# 1 Common libraries

# 1.1 dplyr

**dplyr**: A grammar of data manipulation, providing tools for filtering, selecting, mutating, and summarizing data efficiently. It is designed to work seamlessly with data frames and tibbles.

#### 1.2 caret

caret: Provides tools for pre-processing, feature selection, and resampling.

#### 1.3 kernlab

**kernlab**: Focused on kernel-based machine learning methods, this library supports algorithms like Support Vector Machines (SVM), Kernel Principal Component Analysis (KPCA), and Gaussian Processes.

#### 1.4 neuralnet

**neuralnet**: A package for training and visualizing neural networks in R. It supports deep learning on numeric data and allows customization of activation functions and network architecture.

# 1.5 glmnet

**glmnet**: Implements regularized regression methods such as LASSO and Elastic Net, suitable for linear and logistic regression with high-dimensional data.

#### 1.6 e1071

**e1071**: Provides tools for machine learning, including support for SVMs, Naïve Bayes classifiers, clustering algorithms, and statistical functions. It also includes implementations of fuzzy logic.

#### 2 Models

# 2.1 Linear Regression

```
model = lm ( Fat ~ . , data = data_train)
#0r
model=glm(Fat~., data=data_train, family="gaussian")
```

# 2.2 Logistic Regression (classification)

```
glm(formula = as.factor(species)~., data = a, family = "binomial")
```

#### 2.3 Poisson

```
glm(formula = Visitors~., data = data_b, family = "poisson"(link=log))
```

# 2.4 Lasso (L1) and Ridge (L2)

```
# Import glmnet package
 library(glmnet)
3
 # Create the lasso model by setting alpha = 1.
5 lasso <- glmnet(X train matrix, data train$Fat, alpha = 1, family =</pre>
     "gaussian")
 # Cross validation can also be calculated for lasso and ridge models.
 cv lasso=cv.glmnet(X train matrix, data train$Faat, alpha=1
     ,family="gaussian")
10 # Optimal lambda can be retrieved,
 cv lasso$lambda.min
12
13 # Predictions can be calculated by specifying optimal lambda
 predictions = predict(cv_lasso, s = "lambda.min", newx =
     X test matrix, type = "response")
15
16 # If known model can also be trained on the optimal lambda directly
| lasso <- glmnet(X train matrix, data train$Fat, alpha = 1, family =
     "gaussian", lambda = optimal_lambda)
18
19 # Create a ridge model by setting alpha = 0.
ridge <- glmnet(X train matrix, data train$Fat, alpha = 0, family =</pre>
     gaussian")
```

# 2.5 Principal Component Analysis

```
res <- princomp(dataSC)

# View summary of the PCA
summary(res)

print(res$loadings)
print(res$scores)
```

# 3 Lab 1 Task 1: K-Nearest Neighbor Classification

The data file optdigits.csv contains information about normalized bitmaps of handwritten digits from a preprinted form from a total of 43 people. The data were first derived as 32x32 bitmaps

which were then divided into nonoverlapping blocks of 4x4, and the number of on pixels was counted in each block. This generated a resulting image of size 8x8 where each element is an integer in the range 0..16. Accordingly, each row in the data file is a sequence corresponding to an 8x8 matrix, and the last element shows the actual digit from 0 to 9.

- 1. Import the data into R and divide it into training, validation, and test sets (50%/25%/25%) by using the partitioning principle specified in the lecture slides. Use the training data to fit a 30-nearest neighbor classifier with the function kknn() and kernel="rectangular" from the kknn package, and estimate:
  - Confusion matrices for the training and test data (use table()).
  - Misclassification errors for the training and test data.

Comment on the quality of predictions for different digits and on the overall prediction quality.

- 2. Find any 2 cases of digit "8" in the training data which were easiest to classify and 3 cases that were hardest to classify (i.e., having the highest and lowest probabilities of the correct class). Reshape features for each of these cases as an 8x8 matrix and visualize the corresponding digits (e.g., using the heatmap() function with parameters Colv=NA and Rowv=NA). Comment on whether these cases seem to be hard or easy to recognize visually.
- 3. Fit K-nearest neighbor classifiers to the training data for different values of  $K=1,2,\ldots,30$ , and plot the dependence of the training and validation misclassification errors on the value of K (in the same plot). How does the model complexity change when K increases, and how does it affect the training and validation errors? Report the optimal K according to this plot. Finally, estimate the test error for the model having the optimal K, compare it with the training and validation errors, and make necessary conclusions about the model quality.
- 4. Fit K-nearest neighbor classifiers to the training data for different values of  $K=1,2,\ldots,30$ , compute the error for the validation data as cross-entropy (when computing the log of probabilities, add a small constant within the log, e.g., 1e-15, to avoid numerical problems), and plot the dependence of the validation error on the value of K. What is the optimal K value here? Assuming that the response has a multinomial distribution, why might the cross-entropy be a more suitable choice of error function than the misclassification error for this problem?

```
|18| id1 = setdiff(1:n, id)
 set.seed(12345)
20 | id2 = sample(id1, floor(n * 0.25))
21 data_validation = data_frame[id2, ]
23 id3 = setdiff(id1, id2)
24
 data_test = data_frame[id3, ]
25
 26
27
 # Performs k-nearest neighbor classification of a test set using a
     training set.
model test = kknn(formula = label ~ ., train = data train, test =
     data_test, k = 30, kernel = "rectangular")
  model_train = kknn(formula = label ~ ., train = data_train, test =
     data_train, k = 30, kernel = "rectangular")
31
 # Generate predictions for test and training data
predictions_test = predict(model_test)
34 predictions_train = predict(model_train)
35
 # Generate confusion matrix (classification) comparing predictions
     with true data.
gr conf_matrix_test = table(predictions_test, data_test$label)
conf_matrix_train = table(predictions_train, data_train$label)
 conf matrix test
40 conf matrix train
41
42 # Function to calculate the misclassification rate
43 # X = true labels, X1 = predictions
44 missclass=function(X,X1){
   n=length(X)
45
    return(1-sum(diag(table(X,X1)))/n)
47 }
48 missclass(predictions_test, data_test$label)
49 missclass(predictions_train, data_train$label)
51
  # predictions_test
                        0
                             1
                                 2
                                      3
                                          4
                                              5
                                                       7
                                                           8
 # 0
       77
            0
                 0
                     0
                         0
                              0
                                  0
                                       0
                                           0
                                               0
52
 # 1
        0
           81
                 0
                     0
                         0
                                  0
                                       0
                                           7
                              1
                                               1
53
 # 2
            2
        0
                98
                     0
                         0
                                  0
                                       0
                              1
                                           0
                                               1
 # 3
        0
             0
                 0
                   107
                          0
                              0
                                  0
55
                                               1
 # 4
            0
                 0
                     0
                         94
                              0
                                  0
                                       0
                                           0
                                               0
56
        1
 # 5
                     2
        0
            0
                 0
                         0
                             93
                                  0
                                       0
                                           0
                                               0
57
 # 6
            0
                     0
                          2
                              2
                                 90
                                       0
58
        0
                 0
                                           0
                                               0
 #
   7
        0
            0
                 0
                     0
                          6
                              1
                                  0
                                    111
                                           0
                                               1
59
 # 8
        0
            0
                 3
                     1
                          2
                              0
                                  0
                                       0
                                          70
                                               0
60
 # 9
                          5
        0
             3
                     1
                              5
                                  0
                                       0
                                           0
                                              85
61
                 0
63
 # predictions_train
                                  2
                                       3
                                           4
                                               5
                                                        7
                              1
                                                    6
                                                            8
                          0
64
65 # 0 202
            0
                              0
                                  0
                                       0
                                           0
                 0
                     0
                          1
                                               1
                                  2
66 # 1
        0 179
                 1
                     0
                          3
                              0
                                       3
                                          10
```

```
67 # 2
           11 190
                    0
                         0
                             0
                                 0
                                      0
                                          0
                                              0
        0
                 0 185
  # 3
        0
            0
                         0
                             1
                                 \cap
                                      0
                                          2
                                              5
                     0 159
                                              2
  # 4
        0
            0
                 0
                             0
                                 0
                                      0
                         0 171
  # 5
                 0
                     1
                                 0
70
  # 6
                     0
                         0
                               190
                                      0
                                              0
        0
            0
                 0
                             0
                                          2
71
72 # 7
                         7
                                              3
        0
            1
                 1
                     1
                             1
                                 0 178
                                          0
73
  # 8
        0
            1
                 0
                     0
                         1
                             0
                                 0
                                      1 188
                                              3
  # 9
            3
                 0
                     1
                         4
                             8
                                 0
                                      0
                                          2 183
74
75
76
  # > missclass(predictions test, data test$label)
77
78 # [1] 0.05329154
79 # > missclass(predictions train, data train$label)
  # [1] 0.04500262
81
82 # The overall quality of the models predictions is quite good, which
     can be seen
83 # above. For the test and training data set the misclassification
     rates are only
_{84} # 0.05329154 and 0.04500262 respectively. When checking the confusion
     matrix for
85 # the training dataset there are some values that stand out. These
     are the
86 # numbers 1, 4, 5, 8 and 9. These were wrongly classified the most by
     the model.
  # This is probably because some handwritten numbers look the same
     when using a
88 # low resolution image.
  91
92 # Get the probability matrix from the trained model
93 prob matrix = model train$prob
94
95 # Identify rows with the label "8" in the training dataset
96 label_8_indices = which(data_train$label == "8")
98
  # Extract probabilities of the true label "8" for these rows
  label_8_prob = prob_matrix[label_8_indices, "8"]
99
100
  # Sort the indices by descending probabilities
101
  sorted indices = order(label 8 prob, decreasing = TRUE)
102
103
  # Select the indices of the two easiest and three hardest
104
     classifications
  easiest_to_classify = label_8_indices[sorted_indices[1:2]]
105
  hardest_to_classify =
     label 8 indices[sorted indices[(length(sorted indices)-2):length(sorted indic
107
# Reshape to 8x8 matrices to visualize the digit with heat map
109 for (i in 1:2) {
    easiest case = matrix(as.numeric(data train[easiest to classify[i],
       1:64]), nrow = 8, ncol = 8, byrow=TRUE)
```

```
heatmap(easiest case, Rowv = NA, Colv = NA, main = paste("Easiest
111
       Case", i))
  }
112
  for (i in 1:3) {
113
    hardest_case = matrix(as.numeric(data_train[hardest_to_classify[i],
114
       1:64]), nrow = 8, ncol = 8, byrow=TRUE)
    heatmap(hardest case, Rowv = NA, Colv = NA, main = paste("Hardest
       Case", i))
116 }
117
  # The easiest two classifications are simple to recognize while the
118
     three hardest are almost impossible.
  120
121
  # Decide how many data points from neighboring data you want to
122
     consider when making predictions.
123 \min_{k} = 1
_{124} \mid \max_{k} \mid k = 30
|k_values = min_k:max_k
126 missclass_vector = c()
  missclass_vector_validation = c()
127
128
  for (k_value in k_values) {
129
    model_train = kknn(formula = label ~ ., train = data_train, test =
130
       data_train, k = k_value, kernel = "rectangular")
    model_validation = kknn(formula = label ~ ., train = data_train,
       test = data_validation, k = k_value, kernel = "rectangular")
    predict train = predict(model train)
    predict validation = predict(model validation)
134
135
    prob validation = model validation$prob
137
    missclass_value = missclass(predict_train, data_train$label)
138
    missclass_value_validation = missclass(predict_validation,
139
       data_validation$label)
140
    missclass_vector = append(missclass_vector, missclass_value)
141
    missclass_vector_validation = append(missclass_vector_validation,
142
       missclass value validation)
143
144
  plot(k values, missclass vector, col = "red", xlab = "k (Number of
145
     neighbors)", ylab = "Misclassification rate", main =
     "misclassification rate vs k value", ylim = c(0, 0.05))
  points(k_values, missclass_vector_validation, col = "blue")
147
  missclass vector validation[which.min(missclass vector validation)]
149
  legend("bottomright", legend = c("Training", "Validation"), col =
150
     c("red", "blue"), lty = 1)
151
```

```
452 # When the K values are small the complexity of the model is high and
     there is a big risk for overfitting. While when the K values are
     big the complexity of the model decreases and there is risk for
     underfitting. By looking at the figure we can see that for low K
     values the red and blue model is better while for big K values
     both models is almost equally as bad.
  # The most optimal K value for the blue model is K = 3 then the
154
     misclassification value is 0.02513089 and for the red model is K =
     1 and misclassification value is 0.00. This is because the red
     model is trained and tested on the same dataset. This makes the
     model look very good for low K values but in reality its
     overfitted. The blue model has worse misclassification value but
     its not trained and tested on the same dataset so it shows a more
     realistic rate for different K values.
155
  LOG CONSTANT = 1e-15
158
  cross_entropy = c()
159
  for (k value in k values) {
161
    model_train = kknn(formula = label ~ ., train = data_train, test =
162
       data_train, k = k_value, kernel = "rectangular")
    model validation = kknn(formula = label ~ ., train = data train,
163
       test = data validation, k = k value, kernel = "rectangular")
164
    prob_validation = model_validation$prob
165
166
    # Cross entropy
167
    cross entropy k = 0
168
    for (i in 1:nrow(data validation)) {
169
      true_label = data_validation$label[i] # Gets label for each row
      true prob = prob validation[i, as.numeric(true label)] # Get prob
171
         for current row and true label
      cross_entropy_k = cross_entropy_k - log(true_prob + LOG_CONSTANT)
         # calc cross entropy
    cross_entropy = c(cross_entropy, cross_entropy_k /
174
       nrow(data_validation))
  }
175
176
  plot(k_values, cross_entropy, xlab = "k (Number of neighbors)", ylab
     = "Cross-entropy loss", main = "Cross-Entropy Loss vs. k", ylim =
     c(0, 1.0)
  which.min(cross entropy)
178
179
  # From the figure we can see that the optimal K value is 6, because
     it has the lowest loss. The cross entropy is a more suitable
     choice for the error function because it evaluates the quality of
     the predicted probabilities. Compared to the misclassification
     which only cares if the predicted digit was correct or not, and
     not how high probability that the true digit actually has.
```

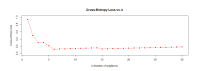


Figure 1: cross entropy vs k



Figure 2: easiest case 1



Figure 3: easiest case 2



Figure 4: hardest case 1



Figure 5: hardest case 2



Figure 6: hardest case 3

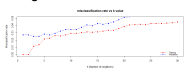


Figure 7: misclassfication vs k

# 4 Lab 1 Task 2: Linear Regression and Ridge Regression

The data file parkinson.csv is composed of a range of biomedical voice measurements from 42 people with early-stage Parkinson's disease recruited to a six-month trial of a telemonitoring device for remote symptom progression monitoring. The purpose is to predict Parkinson's disease symptom score (motor UPDRS) from the following voice characteristics:

- **Jitter**: (%), Jitter(Abs), Jitter:RAP, Jitter:PPQ5, Jitter:DDP Several measures of variation in fundamental frequency.
- **Shimmer**: Shimmer(dB), Shimmer:APQ3, Shimmer:APQ5, Shimmer:APQ11, Shimmer:DDA Several measures of variation in amplitude.
- NHR, HNR Two measures of the ratio of noise to tonal components in the voice.
- RPDE A nonlinear dynamical complexity measure.
- **DFA** Signal fractal scaling exponent.
- PPE A nonlinear measure of fundamental frequency variation.
- 1. Divide the data into training and test sets (60/40) and scale it appropriately. In the following steps, assume that motor\_UPDRS is normally distributed and is a function of the voice characteristics. Since the data are scaled, no intercept is needed in the modeling.
- 2. Compute a linear regression model from the training data, estimate training and test MSE, and comment on which variables contribute significantly to the model.
- 3. Implement the following functions using basic R commands only (no external packages):
  - a) **LogLikelihood**: A function for a given parameter vector  $\theta$  and dispersion  $\sigma$  computes the log-likelihood function  $\log P(T|\theta,\sigma)$  for the stated model and the training data.
  - b) **Ridge**: A function that, for a given vector  $\theta$ , scalar  $\sigma$ , and scalar  $\lambda$ , adds a Ridge penalty  $\lambda \|\theta\|^2$  to the minus log-likelihood.

