaFi/results/features.csv")
#head(features.csv)</pre>
```

Exclude subcorpora (if needed).

```
#selected <- c("shuffled", "random") #estimations.df <- estimations.df[!(estimations.df$subcorpus %in% selected), ]
```

Split into separate files by length of chunks in characters.

```
# choose number of characters
num.char = 10
# subset data frame
estimations.df <- estimations.df[estimations.df$num.char == num.char, ]</pre>
```

Select relevant columns of the data frame, i.e. the measures to be included in classification and the "corpus" or "subcorpus" column.

```
Remove NAs (whole row)
estimations.subset <- na.omit(estimations.subset)
```

Center and scale the data

```
estimations.scaled <- cbind(estimations.subset[1], scale(estimations.subset[2:ncol(estimations.subset)]
```

Create Training and Test Sets

```
# Generating seed
set.seed(1234)
# Randomly generating our training and test samples with a respective ratio of 2/3 and 1/3
datasample <- sample(2, nrow(estimations.scaled), replace = TRUE, prob = c(0.67, 0.33))
# Generate training set
train <- estimations.scaled[datasample == 1, 1:ncol(estimations.scaled)]
# Generate test set
test <- estimations.scaled[datasample == 2, 1:ncol(estimations.scaled)]</pre>
```

Building SVM

The following code to build a linear SVM is adopted from https://rpubs.com/cliex159/865583 (last accessed 17.01.2023).

```
# choose kernel
kernel = "sigmoid" # "linear", "polynomial", "radial", "sigmoid"
# svm classification
svm.model <- svm(as.factor(corpus) ~ .,</pre>
                data = train,
                type = "C-classification",
                kernel = kernel,
                scale = FALSE)
# list components of model
names(svm.model)
## [1] "call"
                           "type"
                                              "kernel"
                                                                 "cost"
## [5] "degree"
                           "gamma"
                                              "coef0"
                                                                 "nu"
## [9] "epsilon"
                                              "scaled"
                           "sparse"
                                                                 "x.scale"
## [13] "y.scale"
                           "nclasses"
                                              "levels"
                                                                 "tot.nSV"
## [17] "nSV"
                           "labels"
                                              "SV"
                                                                 "index"
## [21] "rho"
                           "compprob"
                                              "probA"
                                                                 "probB"
## [25] "sigma"
                           "coefs"
                                              "na.action"
                                                                 "fitted"
## [29] "decision.values" "terms"
# check the levels
svm.model$levels
```

Prediction

```
Make predictions using the svm model.
```

```
svm.prediction <- predict(svm.model, test)</pre>
```

Model evaluation

```
# creating a dataframe from known (true) test labels
test.labels <- data.frame(test$corpus)</pre>
# combining predicted and known classes
class.comparison <- data.frame(svm.prediction, test.labels)</pre>
# giving appropriate column names
names(class.comparison) <- c("predicted", "observed")</pre>
# inspecting our results table
head(class.comparison)
##
        predicted
                     observed
## 5 non-writing non-writing
## 11 non-writing non-writing
## 14 non-writing non-writing
## 16 non-writing non-writing
## 26 non-writing non-writing
## 28 non-writing non-writing
# get confusion matrix
cm <- confusionMatrix(as.factor(class.comparison$predicted),</pre>
                       reference = as.factor(class.comparison$observed))
print(cm)
## Confusion Matrix and Statistics
##
                Reference
##
## Prediction
                 non-writing writing
     non-writing
##
                         265
                                  193
     writing
                          223
                                  282
##
##
##
                  Accuracy: 0.568
##
                    95% CI: (0.536, 0.5996)
       No Information Rate: 0.5067
##
##
       P-Value [Acc > NIR] : 7.94e-05
##
##
                     Kappa: 0.1366
##
##
    Mcnemar's Test P-Value: 0.1551
##
##
               Sensitivity: 0.5430
               Specificity: 0.5937
##
##
            Pos Pred Value: 0.5786
##
            Neg Pred Value: 0.5584
##
                Prevalence: 0.5067
##
            Detection Rate: 0.2752
      Detection Prevalence: 0.4756
##
```

