

Compulsory Exercise 3

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Problem 1 (9P)

In compulsory exercise 2 we used the `College` data from the `ISLR` library, where we wanted to predict `Outstate`.

```
library(ISLR)
library(keras)
set.seed(1)
College$Private = as.numeric(College$Private)
train.ind = sample(1:nrow(College), 0.5 * nrow(College))
college.train = College[train.ind, ]
college.test = College[-train.ind, ]
str(College)
```

```
## 'data.frame':    777 obs. of  18 variables:
## $ Private      : num  2 2 2 2 2 2 2 2 2 2 ...
## $ Apps         : num  1660 2186 1428 417 193 ...
## $ Accept       : num  1232 1924 1097 349 146 ...
## $ Enroll       : num  721 512 336 137 55 158 103 489 227 172 ...
## $ Top10perc    : num  23 16 22 60 16 38 17 37 30 21 ...
## $ Top25perc    : num  52 29 50 89 44 62 45 68 63 44 ...
## $ F.Undergrad  : num  2885 2683 1036 510 249 ...
## $ P.Undergrad  : num  537 1227 99 63 869 ...
## $ Outstate     : num  7440 12280 11250 12960 7560 ...
## $ Room.Board   : num  3300 6450 3750 5450 4120 ...
## $ Books        : num  450 750 400 450 800 500 500 450 300 660 ...
## $ Personal     : num  2200 1500 1165 875 1500 ...
## $ PhD          : num  70 29 53 92 76 67 90 89 79 40 ...
## $ Terminal     : num  78 30 66 97 72 73 93 100 84 41 ...
## $ S.F.Ratio    : num  18.1 12.2 12.9 7.7 11.9 9.4 11.5 13.7 11.3 11.5 ...
## $ perc.alumni  : num  12 16 30 37 2 11 26 37 23 15 ...
## $ Expend       : num  7041 10527 8735 19016 10922 ...
## $ Grad.Rate    : num  60 56 54 59 15 55 63 73 80 52 ...
```

The task here is to fit densely connected neural networks using the package `keras` in order to predict `Outstate`.

a) (2P)

```
y.train = college.train$Outstate
y.test = college.test$Outstate
```

```

x.train <- subset(college.train, select = -c(Outstate))
x.test <- subset(college.test, select = -c(Outstate))

mean <- apply(x.train, 2, mean)
std <- apply(x.train, 2, sd)

x.train <- as.array(scale(x.train, center = mean, scale = std))
x.test <- as.array(scale(x.test, center = mean, scale = std))

str(x.train)

##  num [1:388, 1:17] -1.618 0.616 -1.618 0.616 0.616 ...
##  - attr(*, "dimnames")=List of 2
##    ..$ : chr [1:388] "University of Southern Colorado" "College of Notre Dame" "Salisbury State Univer
##    ..$ : chr [1:17] "Private" "Apps" "Accept" "Enroll" ...
##  - attr(*, "scaled:center")= Named num [1:17] 1.72 3220.68 2117.89 796.6 27.82 ...
##    ..- attr(*, "names")= chr [1:17] "Private" "Apps" "Accept" "Enroll" ...
##  - attr(*, "scaled:scale")= Named num [1:17] 0.447 4282.902 2615.659 914.802 17.649 ...
##    ..- attr(*, "names")= chr [1:17] "Private" "Apps" "Accept" "Enroll" ...

```

b) (2P)

Write down the equation which describes a network that predicts `Outstate` with 2 hidden layers and `relu` activation function with 64 units each. What activation function will you choose for the output layer?

$$y_c(\mathbf{x}) = \phi_o(\beta_{0c} + \sum_{m=1}^4 \beta_{mc} z_m) = \phi_o(\beta_{0c} + \sum_{m=1}^4 \beta_{mc} \phi_{h*}(\gamma_{0m} + \sum_{l=1}^4 \gamma_{lm} \phi_h(\alpha_{0l} + \sum_{j=1}^4 \alpha_{jl} x_j))). \quad (1)$$

c) (3P)

- (i) Train the network from b) for the training data using the library `keras`; use 20% of the training data as your validation subset (1P).
- (ii) Plot the training and validation error as a function of the epochs (1P).
- (iii) Report the MSE of the test set and compare it with methods that you used in Compulsory 2 (1P).