- c) **RidgeOpt**: A function that depends oon the scalar  $\lambda$ , uses function 3b and function optim() with method="BFGS" to find the optimal  $\theta$  and  $\sigma$  for a given  $\lambda$ .
- d) **DF (Degrees of Freedom)**: function that for given scalar  $\lambda$  computes the degrees of freedom of the ridge model based on the training data.
- 4. Using the Ridge optimization function, compute the optimal  $\theta$  parameters for  $\lambda=1,\,\lambda=100$ , and  $\lambda=1000$ . Use the estimated parameters to predict motor\_UPDRS values for training and test data and report the training and test MSE values. Determine which penalty parameter is most appropriate among the selected ones. Compute and compare the degrees of freedom of these models and make appropriate conclusions.

```
_____ TASK 1 _____
 data <- read.csv('parkinsons.csv')</pre>
 labels <- colnames(data)</pre>
 parameters<-c("Jitter...", "Jitter.Abs.", "Jitter.RAP",</pre>
     "Jitter.PPQ5", "Jitter.DDP",
               "Shimmer", "Shimmer.dB.", "Shimmer.APQ3",
                  "Shimmer.APQ5", "Shimmer.APQ11",
               "Shimmer.DDA", "NHR", "HNR", "RPDE", "DFA", "PPE")
 parameters_y<-c(parameters, "motor_UPDRS")</pre>
9
 n=dim(data)[1]
11
12 set.seed (12345)
13 id=sample(1:n, floor(n*0.6))
14 train=data[id,]
15
16 valid=data[-id,]
17
18 library (caret)
19 scaler=preProcess(train)
20 trainS=predict(scaler, train)
21 validS=predict(scaler, valid)
trainS = trainS[, colnames(trainS) %in% parameters_y]
validS = validS[, colnames(validS) %in% parameters y]
25
 trainS_without_y_matrix = as.matrix(subset(trainS,
     select=-motor_UPDRS))
 validS_without_y_matrix = as.matrix(subset(validS,
     select=-motor UPDRS))
28
    TASK 2
29
30
 y = trainS$motor_UPDRS
32 y_validS = validS$motor_UPDRS
34 fit=lm(y ~ . - motor UPDRS, data=trainS)
36 coeff = coefficients(fit)[-1] # model coefficients, intercept not
     needed since the data is scaled
37 confint(fit, level=0.95) # CIs for model parameters
```

```
38 fitted(fit) # predicted values
39 residuals trainS = residuals(fit) # residuals
40
41
42
 predicted_validS = predict(fit, newdata=validS)
 residuals validS <- validS$motor_UPDRS - predicted_validS</pre>
45
46 # ANSWER:
mean(residuals trainS^2) # MSE for trainS = 0.8785431
48 mean(residuals validS^2) # MSE for validS = 0.9354477
49
50 summary (fit)
51
 # ANSWER:
52
_{53}| # summary(fit) will print some stuff. The three stars next to some
     variables mean that there is a high correlation
54 # between them and the target variable motor_UPDRS.
55 # Answer: Jitter.Abs., Shimmer.APQ5, Shimmer.APQ11, NHR, HNR, DFA and
     PPE
56
57
           TASK 3
Loglikelihood <- function(theta, sigma) # linear regression gives
     normal distribution
59 {
    - nrow(trainS) / 2 * log(2 * pi) - nrow(trainS) / 2 * log(sigma^2)
       -1/(2*sigma^2) * sum((y - trainS without y matrix %*% theta)^2)
 }
61
Ridge <- function(theta, lambda, sigma)
64
    -Loglikelihood(theta, sigma) + lambda * sum(theta^2)
65
66 }
67
68 RidgeOpt <- function(lambda)
69
    optim(coeff, fn = Ridge, lambda = lambda, sigma = 1, method =
       "BFGS")
 }
71
73 DF <- function(lambda)
74 {
    X = trainS_without_y_matrix
75
    sum(diag(X \%*\% solve(t(X) \%*\% X + lambda * diag(ncol(X))) \%*\% t(X)))
76
77
 # ____ TASK 4 _____
78
79
80 lambda_1 = RidgeOpt(1)$par
81 lambda_100 = RidgeOpt(100)$par
82 lambda 1000 = RidgeOpt(1000) $par
83
 #
84
85
```

```
86 # Use the estimated parameters to predict the motor_UPDRS values for
     training data
  predict trainS motor UPDRS lambda 1 = trainS without y matrix %*%
     lambda 1
  predict_trainS_motor_UPDRS_lambda_100 = trainS_without_y_matrix %*%
     lambda 100
  predict trainS motor UPDRS lambda 1000 = trainS without y matrix %*%
     lambda 1000
90
  # calculate the MSE
91
92 MSE trainS lambda 1 = mean((predict trainS motor UPDRS lambda 1 -
     y)^2)
93 MSE trainS lambda 100 = mean((predict trainS motor UPDRS lambda 100 -
     y)^2)
  MSE trainS lambda 1000 = mean((predict trainS motor UPDRS lambda 1000
94
     -y)^2
95
96
97
  # Use the estimated parameters to predict the motor UPDRS values for
98
     test data
  predict validS motor UPDRS lambda 1 = validS without y matrix %*%
     lambda 1
  predict_validS_motor_UPDRS_lambda_100 = validS_without_y_matrix %*%
     lambda 100
  predict validS motor UPDRS lambda 1000 = validS without y matrix %*%
101
     lambda 1000
102
  # Calculate the MSE
103
  MSE validS lambda 1 = mean((predict validS motor UPDRS lambda 1 -
104
     y_validS)^2)
MSE validS lambda 100 = mean((predict validS motor UPDRS lambda 100 -
     y validS)^2)
  MSE validS lambda 1000 = mean((predict validS motor UPDRS lambda 1000
106
     - y_validS)^2)
107
DF_lambda_1 = DF(1)
     lambda_100 = DF(100)
  \mathsf{DF}
109
  DF_lambda_1000 = DF(1000)
110
111
112
    -----
113
114 # ANSWER:
MSE trainS lambda 1 # 0.8786332
116 MSE trainS lambda 100 # 0.8850982
MSE_trainS_lambda_1000 # 0.9232605
118
119 MSE_validS_lambda_1 # 0.9349501
120 MSE validS lambda 100 # 0.9324709
MSE validS lambda 1000 # 0.9554836
122
123 DF lambda 1 # 13.86074
124 DF lambda 100 # 9.924887
```

```
DF_lambda_1000 # 5.643925

# We see that MSE_validS_lambda_100 is the lowest which
# means that the lambda = 100 is the most suitable choice. In this case it can be

# seen that a higher degree of freedom is not always the best choice since it

# can lead to overfitting.
```

# 5 Lab 1 Task 3: Logistic Regression and Basis Function Expansion

The data file pima-indians-diabetes.csv contains information about the onset of diabetes within 5 years in Pima Indians given medical details. The variables are (in the same order as in the dataset):

- 1. Number of times pregnant.
- 2. Plasma glucose concentration 2 hours into an oral glucose tolerance test.
- 3. Diastolic blood pressure (mm Hg).
- 4. Triceps skinfold thickness (mm).
- 5. 2-Hour serum insulin ( $\mu$  U/ml).
- 6. Body mass index (weight in kg/(height in m)<sup>2</sup>).
- 7. Diabetes pedigree function.
- 8. Age (years).
- 9. Diabetes (0 = no or 1 = yes).
- Make a scatterplot showing Plasma glucose concentration on Age where observations are colored by Diabetes levels. Do you think that Diabetes is easy to classify by a standard logistic regression model that uses these two variables as features? Motivate your answer.
- 2. Train a logistic regression model with y= Diabetes as target,  $x_1=$  Plasma glucose concentration, and  $x_2=$  Age as features, and make a prediction for all observations by using r=0.5 as the classification threshold.
  - Report the probabilistic equation of the estimated model (i.e., how the target depends on the features and the estimated model parameters probabilistically).
  - Compute the training misclassification error and make a scatterplot of the same kind as in step 1 but showing the predicted values of Diabetes as a color instead.
  - Comment on the quality of the classification by using these results.
- 3. Use the model estimated in step 2 to:
  - a) Report the equation of the decision boundary between the two classes.
  - b) Add a curve showing this boundary to the scatterplot in step 2.

Comment whether the decision boundary seems to capture the data distribution well.

4. Make the same kind of plots as in step 2 but use thresholds r=0.2 and r=0.8. By using these plots, comment on what happens with the prediction when the r value changes.

5. Perform a basis function expansion trick by computing new features:

$$z_1 = x_1^4$$
,  $z_2 = x_1^3 x_2$ ,  $z_3 = x_1^2 x_2^2$ ,  $z_4 = x_1 x_2^3$ ,  $z_5 = x_2^4$ 

Add them to the dataset and compute a logistic regression model with y as target and  $x_1, x_2, z_1, \ldots, z_5$  as features.

- Create a scatterplot of the same kind as in step 2 for this model.
- · Compute the training misclassification rate.
- What can you say about the quality of this model compared to the previous logistic regression model?
- How has the basis expansion trick affected the shape of the decision boundary and the prediction accuracy?

```
library(ggplot2)
 # Set seed to 12345
 set.seed (12345)
6 # Import data
 data <- read.csv("../pima-indians-diabetes.csv", header = FALSE)</pre>
 # Add headers to data
  colnames(data) <- c("times_pregnant",</pre>
                        "plasma",
11
                        "blood_pressure",
12
                        "skinfold_thickness",
13
                        "insulin",
14
                        "bmi",
15
                        "diabetes_pedigree",
16
                        "age",
17
                        "diabetes")
18
19
     TASK 1
20
21
 # Plot data where age is the x axis, plasma is the y axis and color
22
     is based
23 # on diabetes value
 ggplot(data, aes(x = age, y = plasma, color = diabetes)) +
24
    geom_point() + scale_color_gradient(low = "green", high = "red") +
25
    labs(title = "Task 1",
26
         x = "Age",
27
         y = "Plasma Glucose Concentration",
28
         color = "Diabetes") +
29
    theme minimal()
30
 \# By observing the result of the task, we deem that it is difficult to classify diabetes by a standard logistic regression model that
     uses these two variables as features. When looking at the graph it
     becomes quite clear that there isn't a strong enough connection
     between plasma levels and age to determine if a person has
     diabetes or not. There are for example several people between age
     60 and 70 with a plasma level above 150 whom do not have diabetes.
```

```
Meanwhile there are also young people below the age of 30 with a
     plasma level below 100 whom have diabetes. In conclusion, we
     determine that diabetes is hard to classify using these two
     variable. However, we have observed that diabetes seems more
     common for older ages and higher plasma levels.
33
   TASK 2
35
model = glm(diabetes~age + plasma, data=data, family = binomial)
 predictions <- predict(model, type = "response")</pre>
#Initialize all different R values to be used for future plotting.
40 r1 <- 0.5
42
43 #Calculate binary predictions based on all three r values.
44 binary_predictions_1 <- ifelse(predictions >= r1, 1, 0)
46 #Function to calculate missclassification error.
47 missclass = function(X,X1){
  n=length(X)
   return(1-sum(diag(table(X,X1)))/n)
49
50 }
51
52 #Calculate missclassification error.
misclassification error = missclass(binary predictions 1,
     data$diabetes) # 0.2630208
54
55 # ANSWER:
 # the family variable is set to binomial because the response
     variable follows a binomial distribution, since it is a binary
     variable.
 #Plot binary predictions and the decision boundary where r = 0.5.
ggplot(data, aes(x = age, y = plasma, color = binary_predictions_1)) +
   geom_point() + scale_color_gradient(low = "green", high = "red") +
60
    labs(title = "Diabetes Predictions r = 0.5",
61
         x = "Age",
62
         y = "Plasma Glucose Concentration",
63
         color = "Diabetes") +
64
   theme minimal()
65
66
67
68 # ANSWER:
 # Probabilistic model and figure is under the code.
70
71 # _____ TASK 3 _____
72
73 #Obtain coefficients from model.
74 b0 <- coef(model)[1]
75 b1 <- coef(model)[2]
76 b2 <- coef(model)[3]
```

```
78 #Arrange X1 in order from smallest to largest.
79 x1 <- seq(min(data$age), max(data$age))
81 # Calculate x2 values for the decision boundary.
|a_2| + b_0 + b_1 * x_1 + b_2 * x_2 = 0 => x_2
x^2 < -(b^0 + b^1 * x^1) / b^2
  #Create the decision boundary line based on x1 and x2 values
85
decision boundary \leftarrow data.frame(x1 = x1, x2 = x2)
_{88} #Plot binary predictions and the decision boundary where r = 0.5.
  ggplot(data, aes(x = age, y = plasma, color = binary_predictions_1)) +
89
    geom_point() + scale_color_gradient(low = "green", high = "red") +
90
    labs(title = "Diabetes Predictions r = 0.5 + decision boundary",
91
         x = "Age",
92
         y = "Plasma Glucose Concentration",
93
          color = "Diabetes") +
94
    geom_line(data = decision_boundary, aes(x = x1, y = x2), color =
       "blue", linewidth = 1) + # Add decision boundary
    theme minimal()
96
97
  # ANSWER:
  # It seems to catch the data distribution well since people predicted
     to suffer from diabetes or above it or on it and people predicted
     not to suffer from diabetes are on it or below it. The decision
     boundary can be observed in figure.
100
    TASK 4
101
102
  r2 <- 0.2
103
  r3 <- 0.8
104
105
binary predictions 2 <- ifelse(predictions >= r2, 1, 0)
  binary predictions 3<- ifelse(predictions >= r3, 1, 0)
107
108
  ggplot(data, aes(x = age, y = plasma, color = binary_predictions_2)) +
109
    geom_point() + scale_color_gradient(low = "green", high = "red") +
110
    labs(title = "Diabetes Predictions r = 0.2",
111
         x = "Age",
112
         y = "Plasma Glucose Concentration",
113
          color = "Diabetes") +
    theme minimal()
115
116
  ggplot(data, aes(x = age, y = plasma, color = binary_predictions_3)) +
117
    geom point() + scale color gradient(low = "green", high = "red") +
118
    labs(title = "Diabetes Predictions, r = 0.8",
119
         x = "Age",
120
         y = "Plasma Glucose Concentration",
121
         color = "Diabetes") +
122
    theme_minimal()
123
124
125 # ANSWER:
126 # By observing these plots it becomes evident that r decides how
```

```
certain our model has to bee in order to decide on whether a
     person suffers from diabetes or not. In other words, a person is
     predicted to suffer from diabetes if the predicted probability of
     the model for that person is greater than r. This results in a
     greater amount of people predicted to suffer from diabetes by the
     model for lower r values.
127
      TASK 5
128
129
data$z1 = data$plasma^4
data$z2 = data$plasma^3 * data$age
data$z3 = data$plasma^2 * data$age^2
  data$z4 = data$plasma * data$age^3
  data$z5 = data$age^4
134
135
  model2 \leftarrow glm(diabetes \sim plasma + age + z1 + z2 + z3 + z4 + z5,
136
     family = binomial, data = data)
predictions <- predict(model2, newdata = data, type = "response")
139 r <- 0.5
140 binary predictions <- ifelse(predictions >= r, 1, 0)
  ggplot(data, aes(x = age, y = plasma, color = binary_predictions)) +
142
    geom_point() + scale_color_gradient(low = "green", high = "red") +
143
    labs(title = "Diabetes Predictions r = 0.5",
144
         x = "Age",
145
         y = "Plasma Glucose Concentration",
146
         color = "Diabetes") +
147
    theme minimal()
148
149
  misclassification error <- missclass(data$diabetes,
150
     binary_predictions)
  print(misclassification error) # 0.2447917
152
153 # ANSWER:
154 # This is a small improvement compared to the missclassification
     error observed in task 2, but one could argue that the two answers
     are in a similar range and neither one is particularly good. even
     though this new model has a slightly smaller missclassification
     error than the previous one, it has one large issue which is that
     the prediction rate for diabetes decreases for higher ages. This
     seems like an unwanted behaviour since diabetes is usually more
     common in higher ages, but this prediction model basically
     predicts that people above a certain age (around 65) never have
     diabetes.
```