```
##
         Balanced Accuracy: 0.5684
##
##
          'Positive' Class : non-writing
##
# get precision, recall, and f1 from the output list of confusionMatrix()
accuracy <- cm$overall['Accuracy']</pre>
f1 <- cm[["byClass"]]["F1"]</pre>
recall <- cm[["byClass"]]["Recall"]</pre>
precision <- cm[["byClass"]]["Precision"]</pre>
# prepare data frame with results
svm.results <- data.frame(accuracy, precision, recall, f1, row.names = NULL)</pre>
svm.results.rounded <- round(svm.results, 2)</pre>
print(svm.results.rounded)
     accuracy precision recall f1
## 1
         0.57
                   0.58
                          0.54 0.56
Write to file.
write.csv(svm.results.rounded, file = paste(c("~/Github/NaLaFi/results/SVM/results_SVM_",
                                                 num.char, "chars", "_", kernel, ".csv"), collapse = ""),
          row.names = F)
```

Visualization

Build scatter plot of training dataset.

Unigram entropy and TTR

```
# downsample the training dataset (otherwise there is a lot of overplotting)
train.ds <- sample_n(train, 100)</pre>
huni.ttr.plot <- ggplot(data = train.ds, aes(x = huni.chars, y = ttr.chars, color = corpus)) +
    geom_point() +
    labs(x = "Unigram entropy for 100 characters", y = "TTR for 100 characters")
#huni.ttr.plot
# sum classification (with just entropy rate and repetition rate)
svm.model <- svm(as.factor(corpus) ~ huni.chars + ttr.chars,</pre>
                data = train.ds,
                type = "C-classification",
                kernel = "linear",
                scale = FALSE)
w <- t(svm.model$coefs) %*% svm.model$SV
# calculate slope and intercept of decision boundary from weight vector and sum model
slope_1 < -w[1]/w[2]
intercept_1 <- svm.model$rho/w[2]</pre>
#add decision boundary
```

```
plot.decision <- huni.ttr.plot + geom_abline(slope = slope_1, intercept = intercept_1)

#add margin boundaries
plot.margins <- plot.decision +
    geom_abline(slope = slope_1, intercept = intercept_1 - 1/w[2], linetype = "dashed")+
    geom_abline(slope = slope_1, intercept = intercept_1 + 1/w[2], linetype = "dashed")

# add suport vectors
layered.huni.ttr.plot <-
    plot.margins + geom_point(data = train.ds[svm.model$index, ], aes(x = huni.chars, y = ttr.chars), c

# show plot
# layered.huni.ttr.plot</pre>
```

Entropy rate and repetition rate

```
# downsample the training dataset (otherwise there is a lot of overplotting)
train.ds <- sample n(train, 100)</pre>
# create ggplot
hrate.rm.plot <- ggplot(data = train.ds, aes(x = hrate.chars, y = rm.chars, color = corpus)) +</pre>
    geom point() +
    labs(x = "Entropy rate for 100 characters", y = "Repetition rate for 100 characters") +
    theme(legend.position = "none")
#hrate.rm.plot
# sum classification (with just entropy rate and repetition rate)
svm.model <- svm(as.factor(corpus) ~ hrate.chars + rm.chars,</pre>
                data = train.ds,
                type = "C-classification",
                kernel = "linear",
                scale = FALSE)
w <- t(svm.model$coefs) %*% svm.model$SV
# calculate slope and intercept of decision boundary from weight vector and sum model
slope_1 < -w[1]/w[2]
intercept_1 <- svm.model$rho/w[2]</pre>
#add decision boundary
plot.decision <- hrate.rm.plot + geom_abline(slope = slope_1, intercept = intercept_1)</pre>
#add margin boundaries
plot.margins <- plot.decision +</pre>
geom_abline(slope = slope_1, intercept = intercept_1 - 1/w[2], linetype = "dashed")+
geom_abline(slope = slope_1, intercept = intercept_1 + 1/w[2], linetype = "dashed")
# add suport vectors
layered.hrate.rm.plot <-</pre>
    plot.margins + geom_point(data = train.ds[svm.model$index, ], aes(x = hrate.chars, y = rm.chars), c
# show plot
#layered.hrate.rm.plot
```

Combined Plots

Safe complete figure to file

```
ggsave("~/Github/NaLaFi/figures/plots_SVM.pdf", plots.combined, width = 10,
height = 4, dpi = 300, scale = 1, device = cairo_pdf)
```