Hints:

- Use the `optimizer = "rmsprop"`, `epochs=300` and `batch_size=8`
- Make sure that you are caching the results (`cache=TRUE` in the knitr options), because fitting the models takes some time, and you do not want to repeat this each time you compile your file.

```

# Defining the model
model <- keras_model_sequential() %>%
  layer_dense(units = 64, activation = "relu", input_shape = c(17)) %>%
  layer_dense(units = 64, activation = "relu") %>%
  layer_dense(units = 1, activation = "linear")

# Compile the model
model %>% compile(optimizer = "rmsprop", loss = "mse")

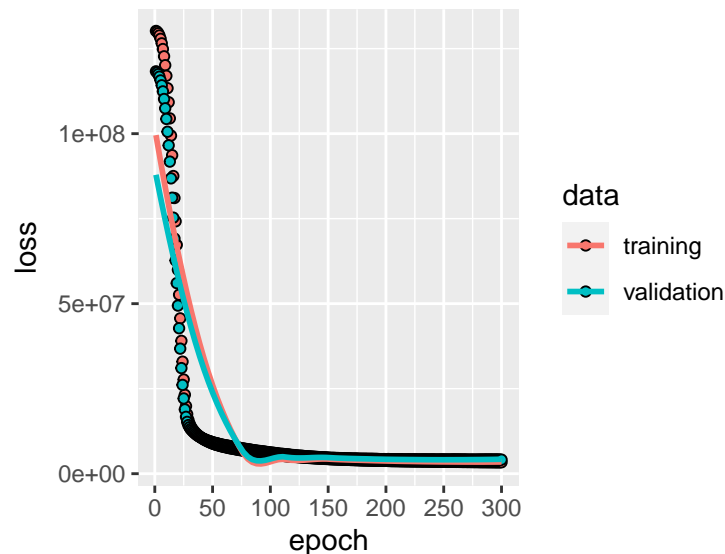
# Fitting the model

```

```
history <- model %>% fit(x.train, y.train, epochs = 300, batch_size = 8, validation_split = 0.2)
str(history)
```

```
## List of 2
## $ params :List of 7
## ..$ batch_size : int 8
## ..$ epochs      : int 300
## ..$ steps       : num 39
## ..$ samples     : int 310
## ..$ verbose     : int 0
## ..$ do_validation: logi TRUE
## ..$ metrics      : chr [1:2] "loss" "val_loss"
## $ metrics:List of 2
## ..$ loss : num [1:300] 1.30e+08 1.30e+08 1.30e+08 1.29e+08 1.28e+08 ...
## ..$ val_loss: num [1:300] 1.18e+08 1.18e+08 1.18e+08 1.17e+08 1.16e+08 ...
## - attr(*, "class")= chr "keras_training_history"
```

```
plot(history)
```



```
# Evaluation and prediction
score <- model %>% evaluate(x.test, y.test)
str(model)
```

```
## Model
## Model: "sequential"
## -----
## Layer (type)                Output Shape          Param #
## -----
## dense (Dense)                (None, 64)            1152
## -----
## dense_1 (Dense)              (None, 64)            4160
## -----
## dense_2 (Dense)              (None, 1)             65
## -----
## Total params: 5,377
## Trainable params: 5,377
## Non-trainable params: 0
```

d) (2P)

Apply one of the regularization techniques you heard about in the course (easiest to use dropout or weight decay with L1/L2 norms). Does this improve the performance of the network? Optional: You might try your own network architecture.

Problem 2 (10P)

In this problem, we will use a real dataset of individuals with the Covid-19 infection. The data were downloaded from <https://www.kaggle.com/shirmani/characteristics-corona-patients> on 30. March 2020, and have only been cleaned for the purpose of this exercise. The dataset consists of 2010 individuals and four columns,

- **deceased**: if the person died of corona (1:yes, 0:no)
- **sex**: male/female
- **age**: age of person (ranging from 2 years to 99 years old)
- **country**: which country the person is from (France, Japan, Korea or Indonesia)

Note that the conclusions we will draw here are probably not scientifically valid, because we do not have enough information about how data were collected.