Probabilistic equation of the model in task 2:

P(diabetes| age, plasma) = 
$$\frac{1}{1 + e^{-(\beta_0 + \beta_1 * age + \beta_2 * plasma)}}$$

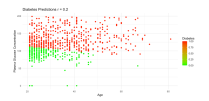




Figure 8: diabetes pred 0.2

Figure 9: diabetes pred 0.5

Figure 10: diabetes pred 0.8



Figure 11: diabetes pred final

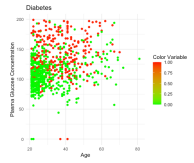


Figure 12: task 1 scatter plot

# 6 Lab 2 Task 1: Tecator Dataset Analysis

The file tecator.csv contains the results of a study aimed at investigating whether a near-infrared absorbance spectrum can predict the fat content of meat samples. For each meat sample, the data consists of a 100-channel spectrum of absorbance records and the levels of moisture (water), fat, and protein. The absorbance is  $-\log_{10}$  of the transmittance measured by the spectrometer. The moisture, fat, and protein levels are determined by analytic chemistry. Divide the data randomly into train and test sets (50/50) using the codes from the lectures.

- 1. Assume that Fat can be modeled as a linear regression in which absorbance characteristics (Channels) are used as features.
  - Report the underlying probabilistic model.
  - Fit the linear regression to the training data and estimate the training and test errors.
  - Comment on the quality of fit and prediction and therefore on the quality of the model.
- 2. Assume now that Fat can be modeled as a LASSO regression in which all Channels are used as features.
  - Report the cost function that should be optimized in this scenario.
- 3. Fit the LASSO regression model to the training data.
  - · Present a plot illustrating how the regression coefficients depend on the log of the penalty factor (log  $\lambda$ ).
  - Interpret this plot. What value of the penalty factor can be chosen if we want to select a model with only three features?
- 4. Repeat step 3 but fit Ridge regression instead of LASSO and compare the plots from steps 3 and 4.
  - · Comment on the conclusions.
- 5. Use cross-validation with the default number of folds to compute the optimal LASSO model.

- Present a plot showing the dependence of the CV score on  $\log \lambda$  and comment on how the CV score changes with  $\log \lambda$ .
- Report the optimal  $\lambda$  and how many variables were chosen in this model.
- Does the information displayed in the plot suggest that the optimal  $\lambda$  value results in a statistically significantly better prediction than  $\log \lambda = -4$ ?
- Finally, create a scatter plot of the original test versus predicted test values for the model corresponding to the optimal  $\lambda$  and comment on whether the model predictions are good.

```
library(glmnet)
 data=read.csv("../tecator.csv", header = TRUE)
 # Set random seed so lab result is predictable
 set.seed(12345)
6
 # Create variable without protein and moisture columns.
 channel data <-data.frame(data[c(2:102)])</pre>
9
10
_{11} | # Split the dataset into training (50%), and test (50%).
                                      # Total nr of rows in dataset
12 n = dim(channel_data)[1]
|13| id = sample(1:n, floor(n * 0.5))
                                      # Randomly selects 50 % rows for
     training
id1 = setdiff(1:n, id)
                                   # Remaining rows
data train = channel data[id,] # Assign rows to training set
data test = channel data[id1, ] # Set remaining rows as test data
 # Task 1
19
20 # Create a linear regression model where Fat is the target
21 # . indicates all other columns are prediction variables
22 model=lm(Fat~., data=data_train)
23 model
24
predict train <- predict(model, data train) # make predictions for
     training data
 predict test <- predict(model,data test) # make predictions for test</pre>
     data
27
28 # calculate difference between predictions and actual values for both
     data sets
29 train_diff <- (data_train$Fat - predict_train)</pre>
test_diff <- (data_test$Fat - predict_test)</pre>
32 # Caclulate Squared Error for both data sets.
33 sum(train diff^2)/dim(data train)[1]
34 sum(test_diff^2)/dim(data_test)[1]
36 # Model is overfitted, probably due to 100 different variables.
37 # n is barley larger than p
38 # mse test = 722.4294
39 # mse_train = 0.005709117
```

```
40
41 # Task 3
42 #Create a matrix without the fat column for the GLMNET model.
43 X_train_matrix <- as.matrix(data_train[c(1:100)])
45 # Create the lasso model by setting alpha = 1.
 lasso <- glmnet(X train matrix, data train$Fat, alpha = 1, family =</pre>
     "gaussian")
47
48 # Look how many features the model has for the lambda = 0.818
_{49} coef(lasso, s = 0.818)
50 plot(lasso, xvar = "lambda")
51
 # By looking at the graph, a model with only three coefficients is
     approximately
_{53} | # log lambda = -0,2 which results in lambda = 0.8187
54
55 # Task 4
56 #Create the ridge model by setting alpha = 0.
57 ridge <- glmnet(X_train_matrix, data_train$Fat, alpha = 0, family =</pre>
     "gaussian")
58 plot(ridge,xvar = "lambda")
59
60 # task 5
61 # Train covariane lasso model
62 cv lasso=cv.glmnet(X train matrix, data train$Faat, alpha=1
     ,family="gaussian")
63 cv lasso
64 cv lasso$lambda.min
65 plot (cv_lasso)
67 coef(cv lasso, s = cv lasso$lambda.min)
68 #8 coefficients are chosen in the model, intercept is NOT included
70 X_test_matrix <- as.matrix(data_test[c(1:100)])</pre>
predictions = predict(cv_lasso, s = "lambda.min", newx =
     X_test_matrix, type = "response")
72
 sum((predictions - mean(data_test$Fat))^2)/sum((data_test$Fat -
     mean(data_test$Fat))^2)
 sum((predictions - data_test$Fat)^2)
75
plot(data_test$Fat, col = "black", pch = 16, ylab = "Fat")
 points(predictions, col="red", pch = 17)
77
| legend("topleft", inset = c(0, -0.24), 
         xpd = TRUE, legend = c("Test data", "Predictions"),
79
         col = c("black", "red"), pch = c(16, 17)
80
81
 )
```

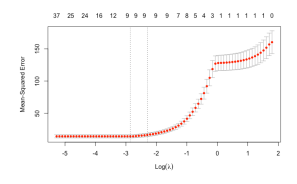
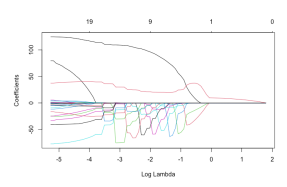


Figure 13: cv lasso

Figure 14: fat scatter



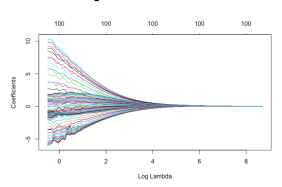


Figure 15: lasso regression

Figure 16: ridge regression plot

# 7 Lab 2 Task 2: Decision Trees and Logistic Regression for Bank Marketing

The data file <code>bank-full.csv</code> is related to direct marketing campaigns of a Portuguese banking institution. The marketing campaigns were based on phone calls. Often, more than one contact with the same client was required to assess if the product (bank term deposit) would be ('yes') or not ('no') subscribed.

# Input Variables

# · Bank client data:

- 1. Age (numeric).
- Job: type of job (categorical: 'admin.', 'blue-collar', 'entrepreneur', 'housemaid',
   'management', 'retired', 'self-employed', 'services', 'student', 'technician',
   'unemployed', 'unknown').
- 3. Marital: marital status (categorical: 'divorced', 'married', 'single', 'unknown'; note: 'divorced' means divorced or widowed).
- 4. Education (categorical: 'basic.4y', 'basic.6y', 'basic.9y', 'high.school', 'illiterate', 'professional.course', 'university.degree', 'unknown').
- 5. Default: has credit in default? (categorical: 'no', 'yes', 'unknown').

- 6. Housing: has housing loan? (categorical: 'no', 'yes', 'unknown').
- 7. Loan: has personal loan? (categorical: 'no', 'yes', 'unknown').

### Related to the last contact of the current campaign:

- 1. Contact: contact communication type (categorical: 'cellular', 'telephone').
- 2. Month: last contact month of the year (categorical: 'jan', 'feb', ..., 'nov', 'dec').
- 3. Day of week: last contact day of the week (categorical: 'mon', 'tue', 'wed', 'thu', 'fri').
- 4. Duration: last contact duration, in seconds (numeric). **Important note:** This attribute highly affects the output target (e.g., if duration=0, then y='no'). Yet, the duration is not known before a call is performed. Also, after the end of the call, y is obviously known. Thus, this input should only be included for benchmark purposes and should be discarded if the intention is to have a realistic predictive model.

#### Other attributes:

- 1. Campaign: number of contacts performed during this campaign and for this client (numeric, includes the last contact).
- 2. Pdays: number of days that passed by after the client was last contacted from a previous campaign (numeric; 999 means client was not previously contacted).
- 3. Previous: number of contacts performed before this campaign and for this client (numeric).
- 4. Poutcome: outcome of the previous marketing campaign (categorical: 'failure', 'nonexistent', 'success').

# Output Variable (Target)

• y: Has the client subscribed a term deposit? (binary: 'yes', 'no').

### **Tasks**

- 1. Import the data to R, remove the variable  $\mathtt{duration}$ , and divide it into training/validation/test sets as 40/30/30. Use data partitioning code specified in Lecture 2a.
- 2. Fit decision trees to the training data by changing the default settings one by one (i.e., not simultaneously):
  - a) Decision tree with default settings.
  - b) Decision tree with smallest allowed node size equal to 7000.
  - c) Decision tree with minimum deviance equal to 0.0005.

Report the misclassification rates for the training and validation data. Which model is the best among these three? Report how changing the deviance and node size affected the size of the trees and explain why.

3. Use the training and validation sets to choose the optimal tree depth in model 2c. Study the trees up to 50 leaves. Present a graph of the dependence of deviances for the training and validation data on the number of leaves. Interpret this graph in terms of the bias-variance tradeoff. Report the optimal number of leaves and identify which variables seem to be most important for decision-making in this tree. Interpret the information provided by the tree structure.

- 4. Estimate the confusion matrix, accuracy, and F1 score for the test data by using the optimal model from step 3. Comment on whether the model has good predictive power and which of the measures (accuracy or F1 score) should be preferred here.
- 5. Perform a decision tree classification of the test data using the following loss matrix:

$$L = \begin{bmatrix} 0 & 5 \\ 1 & 0 \end{bmatrix}$$

Report the confusion matrix for the test data. Compare the results with the results from step 4 and discuss how the rates have changed and why.

6. Use the optimal tree and a logistic regression model to classify the test data using the following principle:

$$Y = \begin{cases} y_{pos} & \text{if } P(Y = y_{pos}|X) > \pi, \\ y_{neg} & \text{otherwise.} \end{cases}$$

where  $\pi = 0.05, 0.1, 0.15, \dots, 0.9, 0.95$ . Compute the TPR and FPR values for the two models and plot the corresponding ROC curves. What are your conclusions? Why might the precision-recall curve be a better option here?

```
library(tree)
 library(rpart)
 # Import bank marketing campaign csv file with str as factors
 data frame = read.csv2("../bank-full.csv", stringsAsFactors = TRUE)
10
 # Remove column "duration" using %in% operator
 data frame = data frame[ , !names(data frame) %in% c("duration")]
13
 # Split the dataset into training (50%), validation (25%), and test
    (25%) sets
_{15}|_{n} = dim(data frame)[1]
                                 # Total nr of rows in dataset
 set.seed(12345)
 id = sample(1:n, floor(n * 0.4))
                                 # Randomly selects 40 % rows for
    training
 data_train = data_frame[id, ]
                                 # Assign rows to training set
18
19
20 id1 = setdiff(1:n, id)
                                 # Remaining rows
21 set.seed (12345)
22 id2 = sample(id1, floor(n * 0.30)) # Randomly selects 30 % rows for
    validation
data validation = data frame[id2, ] # Assign rows to validation set
 id3 = setdiff(id1, id2)
                                  # Remaining rows
25
 data_test = data_frame[id3, ] # Assign rest of 30 % rows to
    testing
27
29 ####################### PART 2 #############################
```

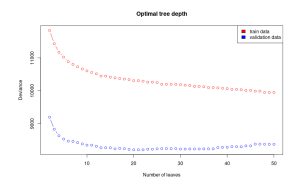
```
31 # Fit decision trees to training dataset
32 tree_a = tree(formula = y ~ ., data = data_train)
s3 tree_b = tree(formula = y ~ ., data = data_train, control =
    tree.control(nrow(data_train), minsize = 7000))
s4 tree_c = tree(formula = y ~ ., data = data_train, control =
    tree.control(nrow(data_train), mindev = 0.0005))
35
36 # Predictions on validation dataset
37 predictions a = predict(tree a, newdata = data validation, type =
predictions b = predict(tree b, newdata = data validation, type =
     "class")
 predictions c = predict(tree c, newdata = data validation, type =
    "class")
40
41 # Misclassification rates for the training dataset
summary(tree_a) # Misclassification error rate: 0.1048
43 summary(tree b) # Misclassification error rate: 0.1048
44 summary(tree_c) # Misclassification error rate: 0.09362
46 # Misclassification rates for the validation dataset
47 misclass = function(X, X1) {
 n = length(X)
48
   return(1 - sum(diag(table(X, X1))) / n)
49
misclass(predictions_a, data_validation$y) # Misclassification error
    rate: 0.1092679
misclass(predictions_b, data_validation$y) # Misclassification error
    rate: 0.1092679
misclass(predictions c, data validation$y) # Misclassification error
    rate: 0.1118484
55 #The best models are A and B, as they have low misclassification
    rates for both the training set and the validation set. The
    problem with Model C is that it has a good misclassification rate
    for the training set but a higher misclassification rate for the
    validation set, which is typical of overfitting.
57 #A larger minimum node size increases the model misclassification
    rate while a lower does not improve performance. Increasing the
    minsize value forces the tree to stop splitting when a node's
    number of obersvations falls below the threshold. This results in
    simpler and more general trees. Lowering the minsize can allow the
    tree to grow very deep, increasing the risk of overfitting. While
    high values the tree is constrained from splitting too much, which
    can lead to underfitting.
59 #A larger minimum deviance makes the model better for the validation
    set but worse for the training set. This is because a higher
    minimum deviance results in a smaller and simpler tree, reducing
    the risk of overfitting but increasing the risk of underfitting.
    Lowering the minimum deviance allows the tree to grow larger and
```

```
more complex, improving performance on the training set but
     increasing the risk of overfitting, which can lead to poor
     generalization and worse performance on the validation set.
 61
62
63
 # Init vector with length 50 to store score
 train_score = rep(0, 50)
64
valid score = rep(0, 50)
67 # Prune the C tree starting with at least 2 node leaves up to 50
# Calculate the deviance on both datasets and save to the score
    vector
69 for (leaves in 2:50) {
    # Prune Tree C to current "leaves" number
70
   pruned_tree = prune.tree(tree_c, best = leaves)
71
72
    # Predictions on validation dataset using pruned tree
73
   predictions_valid = predict(pruned_tree, newdata = data_validation,
74
      type = "tree")
75
    train score[leaves] = deviance(pruned tree)
76
    valid_score[leaves] = deviance(predictions_valid)
77
 }
78
79
 # Graph of the dependence of deviance for the datasets on the number
    of leaves
81 plot(2:50, train_score[2:50], type = "b", col = "red", ylim =
     c(min(valid_score[2:50]), max(train_score[2:50])),
      main = "Optimal tree depth", ylab = "Deviance", xlab = "Number
82
          of leaves")
83 points(2:50, valid_score[2:50], type = "b", col = "blue")
84 legend("topright", c("train data", "validation data"), fill =
    c("red", "blue"))
85
86 # Optimal number of leaves that minimize training deviance.
87 optimal_train = which.min(train_score[2:50])
88 optimal_validation = which.min(valid_score[2:50])
90 optimal_tree = prune.tree(tree_c, best = optimal_validation)
92 # Display optimal tree
93 plot(optimal tree)
94 text(optimal_tree, pretty=0)
 optimal tree
 summary(optimal tree)
96
_{98} # The optimal number of leaves is 47 for the training dataset and 21
    for the validation dataset. The bias-variance tradeoff explains
    how a model's complexity influences prediction accuracy and
    generalization to unseen data. A tree with many leaves (greater
    depth) has low bias but high variance, potentially leading to
     overfitting. A tree with fewer leaves has low variance but high
```

```
bias, which can result in underfitting. As model complexity
     increases, variance rises while bias decreases. In decision trees,
     the number of leaves directly impacts this balance as it defines
     the complexity of the tree. The key is to find the optimum where
     there is a balance between bias and variance so the error is
     minimized, see Figure \ref{fig:task_3_graph}.
  # The variables which are the most important is poutcome which is the
100
     root node but also month, contact and pdays which appear
     frequently in splits in the tree, see Figure
     \ref{fig:task3 optimal tree}.
101
  102
  # Predictions on the test dataset using optimal tree
104
  predications_test = predict(optimal_tree, newdata = data_test, type =
105
     "class")
  # Creating confusion matrix
107
  conf_matrix = table(data_test$y, predications_test)
108
  conf matrix
  # Calculates the accuracy and f1 score from confusion matrix
111
  accuracy_and_f1 = function(conf_matrix) {
112
    TP = conf_matrix["yes", "yes"] # True positives
113
    TN = conf_matrix["no", "no"] # True negatives
114
    FP = conf matrix["no", "yes"] # False positives
115
    FN = conf_matrix["yes", "no"] # False negatives
116
117
    precision = TP / (TP + FP)
118
    recall = TP / (TP + FN)
119
    accuracy = (TN + TP) / (TN + FP + TP + FN)
120
    f1 score = 2 * (precision * recall) / (precision + recall)
    return(c(accuracy = accuracy, f1_score = f1_score))
124
125
126
  # Print result
result = accuracy_and_f1(conf_matrix)
print(result["accuracy"])
print(result["f1 score"])
130
| The model has an accuracy of 0.8923 and an F1-score of 0.2849. The
     high accuracy might seem good, but 'its mainly due to the
     imbalance in the dataset. The F1-score, which considers both
     precision and recall, gives a better picture and shows that the
     model struggles to predict the less common class accurately. This
     shows why accuracy alone 'isnt a reliable metric for imbalanced
     datasets.
132
133
  135
```

```
136 # Loss matrix defined in task
  loss_matrix = matrix(c(0, 1, 5, 0), byrow = TRUE, nrow = 2,
                          dimnames = list(c("no", "yes"), c("no", "yes")))
138
139
  # Predictions on the test dataset using optimal tree
140
  predications_test = predict(optimal_tree, newdata = data_test)
  # Predictions multiplied with loss matrix
143
| losses = predications test %*% loss matrix
146 # For each observation (row) it finds the class (column)
147 # with the smallest loss and returns the index of that class.
| predicted_classes = apply(losses, 1, which.min)
  # Map the correct column names (no, yes) for the predicted classes
150
  predicted_classes = colnames(losses)[predicted_classes]
151
152
# Create confusion matrix
conf_matrix = table(data_test$y, predicted_classes)
155 conf matrix
156
157 # Print result
result = accuracy_and_f1(conf_matrix)
print(result["accuracy"])
print(result["f1_score"])
162 # The model accuracy has decreased to 0.8732, while the F1-score has
     improved to 0.4826. This change is due to the introduction of a
     loss matrix that imposes a heavier penalty on false negatives
     compared to false positives. Correct predictions, represented by
     the diagonal entries of the matrix, are not penalized.
163
  165
  tpr_and_fpr = function(conf_matrix) {
166
    TP = conf_matrix["yes", "yes"] # True positives
167
    TN = conf_matrix["no", "no"] # True negatives
FP = conf_matrix["no", "yes"] # False positives
FN = conf_matrix["yes", "no"] # False negatives
169
170
171
    tpr = TP / (TP + FN) # true positive rate
172
    fpr = FP / (FP + TN) # false positive rate
173
174
    return(c(tpr = tpr, fpr = fpr))
175
  }
176
177
478 # True positive rates and False positive rates
179 tpr_tree = c()
180 | fpr_tree = c()
181 | tpr_lrm = c()
182 fpr_lrm = c()
184 # Logistic regression model
```

```
lsm = glm(formula = y ~ ., data = data_train, family = "binomial")
186
  # Predictions on test data with both models
187
predictions test = predict(optimal tree, newdata = data test, type =
     "vector")
  predictions lrm = predict(lrm, newdata = data test, type = "response")
190
  for (pi in seq(from = 0.05, to = 0.95, by = 0.05)) {
191
    # Optimal tree predictions
192
    y_hat_optimal_tree = ifelse(predictions_test[, "yes"] > pi, "yes",
193
       "no")
    conf matrix = table(data test$y, factor(y hat optimal tree, levels
194
       = c("no", "yes")))
    result = tpr and fpr(conf matrix)
195
    tpr_tree = append(tpr_tree, result["tpr"])
196
    fpr_tree = append(fpr_tree, result["fpr"])
197
198
    # Logistic regression model
199
    y_hat_lrm = ifelse(predictions_lrm > pi, "yes", "no")
200
    conf_matrix = table(data_test$y, y_hat_lrm)
201
    result = tpr_and_fpr(conf_matrix)
202
    tpr_lrm = append(tpr_lrm, result["tpr"])
203
    fpr_lrm = append(fpr_lrm, result["fpr"])
204
205
206
  # Plot ROC curves
207
  plot(fpr_tree, tpr_tree, type = "b", col = "red", ylim =
208
     c(min(tpr_tree), max(tpr_tree))
        ,xlim = c(min(fpr_tree), max(fpr_tree)),
       main = "ROC curves", ylab = "True positive rate", xlab = "False
210
          positive rate")
  points(fpr_lrm, tpr_lrm, type = "b", col = "blue")
legend("bottomright", c("Optimal tree", "Logistic regression model"),
     fill = c("red", "blue"))
213
214 # The ROC curve in Figure \ref{fig:roc_curves} shows that the optimal
     tree model performs slightly better than the logistic regression
     model, with higher True Positive Rate (TPR) across the False
     positive Rate (FPR) values. Since the test data has very
     imbalanced classes, the precision-recall curve would be a better
     choice here. Because it focuses on the minority class, which would
     be more informative for an imbalanced dataset such as this.
```



month: apraug leb lan jul jun.may.nov pdays 2.94.5

yes yes

month: lul jun.may
no no pdays 3.93.5 housing no
no no pdays 3.93.5 housing no
month: langlage 3.94.5 yes
month: langlage

Figure 17: dependency deviation

Figure 18: optimal tree

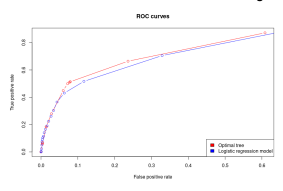


Figure 19: roc curves

# 8 Lab 2 Task 3: Principal Components and Implicit Regularization

The data file communities.csv contains the results of studies of the crime level in the United States based on various characteristics of a given location. The main variable studied is ViolentCrimesPerPop, which represents the total number of violent crimes per 100K population.

#### **Tasks**

- 1. Scale all variables except ViolentCrimesPerPop and implement PCA using the eigen() function.
  - Report how many components are needed to obtain at least 95% of the variance in the data.
  - What is the proportion of variation explained by each of the first two principal components?
- 2. Repeat the PCA analysis using the princomp() function and make the trace plot of the first principal component.
  - · Do many features have a notable contribution to this component?
  - Report which 5 features contribute the most (by absolute value) to the first principal component.

- Comment on whether these features have anything in common and whether they may have a logical relationship to the crime level.
- Provide a plot of the PC scores in the coordinates (PC1, PC2) where the color of the points corresponds to ViolentCrimesPerPop. Analyze this plot (hint: use the ggplot2 package).
- 3. Split the original data into training and test sets (50/50), scale both features and response appropriately, and estimate a linear regression model from the training data where ViolentCrimesPerPop is the target and all other data columns are features.
  - Compute the training and test errors for these data.
  - Comment on the quality of the model.
- 4. Implement a function that depends on the parameter vector  $\theta$  and represents the cost function for linear regression without intercept on the training dataset.
  - Use the BFGS method (optim() function without the gradient specified) to optimize this cost with the starting point  $\theta_0 = 0$ .
  - Compute the training and test errors for every iteration number.
  - Present a plot showing the dependence of both errors on the iteration number.
  - Comment on which iteration number is optimal according to the early stopping criterion.
  - Compute the training and test errors in the optimal model, compare them with the results in step 3, and make conclusions.

#### Hints

- **Hint 1:** Don't store parameters from each iteration (this will require a lot of memory). Instead, compute and store test errors directly.
- **Hint 2:** Discard some initial iterations (e.g., 500) in your plot to make the dependencies visible.

```
data <- read.csv('communities.csv')</pre>
 # task 1
3
5 library(caret)
 scaler<-preProcess(data)</pre>
 dataS<-predict(scaler,data)</pre>
 dataS$ViolentCrimesPerPop <- data$ViolentCrimesPerPop # no scaling on
     ViolentCrimesPerPop according to instructions
|S| < (1/101) * t(as.matrix(dataS)) %*% as.matrix(dataS) # covariance
     matrix, how variables depends on each other
11 eig <- eigen(S)</pre>
12 lambda <- eig$values
13 lambda_perc <- lambda/sum(lambda)*100 # gives the variance of each
     'projection' (how much data the projection can represent)
sprintf("%2.3f", lambda perc)
15 # prints:
16 # [1] "25.025" "16.931" "9.298" "7.554" "5.660" "4.236"
                                                                "3.226"
     "2.969" "2.067" "1.617" "1.572" "1.470"
```

```
17 # [13] "1.414" "1.030" "0.931" "0.890" "0.748"
                                                        "0.705"
                                                                 "0.649"
     "0.638" "0.624" "0.568" "0.542" "0.519"
    [25] "0.504"
                 "0.480" "0.466" "0.451"
                                              "0.431"
                                                        "0.386"
                                                                 "0.365"
18
     "0.351"
             "0.337" "0.311" "0.287"
                                         "0.259"
                                    "0.222"
    [37] "0.256" "0.245" "0.240"
19 #
                                             "0.211"
                                                        "0.205"
                                                                  "0.200"
     "0.190" "0.183" "0.165" "0.160" "0.142"
    [49] "0.138" "0.126" "0.112" "0.107" "0.104"
                                                        "0.100"
                                                                 "0.090"
     "0.081" "0.077" "0.073" "0.069" "0.066"
    [61] "0.064"
                 "0.062" "0.058"
                                    "0.051" "0.049"
                                                        "0.046"
                                                                 "0.044"
     "0.042" "0.040" "0.038" "0.035" "0.034"
    [73] "0.032"
                 "0.031" "0.028" "0.026" "0.024"
                                                        "0.022"
                                                                 "0.021"
     "0.020"
             "0.018"
                      "0.018"
                                "0.016"
                                         "0.015"
   [85] "0.014" "0.013" "0.011" "0.009" "0.008"
                                                        "0.006"
                                                                 "0.006"
     "0.005" "0.004" "0.004" "0.003" "0.002"
 # [97] "0.002" "0.001" "0.001" "0.001" "0.001"
 sum(lambda_perc[1:34]) # 94.9
 sum(lambda_perc[1:35]) # 95.2 => we need 35 variables for a variance
     of 95 %
27
 # task 2
28
29
30
 res <- princomp(dataS)
31 lambda <- res$sdev^2 # variance = standard_deviation^2</pre>
sprintf("%2.3f",lambda/sum(lambda)*100) # prints as above
33 U1<- (res$loadings)[,1]
34 plot(sort(abs(U1)), main="Traceplot, PC1")
35 top 5 <- tail(sort(abs(U1)),5)</pre>
36 top 5
37 # PctPopUnderPov
                       pctWInvInc
                                      PctKids2Par
                                                        medIncome
     medFamInc
 # 0.1737183
                       0.1748076
                                      0.1755406
                                                        0.1818171
38
     0.1831478
 # It seems that poverty is the largest contributing factor
40
41 library (ggplot2)
42 S1 <- (res$scores)[,1]
43 S2 <- (res$scores)[,2]
44
  ggplot() + geom_point(aes(x=S1, y=S2,
45
     color=dataS$ViolentCrimesPerPop)) +
    labs(title = "PC1 and PC2 scores",
46
         x = "PC1"
47
         y = "PC2") +
48
    theme minimal()
49
 # TASK 3
51
52
53 n < - dim (data) [1]
54 set.seed (12345)
|id < -sample(1:n, floor(n*0.5))|
56 train < -data[id,]
57 test<-data[-id,]</pre>
58
```

```
59 scaler <-preProcess(train)</pre>
60 trainS<-predict(scaler, train)
  testS<-predict(scaler,test)</pre>
63 fit <-lm(ViolentCrimesPerPop ~ ., data=trainS)
64 summary <- summary(fit)
66 dim(summary$coefficients)
67
68 MSE train <- mean(residuals(fit)^2) # MSE for train = 0.2752071
  MSE test <- mean((test$ViolentCrimesPerPop - predict(fit,
     newdata=testS))^2) # MSE for test = 0.5408757
70
  # We see that the MSE is quite good for train but quite high for test
     meaning it is alright but not more.
  # Slightly overfitted
  # through the summary(fit) we see that racepctblack, PctWorkMom,
     PctPersDenseHous are significant contributors (***)
  # Task 4 ########
75
76
77 # cost function for linear regression w/o intercept
78 trainS_X <- as.matrix(subset(trainS, select=-ViolentCrimesPerPop))</pre>
rel testS_X <- as.matrix(subset(testS, select=-ViolentCrimesPerPop))</pre>
80 trainS Y <- trainS$ViolentCrimesPerPop
81 testS Y <- testS$ViolentCrimesPerPop
82
83 train_MSE_vec <- c()
  test MSE vec <- c()
84
85
  MSE <- function(theta)
86
87
    MSE train <- mean( (trainS$ViolentCrimesPerPop - trainS X %*%
88
       theta)^2 )
    MSE_test <- mean( ( testS_Y - testS_X %*% theta )^2 )</pre>
89
90
    train_MSE_vec <<- c(train_MSE_vec, MSE_train)</pre>
91
92
    test_MSE_vec <<- c(test_MSE_vec, MSE_test)</pre>
93
    return(MSE_train)
94
  }
95
96
  optim(rep(0, 100), fn = MSE, method = "BFGS")
97
98
  interval <- 1:length(train_MSE_vec)</pre>
99
  ggplot() +
100
    geom_point(aes(interval, train_MSE_vec[interval]), color="red") +
101
    geom_point(aes(interval, test_MSE_vec[interval]), color="blue") +
102
    ylim(0,1) +
    labs(title = "MSE train (red) and test (blue)", x = "Iteration", y
104
       = "MSE Score")
106 # early stopping criterion says to stop at the lowest test MSE
```

```
optimal_MSE_train <- train_MSE_vec[which.min(test_MSE_vec)] # = 0.3032999

optimal_MSE_test <- min(test_MSE_vec) # = 0.4002329

# We see that this is a slight elevation from the previous MSE of 0.275 for test

# but a fairly good decrease from 0.541 for the test MSE.

# Overall this is probably a better model
```