Load your data into R using the following code:

```
# id <- "1CA1RPRYqU9oTIaHfSroitnWrI6WpUeBw" # google file ID
# d.corona <- read.csv(sprintf("https://docs.google.com/uc?id=%s&export=download", id), header=T)
```

a) Inspecting your data (1P)

Inspect the data by reporting **tables** for

- the number of deceased for each country,
- the number of deceased for each sex, and
- for each country: the number of deceased, separate for each sex.

b) Multiple choice (2P)

Answer the following multiple choice questions by using the data above to model the probability of deceased as a function of **sex**, **age** and **country** (with France as reference level; no interactions).

Which of the following statements are true, which false?

- Country is not a relevant variable in the model.
- The slope for indonesia has a large p -value, which shows that we should remove the Indonesian population from the model, as they do not fit the model as well as the Japanese population.
- Increasing the age by 10 years, $x_{age}^* = x_{age} + 10$, and holding all other covariates constant, the odds ratio to die increases by a factor of 1.97.
- The probability to die is approximately 3.12 larger for males than for females.

c) (1P)

Create a plot of probabilities to die of coronavirus as a function of age, separately for the two sexes and each country.

Hints:

- Make one plot and add lines for each country/sex.
- A useful function to generate gridded data for prediction is `expand.grid()`. For example `newdata = expand.grid(sex="male", age= seq(20,100,1) ,country="France")` generates a grid for males in France over a range of ages between 20 and 100.

d) (3P)

As a statistician working on these data, you are asked the following questions:

- (i) Have males generally a higher probability to die of coronavirus than females?
- (ii) Is age a greater risk factor for males than for females?
- (iii) Is age a greater risk factor for the French population than for the Korean population?

Answer the questions by fitting appropriate models (1P each).

e) Interpret your model (1P)

According to your model fitted in part b), it looks like the French population is at a much higher risk of dying from Covid-19 than the other countries. Do you trust this result? How could it be influenced by the way the data were collected?

f) Multiple choice (2P)

Which of the following statements are true, which false?

Consider the classification tree below to answer:

- (i) The probability of dying (`deceased = 1`) is about 0.46 for a French person with age above 91.
- (ii) Age seems to be a more important predictor for mortality than sex.

Consider the LDA code and output below:

- (iii) The “null rate” for misclassification is 2.24%, because this is the proportion of deaths among all cases in the dataset. No classifier should have a higher misclassification rate.
- (iv) LDA is not a very useful method for this dataset, among other reasons because it does not estimate probabilities, but also because the misclassification error is too high.

```
# library(MASS)
# table(predict=predict(lda(deceased ~ age + sex + country, data=d.corona))$class, true = d.corona$deceased)
```

Problem 3 (14P)

The `d.support` dataset (source *F. E. Harrell, Regression Modeling Strategies*) contains the total hospital costs of 9105 patients with certain diseases in American hospitals between 1989 and 1991. The different variables are

Variable	Meaning
<code>totcst</code>	Total costs
<code>age</code>	Age of the patients
<code>'dzgroup</code>	' Disease group
<code>num.co</code>	Number of co-morbidities
<code>edu</code>	Years of education
<code>scoma</code>	Measure for Glasgow coma scale
<code>income</code>	Income
<code>race</code>	Rasse
<code>meanbp</code>	Mean blood pressure
<code>hrt</code>	Heart rate
<code>resp</code>	Respiratory frequency
<code>temp</code>	Body temperature
<code>pafi</code>	PaO2/FiO2 proportion (blood-gas mixture)