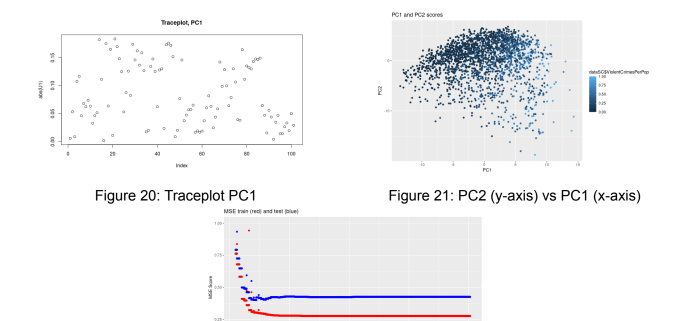


Figure 22: task 4: train = red, test = blue

# 9 Lab 3 Task 1: Kernel Method for Predicting Hourly Temperatures in Sweden

You are tasked with implementing a kernel method to predict hourly temperatures for a specific date and location in Sweden using the provided files stations.csv and temps50k.csv. These files contain data on weather stations and temperature measurements from the Swedish Meteorological and Hydrological Institute (SMHI).

# **Objective**

Provide a temperature forecast for a given date and location in Sweden. The forecast should include predicted temperatures from 4 am to midnight at 2-hour intervals.

# Methodology

Use a kernel composed of three Gaussian kernels:

- 1. **Physical Distance Kernel:** Accounts for the physical distance between a station and the point of interest. Use the distHaversine function from the geosphere package.
- 2. **Date Distance Kernel:** Accounts for the temporal distance between the measurement date and the prediction date.
- 3. **Time Distance Kernel:** Accounts for the temporal distance between the measurement time and the prediction time.

Choose appropriate smoothing coefficients ( $h_{\text{distance}}$ ,  $h_{\text{date}}$ ,  $h_{\text{time}}$ ) for each kernel manually. Show kernel values as a function of distance to demonstrate how closer points have higher values.

# **Steps**

- 1. Filter out temperature measurements that are posterior to the prediction date and time.
- 2. Compute kernel values for the physical distance, date distance, and time distance.
- 3. Combine the three kernels:
  - · First, by summing them.
  - Then, by multiplying them.
- 4. Generate forecasts for temperatures at the specified times.
- 5. Compare results obtained using the summation and multiplication methods and explain differences.

#### **Template Code**

Below is the R template code:

```
set.seed(1234567890)
 library(geosphere)
3
 # Load data
5 stations <- read.csv("stations.csv", fileEncoding = "latin1")
temps <- read.csv("temps50k.csv")</pre>
 st <- merge(stations, temps, by = "station_number")</pre>
9 # Smoothing coefficients (to be determined manually)
10 h_distance <- # Physical distance smoothing coefficient
11 h date <-
               # Date smoothing coefficient
12 h time <-
               # Time smoothing coefficient
13
# Prediction parameters
a <- 58.4274 # Latitude of the point to predict
17 date <- "2013-11-04" # Date to predict
18 times <- c("04:00:00", "06:00:00", "08:00:00", ..., "24:00:00") #
    Times
19
```

```
# Initialize temperature predictions
temp <- vector(length = length(times))

# Kernel implementation and prediction (students' code here)

# Visualization
plot(temp, type = "o", xlab = "Time", ylab = "Temperature (C)",
main = "Predicted Temperatures")
```

### **Analysis**

After implementing the above, repeat the exercise by combining the kernels through multiplication instead of summation. Discuss the differences in results and provide insights into why the two methods yield different outcomes.

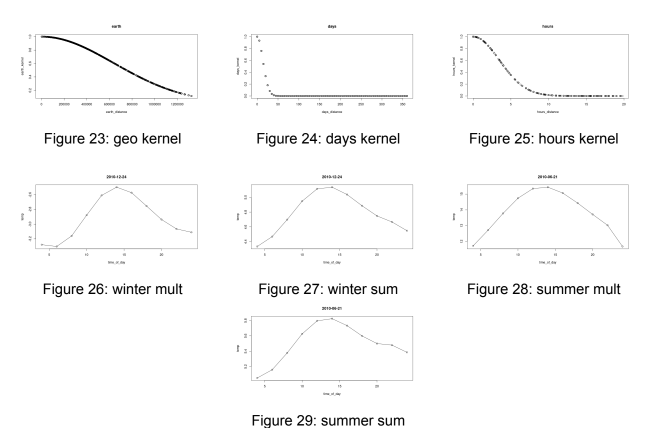
```
# setwd(paste0(getwd(),"/tdde01-machine-learning/lab3/"))
     set.seed(1234567890)
 3
     library(geosphere)
     stations <- read.csv("stations.csv", fileEncoding = "latin1")</pre>
     temps <- read.csv("temps50k.csv")</pre>
 s st <- merge(stations, temps, by="station_number")</pre>
_{10} h pos <- c(58.41086,15.62157) # Linkoping's latitude and longitude
11 h_time <- "15:10:22"
12 h_date <- "2013-11-04"
times <- c("04:00:00", "06:00:00", "08:00:00", "10:00:00", "12:00:00", "13:00:00", "13:00:00", "14:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "15:00", "1
                                   "14:00:00", "16:00:00", "18:00:00", "20:00:00", "22:00:00",
                                   "24:00:00")
15
16
     17
18
    ## finding the width of the first kernel
19
20 # max_lat_long <- c( max(st$latitude), max(st$longitude) )</pre>
    # min_lat_long <- c( min(st$latitude), min(st$longitude) )</pre>
     # max dist = distHaversine(
               max_lat_long,
23 #
24 #
                min_lat_long
25 #
     # 1 <- 6378137 / 10 # earth radius / 10
     \# \exp(-(\max_{i=1}^{n} dist^2)) / (2 * 1^2)) \# gives practically zero
             which is good
     earth distance <- c()</pre>
     gaussian kernel earth <- function(x, data)</pre>
30
     {
31
          x prim <- matrix( c(data$latitude, data$longitude), nrow =</pre>
32
                   length(data$longitude), ncol=2)
33
          1 = 6378137 / 10 # standard deviation or in this case width = earth
34
                  radius / 10
```

```
distance <- c()</pre>
35
    for (i in 1:dim(x_prim)[1])
36
37
      distance <- c(distance, distHaversine(x, x_prim[i,]))</pre>
38
39
40
    earth_distance <<- distance</pre>
41
    exp( - ( ( distance )^2 ) / ( 2 * 1^2 ) )
  }
42
43
  #
44
45
  days distance <- c()</pre>
  gaussian_kernel_days <- function(x, data)</pre>
47
  {
48
    x_prim <- data$date</pre>
49
    1 = sqrt(365 / 2) # standard deviation in gaussian kernel =
50
       sqrt(variance)
    x <- rep(x, length(x_prim))
51
    x_prim <- x_prim</pre>
52
53
    days diff <- as.numeric( difftime(x, x prim) ) %% 365
54
    days_distance <<- days_diff</pre>
55
56
    exp( - ( ( days_diff )^2 ) / ( 2 * 1^2 ) )
57
  }
58
59
  #
60
61
  hours distance <- c()
62
  gaussian kernel hours <- function(x, data)</pre>
63
64
    x prim <- as.POSIXct(data$time, format = "%H:%M:%S")</pre>
65
    x \leftarrow as.POSIXct(rep(x, length(data$time)), format = "%H:%M:%S")
66
    1 = sqrt(24 / 2) # standard deviation in gaussian kernel =
67
        sqrt(variance)
    time_diff <- abs( as.numeric( difftime(x,x_prim, units="hours") ) )</pre>
68
       %% 24
    hours_distance <<- time_diff
69
    exp(-((time diff^2)/(2*1^2)))
70
  }
71
72
  delete posterior dates <- function(current date, data)</pre>
73
74
    delete_index <- c()</pre>
75
    for (i in 1:dim(data)[1])
76
77
78
       if (difftime(current date, data$date[i]) <= 0)</pre>
79
         delete index <- c(delete index, i)</pre>
80
      }
81
    }
82
```

```
data[-delete_index,]
  }
84
85
  # delete_posterior_dates("1960-04-20", st)
86
87
88
  # further tweaking of the width (1)
89
  # dist_kernel <- gaussian_kernel_earth( h_pos, st )</pre>
  # mean( dist_kernel )
90
91 # max( dist kernel )
92 # min( dist kernel )
93 #
  # days_kernel <- gaussian_kernel_days(h_date, st)</pre>
94
95 # mean(days kernel)
  # max(days_kernel)
  # min(days_kernel)
97
  # #
98
  # # hours_kernel <- gaussian_kernel_hours(times[1], st)</pre>
99
  # mean(hours_kernel)
  # max(hours kernel)
  # min(hours_kernel)
102
  total kernel <- function(pos, date, times, data, mult add)
104
105
    data <- delete_posterior_dates(date, data)</pre>
106
107
    dist kernel <- gaussian kernel earth( pos, data )
108
    days kernel <- gaussian kernel days (date, data)
109
110
    total_kernel_vector <- c()
111
    for (i in 1:length(times))
112
113
       time kernel <- gaussian kernel hours( times[i], data )
114
       # print(time_kernel)
       if (mult add == "add")
116
         total_kernel_vector <- c(total_kernel_vector, dist_kernel +</pre>
117
            days_kernel + time_kernel)
       else
         total_kernel_vector <- c(total_kernel_vector, dist_kernel *</pre>
119
            days_kernel * time_kernel)
120
    total matrix <- matrix(total kernel vector, nrow = dim(data)[1],
121
        ncol = length(times))
    temp_pred <- c()
123
    air temperatures <- data$air temperature
124
    for (i in 1:length(times))
125
126
127
       # Nadaraya-Watson weighted average, eq 2.41, page 35, course book
128
       temp_pred <- c(temp_pred, (air_temperatures %*% total_matrix[,i])
           sum(total_matrix[,i]) )
    }
129
    temp_pred
130
131 }
```

```
132
  134
  earth_kernel <- gaussian_kernel_earth( h_pos, st )</pre>
135
  days_kernel <- gaussian_kernel_days( h_date, st )</pre>
hours_kernel <- gaussian_kernel_hours( times[1], st )
138
  plot( earth_distance, earth_kernel, main = "earth")
139
  plot( days distance, days kernel, main = "days" )
  plot( hours distance, hours kernel, main = "hours")
143 time of day <- c()
144 for (i in 1:length(times))
145
    time of day <- c(time of day, (strsplit(times[i], split =
146
       ":"))[[1]][1] )
147
  time_of_day <- as.numeric(time_of_day)</pre>
  times <- c("04:00:00", "06:00:00", "08:00:00", "10:00:00", "12:00:00",
              "14:00:00", "16:00:00", "18:00:00", "20:00:00", "22:00:00",
150
              "24:00:00")
151
152
  pos <- c(58.41086,15.62157)
153
  date <- "2010-06-21"
154
  temp <- total_kernel(pos, date, times, st, "add")</pre>
  plot(time of day, temp, type="o", main=date)
157
  print(date)
158
  date <- "2010-12-24"
159
  temp <- total kernel(pos, date, times, st, "add")</pre>
  plot(time_of_day, temp, type="o", main=date)
161
162 print (date)
164 date <- "2010-06-21"
temp <- total_kernel(pos, date, times, st, "mult")
plot(time_of_day, temp, type="o", main=date)
  print(date)
168
  date <- "2010-12-24"
169
  temp <- total_kernel(pos, date, times, st, "mult")</pre>
  plot(time_of_day, temp, type="o", main=date)
172 print (date)
173
  # We see that the sum kernel seems to always give values around 4
174
     degree celsius
  # whereas the multiplication kernel seems to give more accurate
175
     values.
176 # This is thought to be because multiplying values between 0 and 1
     really accentuates
477 # the 'strong' values close to one and it really diminishes the
     values close to 0,
178 # whereas when summing the values it is possible that two values even
     each other out
```

# making it essentially only depend on the last value which might not contribute majorly to the result.



# 10 Lab 3 Task 2 SVM Model Selection for Spam Dataset

The code in the file Lab3Block1 2021 SVMs St.R performs SVM model selection to classify the spam dataset. To do so, the code uses the function ksvm from the R package kernlab, which also includes the spam dataset. All the SVM models to select from use the radial basis function kernel (also known as Gaussian) with a width of 0.05. The C parameter varies between the models.

Run the code in the file Lab3Block1 2021 SVMs St.R and answer the following questions:

- 1. Which filter do you return to the user? filter0, filter1, filter2 or filter3? Why?
- 2. What is the estimate of the generalization error of the filter returned to the user? err0, err1, err2 or err3? Why?
- 3. Once an SVM has been fitted to the training data, a new point is essentially classified according to the sign of a linear combination of the kernel function values between the support vectors and the new point. You are asked to implement this linear combination for filter3.
  - You should make use of the functions alphaindex, coef, and b that return the indexes of the support vectors, the linear coefficients for the support vectors, and the negative intercept of the linear combination.

- See the help file of the kernlab package for more information.
- You can check if your results are correct by comparing them with the output of the function predict where you set type = "decision".
- Do so for the first 10 points in the spam dataset.
- Feel free to use the template provided in the file Lab3Block1 2021 SVMs St.R.

```
library(kernlab)
  set.seed(1234567890)
 data(spam)
 foo <- sample(nrow(spam))</pre>
 spam <- spam[foo,]</pre>
 tr <- spam[1:3000, ]
8 va <- spam[3001:3800, ]
9 trva <- spam[1:3800,]
10 te <- spam [3801:4601, ]
11 by <- 0.3
12 err_va <- NULL
13 for(i in seq(by,5,by)){
    filter <- ksvm(type~.,data=tr,kernel="rbfdot",</pre>
14
    kpar=list(sigma=0.05),C=i,scaled=FALSE)
15
    mailtype <- predict(filter, va[,-58])</pre>
16
    t <- table(mailtype, va[,58])
    err_va < -c(err_va, (t[1,2]+t[2,1])/sum(t))
18
19 }
20 filter0 <- ksvm(type~.,data=tr,kernel="rbfdot",</pre>
21 kpar=list(sigma=0.05), C=which.min(err_va)*by, scaled=FALSE)
22 mailtype <- predict(filter0, va[, -58])</pre>
23 t <- table(mailtype, va[,58])
|err0| < (t[1,2]+t[2,1])/sum(t)
25 err0
26 filter1 <- ksvm(type~.,data=tr,kernel="rbfdot",
27 kpar=list(sigma=0.05), C=which.min(err_va)*by, scaled=FALSE)
28 mailtype <- predict(filter1,te[,-58])</pre>
29 t <- table (mailtype, te[,58])
|err1| < (t[1,2]+t[2,1])/sum(t)
31 err1
filter2 <- ksvm(type~.,data=trva,kernel="rbfdot",</pre>
33 kpar=list(sigma=0.05), C=which.min(err va)*by, scaled=FALSE)
34| mailtype <- predict(filter2,te[,-58])</pre>
35 t <- table(mailtype, te[,58])
|err2| < (t[1,2]+t[2,1])/sum(t)
37
filter3 <- ksvm(type~.,data=spam,kernel="rbfdot",
sel kpar=list(sigma=0.05), C=which.min(err_va)*by, scaled=FALSE)
40 mailtype <- predict(filter3, te[,-58])
41 t <- table (mailtype, te[,58])
|42| \text{ err3} < (t[1,2]+t[2,1])/\text{sum}(t)
43 err3
 # Questions
45
46
```

```
47 # 1. Which filter do we return to the user ? filter0, filter1,
     filter2 or filter3? Why?
48 # 2. What is the estimate of the generalization error of the filter
     returned to the user? err0, err1, err2 or err3? Why?
49 # 3. Implementation of SVM predictions.
51
 sv<-alphaindex(filter3)[[1]]</pre>
 co<-coef(filter3)[[1]]</pre>
52
53 inte<- - b(filter3)</pre>
55 k < - NULL
for(i in 1:10){ # We produce predictions for just the first 10 points
     in the dataset.
    k2<-NULL
    # Calculate k values for each support vector.
58
    for(j in 1:length(sv)){
59
      # Calculate the difference between point and current support
60
          vector point
      diff <- spam[i, -58] - spam[sv[j],-58]</pre>
61
      # Calculate the value of the gaussian kernel for the current
62
         points
      k \text{ val} \leftarrow \exp(-0.05 * \text{sum}(\text{diff}^2))
63
      # Add to vector of all k_values of the current point
64
      k2 \leftarrow c(k2, k_val)
65
    }
66
    # Calculate decision function for the current point
    k \leftarrow c(k, sum(co * k2) + inte)
68
 }
69
 print(k)
 print(spam[1:10, 58])
71
predict(filter3, spam[1:10, -58], type = "decision")
```

#### Answers:

1.

- Filter0: trained on the training data, tested on validation data.
- Filter1: trained on the training data, tested on test data.
- Filter2: trained on training + validation data, tested on test data.
- Filter3: trained on the entire dataset, tested on test data.

A model performs better if it is trained on more data. As long as that data is accurate. Filter 0 and 1 are trained on only the training set, but we want our model to be trained on as much data as possible. Filter 2 is trained on training + validation data. However, filter3 is trained on the entirety of the spam data set. Therefore it's the most appropriate filter to return to someone who needs to use a model for making predictions on new data.

2.

- err0: obatained by filter0 which is trained on training data and validated on validation data.
   not appropriate becasue not trained on validation data. Validation data was used to choose best C. So it has already "seen" some of the data on which it was tested.
- err1: obatained by filter1 which is trained on training data and validated on tested data. Not

appropriate becasue not trained on validation data, I.E not the most amount of data possible which isn't test data.

- err2: obatained by filter2 which is trained on training + validation and test data. Thus an appropriate generalization error because the model is trained on the largest possible amount of data without having "seen" any of the test data.
- err3: obatained by filter3 which is trained on all data and validated on test data. Not appropriate because it has "seen" the test data.

Based on the reasoning above, err2 is the estimate of the generalization error of the filter returned to the user. The other errors are validated on the wrong dataset, are not trained on a satisfiable amount of data or have been trained on the data on which it is being validated.

The decision function for the SVM for a point  $x_*$  is the following.

$$\hat{y}(x_*) = \operatorname{sign}(\hat{\alpha}^\top K(X, x_*) + \beta)$$

Where  $\hat{\alpha}^{\top}$  is a vector where non-zero values represent the support vectors.  $K(X, x_*)$  is a vector created by applying the kernel on  $x_*$  and each of the training points in X.

$$K(X, x_*) = \exp(-\sigma ||X - x_*||^2)$$

 $\beta$  is the intercept of the SVM.

ANSWER TASK 3: Using the decision function, we can classify an email as spam accordingly.

$$class = \begin{cases} \mathsf{nonspam}, & \mathsf{if} \ \mathsf{sign}(\hat{y}(x_*)) = -\\ \mathsf{spam}, & \mathsf{if} \ \mathsf{sign}(\hat{y}(x_*)) = + \end{cases}$$

The result of the SVM predictions on the first 10 points in the spam dataset can be seen in table 1

Data Point	Prediction (Class)	Decision Value ( $\hat{y}(x_*)$ )
1	Nonspam	-1.0703
2	Spam	1.0003
3	Spam	0.9996
4	Nonspam	-0.9999
5	Nonspam	-0.9995
6	Spam	1.0001
7	Nonspam	-0.8586
8	Nonspam	-0.9997
9	Spam	0.9998
10	Nonspam	-1.0001

Table 1: SVM Predictions for the First 10 Data Points in the spam dataset

### 11 Lab 3 Task 3: Neural Networks

This assignment is to be solved using the neuralnet package.

#### **Tasks**

- 1. Train a neural network to learn the trigonometric sine function.
  - Sample 500 points uniformly at random in the interval [0, 10].
  - Apply the sine function to each point. The resulting value pairs are the data points available to you.
  - Use 25 of the 500 points for training and the rest for testing.
  - Use one hidden layer with 10 hidden units. You do not need to apply early stopping.
  - Plot the training and test data, and the predictions of the learned neural network on the test data.
  - Comment on your results.
- 2. In question (1), you used the default logistic (a.k.a. sigmoid) activation function, i.e., act.fct = "logistic".
  - Repeat question (1) with the following custom activation functions:

$$h_1(x) = x, \quad h_2(x) = \max\{0, x\}, \quad h_3(x) = \ln(1 + e^x)$$

(a.k.a. linear, ReLU, and softplus).

- See the help file of the neuralnet package to learn how to use custom activation functions.
- Plot and comment on your results.
- 3. Sample 500 points uniformly at random in the interval [0, 50] and apply the sine function to each point.
  - Use the neural network learned in question (1) to predict the sine function value for these new 500 points.
  - You should get mixed results. Plot and comment on your results.
- 4. In question (3), the predictions seem to converge to some value.
  - Explain why this happens. To answer this question, you may need to access the weights of the learned neural network.
  - You can do this by running nn or nn\$weights, where nn is the learned neural network.
- 5. Sample 500 points uniformly at random in the interval  $\left[0,10\right]$  and apply the sine function to each point.
  - Use all these points as training points for learning a neural network that predicts x from  $\sin(x)$ , i.e., unlike before when the goal was to predict  $\sin(x)$  from x.
  - Use the learned neural network to predict the training data.
  - You should get bad results. Plot and comment on your results.
  - **Help:** Some people get a convergence error in this question. This can be solved by stopping the training before reaching convergence by setting threshold = 0.1.

```
_{7} n = 500
8 set.seed (1234567890)
points = runif(n, min=0, max=10)
data <- data.frame(points, sin=sin(points))</pre>
12 # Split the dataset into 25 points for training and rest for testing
training_amount = 25
train data = data[1:training amount, ]
test data = data[training amount:n,]
16
47 # Random initialization of the weights in the interval [-1, 1]
18 winit = runif(10, -1, 1)
nn = neuralnet(formula = sin ~ ., data = train data, hidden = 10,
     startweights = winit)
20
21 # Plot of the training data (black), test data (blue), and
    predictions (red)
plot(train_data, cex=2, main = "Predictions of sin(x) from x")
points(test data, col = "blue", cex=1)
points(test_data[, 1], predict(nn, test_data), col="red", cex=1)
 legend("bottomright",
26
        legend = c("Training Data", "Test Data", "Predictions"),
27
        col = c("black", "blue", "red"),
28
        pch = 1,
        pt.cex = c(2, 1, 1))
30
31
_{
m 32} # it can be seen that the model's predictions for the sine points
    using the test data are highly accurate. The predicted values
     closely overlap with the actual test data, indicating strong model
    performance.
 35
36 h1 = function(x) x # Linear
_{37} h2 = function(x) ifelse(x >= 0, x, 0) # ReLU
_{38} h3 = function(x) log(1 + exp(x)) # Softplus
nn1 = neuralnet(formula = sin ~ ., data = train_data, hidden = 10,
    act.fct = h1, startweights = winit)
40 nn2 = neuralnet(formula = sin ~ ., data = train data, hidden = 10,
    act.fct = h2, startweights = winit)
nn3 = neuralnet(formula = sin ~ ., data = train_data, hidden = 10,
    act.fct = h3, startweights = winit)
plot(train_data, cex=2, main = "Predictions of sin(x) from x")
44 points(test_data, col = "blue", cex=1)
45 points(test_data[, 1], predict(nn1, test_data), col="red", cex=1)
46 points(test_data[, 1], predict(nn2, test_data), col="green", cex=1)
points(test_data[, 1], predict(nn3, test_data), col="orange", cex=1)
48
49 legend ("bottomright",
        legend = c("Training Data", "Test Data", "NN1 (Linear)", "NN2
```

```
(ReLU)", "NN3 (Softplus)"),
         col = c("black", "blue", "red", "green", "orange"),
51
        pch = 1,
52
        pt.cex = c(2, 1, 1, 1, 1))
53
 # Figure shows that the Linear model performs poorly in predicting
    \sin(x) from the data points, as a linear model cannot fit a
    sinusoidal curve. The ReLU activation function shows promising
    results for the initial data points but becomes worse after four
    points. This decline occurs because ReLU is a piecewise linear
    function, meaning it is linear in segments. Therefore, it
    struggles to capture the smooth, continuous oscillations of a
    sinusoidal function. On the other hand, the Softplus activation
    function allows the neural network to perfectly predict the test
    data. This is because Softplus is a smooth approximation of ReLU,
    making it well-suited for modeling non-linear functions.
 58
 set.seed (1234567890)
 points = runif(n, min=0, max=50)
 data <- data.frame(points, sin=sin(points))</pre>
61
62
 predictions = predict(nn, data)
63
64
 plot(data, cex=2, ylim = c(min(predictions), max(predictions)), main
    = "Predictions of sin(x) from x")
 points(data[, 1], predictions, col="red", cex=1)
 legend("topright",
68
        legend = c("New Data", "Predictions"),
69
        col = c("black", "red"),
70
        pch = 1,
71
        pt.cex = c(2, 1)
72
73
 # As shown in Figure the original NN model predicts the new data
    points perfectly until the after approximately x = 10 points. This
    is because the NN model is not trained for points larger than 10,
    leaving it unable to generalize the behavior of the sine function
    outside its training range.
76 ####################### Part 4 ################################
78 nn $weights
79
 #[[1]]
80
81 # [[1]] [[1]]
82
              [,1]
                         [,2]
                                    [,3]
                                               [,4]
                                                         [,5]
               [,7]
                         [,8]
                                    #[,9]
                                               [,10]
83 #[1,] 3.3046893 -0.3271777
                               0.4043669 -0.7669631 11.955191
     -0.2057986 -6.472200 7.904868 #-0.5708964 0.04342472
84 #[2,] -0.8033506 -0.8324709 -0.1499125 -0.8256431 -1.802597
    0.7993943 3.082136 -2.320714 #0.1543628 0.76288667
```

```
#[[1]][[2]]
         0.7993476
88 #
    [1,]
89 # [2,] -4.9649102
90 # [3,] -4.8295057
91 # [4,] 22.1703831
92 # [5,]
        -5.5590648
93 # [6,] -4.3747945
94 # [7,]
         0.3489759
95 # [8,] -0.7224382
96 # [9.]
         1.8612110
97 # [10,] -9.4390597
        0.4400295
  #[11,]
98
99
  #When examining the NN weights, especially in the second layer, we
100
     can observe that most of them are #negative. This dominance of
     negative weights will influence the networks output, especially
     for #outputs outside the training range. When the NN encounters
     data it hasnt been trained on, it relies #heavily on the these
     learned weights to make predictions. The effect of this causes the
     network's #output to converge to a negative value for such unseen
101
  102
  set.seed (1234567890)
104
  points = runif(n, min=0, max=10)
  data <- data.frame(points, sin=sin(points))</pre>
107
nn = neuralnet(formula = points ~ ., data = data, hidden = 10,
     startweights = winit, threshold = 0.1)
|plot(x = data[, 2], y = data[, 1], cex=2, main = "Predictions of x]
                 , xlab = "sin", ylab = "points")
     from sin(x)"
  points(x = data[, 2], y = predict(nn, data), col = "red", cex = 1)
113
  legend("bottomleft"
         legend = c("Training Data", "Predictions"),
114
         col = c("black", "red"),
115
         pch = 1,
         pt.cex = c(2, 1)
117
118
  # The result of the NN trying to predict x from sin(x) can be seen in
119
     Figure \ref{fig:a3 t5}. This happens because its easier to predict
     sin(x) from x because the sine function provides a predictable
     output for each input x. The other way around is much harder, for
     the NN to learn the inverse of sine function, which is periodic,
     meaning that multiple values of x produce the same sin(x).
```

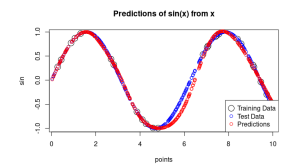


Figure 30: task 1

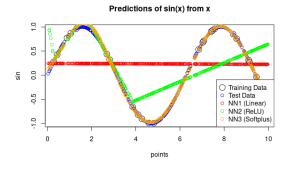


Figure 31: task 2

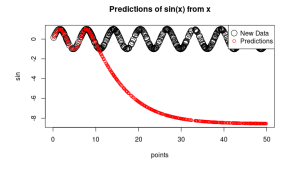


Figure 32: task 3

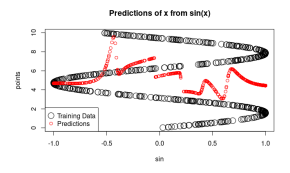


Figure 33: task 5

# 12 Exam 1

### 12.1 Assignment 1 (10p)

File Bikes.csv contains counts of public bicycles rented per hour in the Seoul Bike Sharing System, with corresponding weather data and holiday information.

- 1. Divide the data randomly into training and test data (70/30) and scale them appropriately. Compute a LASSO regression model by cross-validation in which Rented Bikes is the target variable and all remaining variables are features, and present a dependence of the cross-validation error on the penalty parameter. Which value of the penalty parameter is the optimal one? Interpret this model in terms of bias-variance tradeoff. Finally, report the equation showing how the predicted number of Rented bikes (scaled) depends on the features (scaled) in the estimated LASSO model corresponding to "lambda.1se" penalty value. (4p)
- Consider lambda. 1se LASSO model, use the training error MSE as an estimate of the target variance parameter and report the 95% prediction interval for the first observation (first row) in the test data. Explain the model assumptions making these computations possible. (2p)
- 3. Consider the same partitioned and scaled data as in step 1. Assume now that Dew Point Temperature is related to the features Humidity and Visibility as a linear model without intercept, and assume that the loss and the error functions are given by the following

formula:

$$L(y, \hat{y}) = E(y, \hat{y}) = |y - \hat{y}|$$

Implement a code optimizing the cost function by the BFGS optimizer, and plot a dependence of the cost values and the test errors on the iteration number. Is early stopping needed? Report the optimal iteration number. Make 3 scatter plots of (Humidity, Visibility) where observations are colored by:

- a) original target values,
- b) predicted target values from the optimal model,
- c) target values from the modelcorresponding to 5 iterations, and compare the plots with respect to the complexity of the model and quality of prediction. **(4p)**

Hint: to make scatter plots, you may use this kind of code from package ggplot2:

```
df = data.frame(x = your_variable1, y = your_variable2, color = your_variable3)
ggplot(df, aes(x = x, y = y, color = color)) + geom_point()
```

### 12.2 Assignment 1 Answers

```
n=dim(data)[1]
 set.seed (12345)
id=sample(1:n, floor(n*0.6))
4 train=data[id,]
5 test=data[-id,]
7 library (caret)
9 #Assignment 1 Part 1
scaler=preProcess(train)
trainS=predict(scaler, train)
testS=predict(scaler, test)
13
 covariates=trainS[,-1]
 response=trainS[,1]
15
16
17 library (glmnet)
18
19 set.seed (12345)
20 model = cv.glmnet(as.matrix(covariates),
                   response, alpha=1,family="gaussian")
21
22 model $lambda.min
 coef(model, s="lambda.1se")
24
25
26 ## 11 x 1 sparse Matrix of class "dgCMatrix"
 ##
27
28 ## (Intercept)
                            -8.742707e-17
29 ## Hour
                             2.264222e-01
30 ## Temperature
                             2.852890e-01
31 ## Humidity
                            -9.808210e-02
32 ## Wind.speed
```

```
33 ## Visibility
34 ## Dew.point.temperature
35 ## Solar.Radiation
36 ## Rainfall
                             -1.702395e-02
37 ## Snowfall
                             -6.135284e-02
38 ## Holiday
                              1.086443e-01
39
40 #Assignment 1 Part 2
41 Ypred=predict(model, as.matrix(covariates), s="lambda.1se")[,1]
42 MSE=mean((Ypred-response)^2)
43
44 testX = testS[1, -1]
45 Ytest=predict(model, as.matrix(testX), s="lambda.1se")
  CI=c(Ytest-1.96*sqrt(MSE), Ytest+1.96*sqrt(MSE))
47
  print(CI)
48
49
  ## [1] -2.031268 1.240571
51
52 #Assignment 1 part 3
53
  costF<-function(Y, Yfit){</pre>
54
   R=abs(Y-Yfit)
55
    return(mean(R))
56
57 }
|x=as.matrix(trainS[,c(4,6)])|
|xt=as.matrix(testS[,c(4,6)])|
62 Fs=list()
_{63}|k=0
64 TestE=list()
65 Theta=list()
66
  myCost= function(theta){
67
    f=costF(trainS$Dew.point.temperature, x%*%theta)
68
    .GlobalEnv$k= .GlobalEnv$k+1
69
    .GlobalEnv$Fs[[k]]=f
70
    .GlobalEnvTestE[[k]] = costF(testSDew.point.temperature, xt%*%theta)
71
    .GlobalEnv$Theta[[k]]=theta
72
    return(f)
73
  }
74
75
  res=optim(rep(0,2), fn=myCost, method="BFGS")
76
77
78
  plot((as.numeric(Fs)), type="1", col="blue")
  points((as.numeric(TestE)), type="1", col="red")
  which.min(TestE)
82
83
84 ## [1] 49
85
```

```
86 | Pred1=xt%*%Theta[[which.min(TestE)]]
  Pred2=xt%*%Theta[[5]]
89
  df=data.frame(x=xt[,1], y=xt[,2], color=testS$Dew.point.temperature)
  ggplot(df,aes(x=x,y=y,color=color))+geom_point()
  df=data.frame(x=xt[,1], y=xt[,2], color=Pred1)
93
  ggplot(df,aes(x=x,y=y,color=color))+geom point()
  df=data.frame(x=xt[,1], y=xt[,2], color=Pred2)
  ggplot(df,aes(x=x,y=y,color=color))+geom point()
97
98
  #Note If the student makes these plots with training data or full
     data, no point reduction should be done since assignment does not
     specify which data to use in plots
100
101 The optimal prediction model captures the trend and the original
     scale well, while the model with 4 iterations captures the trend
     but not the scale, all predictive values are near zero implying
     that the model is too simple for this data (underfitted).
```

# 12.3 Exam 1: Assignment 2 (10p)

#### 12.3.1 EXERCISE 1 – 5 POINTS

In January 2023, the students were asked to implement the backpropagation algorithm for training a neural network for regression as it appears in the course textbook and slides. The solution is available to you in the file TDD0211January2023.R. Now, you are asked to incorporate dropout into this solution. Recall that dropout is a regularization technique whose detailed description you can find in the course textbook and slides.

Run your implementation with a dropout rate r=1 equal to  $0,\,0.1$ , and 0.05, i.e.,  $r=1,\,0.99,\,0.95$ . Comment on the results.

#### 12.3.2 EXERCISE 2 - 5 POINTS

You are asked to implement the perceptron algorithm. This algorithm for binary classification is a predecessor of modern neural networks.

Consider a binary classification problem with class labels  $t \in \{-1, +1\}$ . Then, the class label assigned to a point **z** is given by:

$$y(\mathbf{w}, \mathbf{z}) = \begin{cases} +1 & \text{if } \mathbf{w}^T \mathbf{z} \geq 0 \\ -1 & \text{if } \mathbf{w}^T \mathbf{z} < 0 \end{cases}$$

The values of **w** are iteratively determined with the help of the learning data  $\{(\mathbf{x}_1, t_1), \dots, (\mathbf{x}_N, t_N)\}$ . Specifically, the *i*-th weight is updated in each iteration as follows:

$$w_i^{(t+1)} = w_i^{(t)} + \alpha \sum_{n=1}^{N} \left( t_n - y(\mathbf{w}^{(t)}, \mathbf{z}_n) \right) z_{n,i}$$

where  $\alpha$  is the learning rate. Finally, you can assume that  $\mathbf{z}_n$  and  $\mathbf{w}$  are of dimension two, i.e.,  $\mathbf{z}_n = (x_{n,1}, x_{n,2})$  and  $\mathbf{w} = (w_1, w_2)$ .

You can stop the learning process after 100 iterations. You can use an alpha value of 0.0001. Plot the non-misclassification rate on the dataset below as a function of the number of iterations.

```
set.seed(1234)

x <- array(NA, dim = c(100, 2))

t <- array(NA, dim = c(100))

x[,1] <- runif(100,0,3)

x[,2] <- runif(100,0,9)

t <- ifelse(x[,2]<(x[,1]^2),-1,1)

plot(x[,1],x[,2],col=t+2)</pre>
```

Finally, explain when the perceptron algorithm works best and why it works in those cases.

### 12.4 Assignment 2 Answers

```
# exercise 1 - 5 p
2 # 3 p for implementation
3 # 1 p if it runs correctly
_{4}| # 1 p for answering the question correctly
# lines corresponding to dropout implementation are marked with ###,
     the rest of the code was provided in the exam
 set.seed(1234)
 # produce the training data in dat
10
|x| < runif(500, -4, 4)
|y| < - \sin(x)
13 dat \leftarrow cbind(x,y)
14 plot (dat)
 gamma <- 0.01
 r <- 1 ### dropout rate 1-r
17
18
19 h <- function(z){</pre>
  # activation function (sigmoid)
```

```
return(1/(1+exp(-z)))
24
25
      hprime <- function(z){
26
27
28
             # derivative of the activation function (sigmoid)
29
            return(h(z) * (1 - h(z)))
30
31 }
32
33 yhat <- function(x){</pre>
34
             # prediction for point x
35
36
37 q0 <- x
|z| = |z| < - w1 \% * % q0 + b1
|q1 < -as.matrix(apply(z1,1,h), nrow = 2, ncol = 1)
|z| = |z| 
41 return(z2)
42 }
43
      yhat2 <- function(x){</pre>
44
45
             ### final prediction for point x (it involves r)
46
47
48
            q0 <- x
            z1 \leftarrow (r*w1) %*% q0 + b1
49
            q1 \leftarrow as.matrix(apply(z1,1,h), nrow = 2, ncol = 1)
50
            z2 \leftarrow (r*w2) %*% q1 + b2
51
             return(z2)
52
53 }
55 MSE <- function() {
56
            # mean squared error
57
58
59
            res <- NULL
             for(i in 1:nrow(dat)){
60
                   res <- c(res,(dat[i,2] - yhat(dat[i,1])) ^ 2)
61
62
             return(mean(res))
63
      }
64
65
      # initialize parameters
67
68 w1 <- matrix(runif(2,-.1,.1), nrow = 2, ncol = 1)
69 b1 <- matrix(runif(2,-.1,.1), nrow = 2, ncol = 1)
v_1 = v_2 < matrix(runif(2, -.1, .1), nrow = 1, ncol = 2)
72 b2 <- matrix(runif(1,-.1,.1), nrow = 1, ncol = 1)
73
74 res <- NULL
```

```
75 for (i in 1:100000) {
     if(i %% 1000 == 0){
       res <- c(res, MSE())
77
78
79
    # forward propagation
80
81
     j <- sample(1:nrow(dat),1)</pre>
82
     q0 <- dat[j,1]
83
     m0 \leftarrow sample(c(0,1),1,replace = TRUE, c(1-r,r)) ### drop input unit
     z1 \leftarrow w1 \% \% (q0 * m0) + b1
85
     q1 \leftarrow as.matrix(apply(z1,1,h), nrow = 2, ncol = 1)
86
     m1 \leftarrow sample(c(0,1),2,replace = TRUE, c(1-r,r)) ### drop hidden
87
     z2 \leftarrow w2 \% \% (q1*m1) + b2
88
89
90
     # backward propagation
     dz2 < - - 2 * (dat[j,2] - z2)
92
     dq1 \leftarrow t(w2) \% \% dz2
93
     dz1 \leftarrow dq1 * hprime(z1)
     dw2 \leftarrow dz2 %*% t(q1)
95
     db2 <- dz2
96
     dw1 <- dz1 %*% t(q0)
97
     db1 <- dz1
98
99
     # parameter updating
100
101
     w2 \leftarrow w2 - (gamma * dw2 * m1) ### update non-dropped hidden units
102
     b2 <- b2 - gamma * db2
103
     w1 <- w1 - (gamma * dw1 * c(m0,m0) * m1) ### update non-dropped
104
        hidden units
     b1 <- b1 - gamma * db1
106
  plot(res, type = "1")
107
108
  plot(dat)
points(dat[,1],lapply(dat[,1],yhat2),col="red") ### apply final
      prediction
111
112 # Lower r value implies larger regularization and, thus, worse fit of
      the training data. The ultimate goal of dropout is
113 # to get a better fit of the test data (i.e., low generalization
      error). However, this is too simple an example to observe it.
114
115 # exercise 2 - 5 p
116 # 3 p for implementation
117 # 1 p if it runs correctly
118 # 1 p for answering the question correctly
119
120 set.seed (1234)
|x| < - array(NA, dim = c(100,2))
```

```
|t| < - array(NA, dim = c(100))
|x[,1]| < runif(100,0,3)
|x[,2]| < runif(100,0,9)
t < \text{ifelse}(x[,2] < (x[,1])^2,-1,1)
|plot(x[,1],x[,2],col=t+2)|
128
129
  W < -c(0,0)
  alpha <- 0.0001
130
131
  res <- NULL
  for(i in 1:100){
133
    res <- c(res, sum(t != (2 * ((w %*% t(x[,1:2]))>0) - 1))) # note
134
        that t is both class label and transpose operation :-(
135
    w[1] \leftarrow w[1] + alpha * sum((t - w %*% t(x[,1:2])) * x[,1])
136
    w[2] \leftarrow w[2] + alpha * sum((t - w %*% t(x[,1:2])) * x[,2])
137
138
  plot(x[,1],x[,2],col=(2 * ((w %*% t(x[,1:2]))>0) - 1)+2)
  plot(res, type = "1")
141
142
  # It works for linearly separable datasets. If the true label is
     larger than the prediction, then the weights get
  # increased for positive input values and decreased for negative
     input values. The opposite when the true label is smaller.
```

### 13 Exam 2

### **13.1** Assignment 1 (10p)

File Rice.csv contains a total of 3810 rice grain's images taken for the two species (Cammeo and Osmancik), which were processed and feature inferences were made. 7 morphological features were obtained for each grain of rice which are the variables in the data set.

- 1. Divide the data randomly into training and test data (70/30). Assume that columns Area, ..., Extent are denoted as  $x_1,...,x_p$ . Perform basis function expansion with basis functions  $\phi_1=x_1,...,\phi_p=x_p,\phi_{p+1}=x_1^2,...,\phi_{2p}=x_p^2$  and fit two logistic regression models: one with features  $x_1,...,x_p$  and another with  $\phi_1,...,\phi_{2p}$ , both with target Class to the training data. Report the training and test misclassification errors for both models and report which model is better and why. Finally, report the estimated probabilistic model for the model with features  $x_1,...,x_p$ . (4p)
- 2. Use cross-validation to fit a decision tree model to the training data with target Class and all other columns  $(x_1, ..., x_p)$  as features and report the optimal number of leaves and training and test misclassification errors for the optimal tree. Comment on the prediction quality of the optimal tree. After this, write a loop that grows decision trees with parameter mindev = 0.001, 0.002, ... 0.01 (without cross-validation) and estimates training and test misclassification errors for these trees. Plot the dependence of these errors on mindev, comment on the trends observed in the plot and comment how the trends are expected to look in theory and why. (4p)

3. Use the model estimated by the cross-validation in step 2 and assume that a user wants to prune this tree. If we use misclassification error as impurity measure, which leaves must be pruned first, 14 and 15, or 4 and 5? Report necessary mathematical calculations to support your answer. (2p)

### 13.2 Assigment 1: Answers

```
#Part 1
  df=read.csv("Rice.csv", stringsAsFactors = T)
  library(dplyr)
6 \mid n = dim(df)[1]
7 set.seed (12345)
| id=sample(1:n, floor(n*0.7)) |
g train=df[id,]
10 test=df[-id,]
12
  train1=train%>%mutate_if(is.numeric, list(x2=function(x) x^2))
13
test1=test%>%mutate_if(is.numeric, list(x2=function(x) x^2))
16 m1=glm(Class~., data=train, family = binomial)
17
m2=glm(Class~., data=train1, family=binomial)
19
20 missclass=function(X,X1){
    n=length(X)
21
    return(1-sum(diag(table(X,X1)))/n)
22
23
24
25
26
  missclass(train$Class, predict(m1, newdata = train,
     type="response") > 0.5)
28
  # [1] 0.06861642
30
  missclass(test$Class, predict(m1, newdata = test,
31
     type="response") > 0.5)
32
  # [1] 0.07524059
33
34
  missclass(train$Class, predict(m2, newdata = train1,
     type="response") > 0.5)
36
  # [1] 0.06786652
37
38
  missclass(test$Class, predict(m2, newdata = test1,
     type="response") > 0.5)
40
41 # [1] 0.07611549
```

```
42
 #We can see that after basis function expansion the training error
     gets smaller and test error gets larger which indicates
     overfitting. The model with original features is best one
44
45
 m 1
46
47
  ##
 ##
            glm(formula = Class ~ ., family = binomial, data = train)
     Call:
48
 ##
49
 ##
    Coefficients:
50
                                                       Perimeter
 ##
           (Intercept)
                                        Area
51
     Major_Axis_Length
              -7.910612
                                   0.006971
 ##
                                                        0.107500
     -0.112395
                                                     Convex_Area
 ## Minor_Axis_Length
                               Eccentricity
53
                  Extent
               0.553997
 ##
                                  -3.747786
                                                       -0.011754
     0.315431
 ##
55
 ## Degrees of Freedom: 2666 Total (i.e. Null); 2659 Residual
 ## Null Deviance:
                            3656
57
 ## Residual Deviance: 920.9
                                    AIC: 936.9
58
59
60 levels (train $Class)
 ## [1] "Cammeo"
                   "Osmancik"
62
63
64
```

The probabilistic model is

$$P(\mathsf{Class} = \mathsf{Osmancik}) = \frac{1}{1 + \exp(z)}$$

where

 $z=7.910612-0.006971\cdot \text{Area} -0.107500\cdot \text{Perimeter} +0.112395\cdot \text{MajorAxisLength} -0.553997\cdot \text{MinorAxisLength} +3.7910612-0.006971\cdot \text{Area} -0.107500\cdot \text{Perimeter} +0.112395\cdot \text{MajorAxisLength} -0.553997\cdot \text{MinorAxisLength} +3.7910612-0.006971\cdot \text{Area} -0.107500\cdot \text{Perimeter} +0.112395\cdot \text{MajorAxisLength} -0.553997\cdot \text{MinorAxisLength} +3.7910612-0.006971\cdot \text{Area} -0.107500\cdot \text{Perimeter} +0.112395\cdot \text{MajorAxisLength} +0$ 

```
#Part 2
 library(tree)
 fit=tree(Class~., data=train)
 set.seed (12345)
5
6 cv.res=cv.tree(fit)
 finalTree0=prune.tree(fit, best=cv.res$size[which.min(cv.res$dev)])
 Yfit1=predict(finalTree0, newdata=train,
               type="class")
 Yfit2=predict(finalTree0, newdata=test,
10
                type="class")
11
12
13 cv.res$size[which.min(cv.res$dev)]
14
```

```
15 # [1] 5
16
 missclass(train$Class,Yfit1)
18
19 # [1] 0.06861642
20
21
 missclass(test$Class,Yfit2)
22
 # [1] 0.07786527
23
24
 #The tree has a good prediction quality since test error is only
25
     around 7 percent.
26
 mind=seq(0.001, 0.01, 0.001)
27
 1M=length(mind)
28
 TrE=numeric(1M)
30 TeE=numeric(1M)
31
 for (i in 1:1M){
32
    fit=tree(Class~., data=train,
33
       control=tree.control(nobs=nrow(train),mindev=mind[i]) )
    TrE[i]=missclass(predict(fit, newdata=train, type="class"),
       train$Class)
    TeE[i]=missclass(predict(fit, newdata=test, type="class"),
35
       test $Class)
36
 }
37
38
 plot(mind, TrE, col="blue", ylim=c(0.05,0.15))
40 points (mind, TeE, col="red")
```

```
#Part 3
 print(finalTree0)
3
    node), split, n, deviance, yval, (yprob)
5
 ##
           * denotes terminal node
6
 ##
      1) root 2667 3656.0 Osmancik ( 0.437945 0.562055 )
 ##
8
        2) Major Axis Length < 191.075 1488 657.3 Osmancik ( 0.057796
 ##
9
     0.942204)
 ##
          4) Major_Axis_Length < 181.225 1116
                                                141.6 Osmancik (
10
     0.011649 0.988351 ) *
          5) Major Axis Length > 181.225 372
                                              368.4 Osmancik ( 0.196237
 ##
11
     0.803763) *
        3) Major Axis Length > 191.075 1179 670.3 Cammeo ( 0.917727
 ##
12
     0.082273 )
 ##
          6) Perimeter < 467.639 198
                                       258.4 Cammeo ( 0.641414 0.358586
13
                                       240.1 Cammeo ( 0.973496 0.026504 )
 ##
          7) Perimeter > 467.639 981
14
 ##
           14) Major Axis Length < 205.866 409
                                                 188.2 Cammeo ( 0.938875
15
     0.061125 ) *
16 ##
                                                 14.7 Cammeo ( 0.998252
           15) Major_Axis_Length > 205.866 572
```

```
0.001748 ) *

ImpurityDiff0=0.057796*1488-(0.011649*1116 +0.196237*372)

ImpurityDiff0

## [1] 1.421085e-14

ImpurityDiff1=0.026504*981 -(0.061125*409+0.001748*572 )

ImpurityDiff1

## [1] 0.000443

#4 and 5 need to be pruned first since provide smallest impurity decrease
```

## 13.3 Assignment 2 (10p)

You are asked to implement the backpropagation algorithm for training a neural network for regression as it appears in the course textbook and slides. To do so, fill in the code provided below. The actual network has one hidden layer with two units. x denotes inputs, y denotes targets, y denotes activation units, and z denotes hidden units. As the result of applying the activation function to z,  $\sigma(z)$  denotes the squashed error, and y denotes the loss (mean squared error). The steps involved in this task are: (1) forward propagation, (2) backward propagation, (3) parameter update as marked in substeps 1-3B. Except the one indicated with a \*, that is element-wise product. (3B). Notice the use of matrix transposition in some steps (3B) in R. Comment your code and add results. The exercise will be graded as follows: Forward propagation 2p, backward propagation 4p, parameter updating 1p, and results 3p.

### Steps:

· Forward propagation.

$$q^{(0)} = x$$

$$z^{(1)} = W^{(1)}q^{(0)} + b^{(1)}$$

$$q^{(1)} = h(z^{(1)})$$

$$z^{(2)} = W^{(2)}q^{(1)} + b^{(2)}$$

$$J(\theta) = (y - z^{(2)})^2$$

Backward propagation.

$$dz^{(2)} = -2(y - z^{(2)})$$

$$dq^{(1)} = W^{(2)T}dz^{(2)}$$

$$dz^{(1)} = dq^{(1)} \odot h'(z^{(1)})$$

$$dW^{(2)} = dz^{(2)}q^{(1)T}$$

$$db^{(2)} = dz^{(2)}$$

$$dW^{(1)} = dz^{(1)}q^{(0)T}$$

$$db^{(1)} = dz^{(1)}$$

Parameter updating.

$$\begin{split} W_{t+1}^{(2)} &= W_t^{(2)} - \gamma dW^{(2)} \\ b_{t+1}^{(2)} &= b_t^{(2)} - \gamma db^{(2)} \\ W_{t+1}^{(1)} &= W_t^{(1)} - \gamma dW^{(1)} \\ b_{t+1}^{(1)} &= b_t^{(1)} - \gamma db^{(1)} \end{split}$$

### **Code Template:**

You are requested to use the template below. Note that the algorithm performs 100000 iterations. In each iteration, one randomly selected training point is used to update the parameters (i.e., this essentially corresponds to stochastic gradient descent with a mini-batch of size 1).

```
# produce the training data in R
  x = runif(500, -4, 4)
|y| = \sin(x)
|dat| = cbind(x,y)
5 plot(dat)
  gamma = 0.01
  h = function(z) {
   # activation function (sigmoid)
    return(1/(1+exp(-z)))
11
  }
12
13
14 hprime = function(z){
    # derivative of the activation function (sigmoid)
15
    return(h(z) * (1 - h(z)))
16
17
18
19 yhat <- function(x){</pre>
  # prediction for point x
21 }
22
23 MSE <- function(res){
  # mean squared error
25 }
26
# initialize parameters
28 res <- NULL
29 for (it in 1:100000) {
    if(it \%\% 10000 == 0){
30
      res <- c(res, MSE())
31
32
33
    # forward propagation
34
    j <- sample(1:nrow(dat),1)</pre>
36
    x0 <- dat[j,1]
37
    # backward propagation
```

```
# parameter updating

to plot(res, type = "l")

ploat(dat)
points(dat[,1],apply(dat[,1],1,yhat))
```

### 13.4 Assignment 2: Answers

```
set . seed (1234)
        # produce the training data in dat
  5 x <- runif (500, -4,4)
        y \leftarrow sin(x)
        dat <- cbind(x,y)</pre>
  8 plot(dat)
10 gamma <- 0.01
12 h <- function(z){
13
                # activation function (sigmoid)
14
15
                return(1/(1+exp(-z)))
16
        }
17
18
hprime <- function(z){</pre>
20
                 # derivative of the activation function (sigmoid)
21
22
                return(h(z) * (1 - h(z)))
23
        }
24
26 yhat <- function(x){</pre>
27
                # prediction for point x
28
29
        q0 <- x
30
z_1 | z_1 < - w_1 | ** q_0 + b_1
|q1| < as.matrix(apply(z1,1,h), nrow = 2, ncol = 1)
|z| |z| < - w2 | |x| |
34 return(z2)
35 }
37 MSE <- function(){
38
                # mean squared error
39
40
                res <- NULL
41
                for(i in 1:nrow(dat)){
42
                         res <- c(res,(dat[i,2] - yhat(dat[i,1])) ^ 2)
43
```

```
return(mean(res))
46
47
48 # initialize parameters
50 w1 <- matrix(runif(2,-.1,.1), nrow = 2, ncol = 1)
|b1| < matrix(runif(2, -.1, .1), nrow = 2, ncol = 1)
|w2| < -matrix(runif(2, -.1, .1), nrow = 1, ncol = 2)
|b2| < -matrix(runif(1, -.1, .1), nrow = 1, ncol = 1)
55
56 res <- NULL
  for(i in 1:100000){
57
    if(i %% 1000 == 0){
58
      res <- c(res, MSE())
59
60
61
    # forward propagation
62
63
    j <- sample(1:nrow(dat),1)</pre>
64
    q0 <- dat[j,1]
65
    z1 \leftarrow w1 \% \% q0 + b1
66
    q1 \leftarrow as.matrix(apply(z1,1,h), nrow = 2, ncol = 1)
67
    z2 \leftarrow w2 \% *\% q1 + b2
68
69
    # backward propagation
70
71
    dz2 \leftarrow -2 * (dat[j,2] - z2)
72
    dq1 <- t(w2) %*% dz2
73
    dz1 \leftarrow dq1 * hprime(z1)
74
    dw2 \leftarrow dz2 %*% t(q1)
75
    db2 \leftarrow dz2
76
    dw1 < - dz1 % * % t(q0)
77
    db1 <- dz1
78
79
    # parameter updating
80
81
    w2 \leftarrow w2 - gamma * dw2
82
    b2 <- b2 - gamma * db2
83
    w1 <- w1 - gamma * dw1
    b1 <- b1 - gamma * db1
85
86
87 | plot(res, type = "1")
89 plot (dat)
points(dat[,1],lapply(dat[,1],yhat),col="red")
```

# 14 Theory

#### 14.1 Linear Model

The general formula for a linear model is:

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \dots + \beta_p x_p + \epsilon$$

In matrix form, the same model can be expressed as:

$$\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\epsilon}$$

Where:

- y: Dependent variable (response).**X**: Design matrix of predictors, of size  $n \times (p+1)$ , where the first column is typically ones (for the intercept).
- $\beta_0$ : Intercept.
- $\beta_1, \beta_2, \dots, \beta_p$ : Coefficients for the independent variables  $x_1, x_2, \dots, x_p$ .  $\beta$ : Coefficient vector, including the intercept, of size  $(p+1) \times 1$ .
- $\epsilon$ : Error term, capturing the variation not explained by the model.  $\epsilon$ : Vector of error terms, of size  $n \times 1$ .
- **y**: Vector of responses, of size  $n \times 1$ .

### 14.2 Logistic Regression

The probability of the positive class is given by:

$$p = \frac{1}{1 + e^{-(\beta_0 + \beta_1 x_1 + \dots + \beta_p x_p)}}$$

In terms of the logit function:

$$logit(p) = ln\left(\frac{p}{1-p}\right) = \beta_0 + \beta_1 x_1 + \dots + \beta_p x_p$$

In matrix form:

$$p = \frac{1}{1 + e^{-\mathbf{X}\beta}}$$

### 14.3 Bias Variance tradeoff

#### 1. Bias:

- Bias refers to the error introduced by approximating a real-world problem, which may be highly complex, by a simplified model.
- High bias implies that the model is too simplistic and may not capture the underlying patterns in the data.
- This can lead to systematic errors, and the model may consistently underperform.

#### 2. Variance:

- Variance refers to the model's sensitivity to small fluctuations or noise in the training data.
- High variance implies that the model is too complex and has learned to capture the noise in the training data rather than the underlying patterns.
- This can lead to overfitting, where the model performs well on the training data but poorly on new, unseen data.

The tradeoff arises because increasing the complexity of a model typically reduces bias but increases variance, and vice versa. The goal is to find the right balance that minimizes the overall prediction error on new, unseen data. Bayes variance: the more leaves, the higher variance but smaller bias. If our model is too simple and has very few parameters then it may have high bias and low variance. On the other hand if our model has large number of parameters then it's going to have high variance and low bias. So we need to find the right/good balance without overfitting and underfitting the data.

Bias is the difference between the average prediction of our model and the correct value which we are trying to predict. Model with high bias pays very little attention to the training data and oversimplifies the model. It always leads to high error on training and test data.

Variance is the variability of model prediction for a given data point or a value which tells us spread of our data. Model with high variance pays a lot of attention to training data and does not generalize on the data which it hasn't seen before. As a result, such models perform very well on training data but has high error rates on test data.

### 14.4 Precision and recall

The formulas for Precision and Recall are as follows:

Precision measures the proportion of positive predictions that were actually correct. It focuses on the accuracy of the positive predictions made by the model.

$$\mbox{Precision} = \frac{\mbox{True Positives}}{\mbox{True Positives} + \mbox{False Positives}}$$

Recall measures the proportion of actual positive instances that were correctly identified by the model. It focuses on the completeness of the positive predictions.

$$\mbox{Recall} = \frac{\mbox{True Positives}}{\mbox{True Positives} + \mbox{False Negatives}}$$

# 14.5 F1 score

Good measure of performance if the dataset is imbalanced. Measures how good the model is at guessing "positive". The formula for the F1 score is given by:

$$F_1 = 2 \cdot \frac{\mathsf{Precision} \cdot \mathsf{Recall}}{\mathsf{Precision} + \mathsf{Recall}}$$