Data are loaded as follows (and we reduce the number of patients to the 4960 complete cases with total costs larger than 0):

```
# id <- "1heRtzi8vBoBGMaM2-ivBQI5Ki3HgJTm0" # google file ID
# d.support <- read.csv(sprintf("https://docs.google.com/uc?id=%s&export=download", id),header=T)
# # We only look at complete cases
# d.support <- d.support[complete.cases(d.support),]
# d.support <- d.support[d.support$totcst >0 ,]
```

We would like to build models that help us to understand which predictors are mostly driving the total cost, but also models for prediction.

a) (1P)

Before we start analysing the data, visualize the distributions of all continuous or integer variables with histograms. Suggest a transformation for the response variable `totcst` (hint: it is a *standard transformation* that we have used earlier in the course). Important: **you should fit all models with the transformed version of the response variable `totcst` from now on. Leave all other variables untransformed.**

b) (3P)

Fit a multiple linear regression model with the six covariates `age`, `temp`, `edu`, `resp`, `num.co` and `dzgroup` and the (transformed version of the) response `totcst`.

- How much/by which factor are the total costs expected to change when a patient's age increases by 10 years, given that all other characteristics of the patient are the same? Use the transformed response to fit the model, but report the result on the original (back-transformed) scale of the response. (1P)
- Do a residual analysis using the Tukey-Anscombe plot and the QQ-diagram. Are the assumptions fulfilled? (1P)
- Does the effect of age depend on the disease group? Do a formal test and report the p -value. (1P)

c) (3P)

In order to build a more robust model for inference and prediction of the total costs, continue using ridge regression. Create a training set with 80% of the data and a test set with the remaining 20% (1P). Run

cross-validation to find the largest value of λ such that the error is within 1 standard error of the smallest λ (1P). Report the test MSE of the ridge regression where you used the respective λ (1P).

Be careful: we still use the same transformation for the response as in b) – you should report the MSE using the transformed version of `totcst` (i.e., do **not back-transform** the MSE to the original scale).

R-hints:

```
set.seed(12345)
# train.ind = sample(1:nrow(d.support), ... *nrow(d.support))
# d.support.train = d.support[... ,]
# d.support.test = d.support[... ,]
```

d) (3P)

Now assume that our sole aim is prediction. In the course you heard about *partial least squares (PLS)*. It is a smart approach that uses the principal component regression idea, but finds the components that are best correlated with the response.

Proceed as follows:

- (i) Run a PLS regression (don't forget to scale the variables, `scale=TRUE`) (1P).
- (ii) Choose an optimal number of principal components (PCs) using cross-validation (1P).
- (iii) Report the MSE of the test set when using the respective set of PCs and compare to the result from ridge regression. Conclusion? (1P)

e) (4P)

Now choose two other methods that you know from the course and try to build models with even lower test MSEs than those found so far (imagine that this is a competition where the lowest test MSE wins). Use the same training and test dataset as generated above. And remember that we are still *always* working with the transformed version of the response variable (`totcst`). In particular, use

- (i) One model that involves non-linear transformations of the covariates (e.g., splines, natural splines, polynomials etc) that are combined to a GAM (2P).
- (ii) One model/method based on regression trees (2P).

Very briefly discuss or explain your choices (1-2 sentences each).

Problem 4 (Mixed questions; 6P)

a) 2P

We look at the following cubic regression spline model:

$$Y = \begin{cases} \beta_0 + \beta_1x + \beta_2x^2 + \beta_3x^3 + \epsilon, & \text{if } x \leq 1, \\ \beta_0 + \beta_1x + \beta_2x^2 + \beta_3x^3 + \beta_4(x-1)^3 + \epsilon, & \text{if } 1 < x \leq 2, \\ \beta_0 + \beta_1x + \beta_2x^2 + \beta_3x^3 + \beta_4(x-1)^3 + \beta_5(x-2)^3 + \epsilon, & \text{if } x > 2. \end{cases}$$

Write down the basis functions (1P) and the design matrix (1P) of this model.

b) Multiple choice - 2P

Inference vs prediction: Which of the following methods are suitable when the aim of your analysis is inference?

- (i) Lasso and ridge regression
- (ii) Multiple linear regression with interaction terms
- (iii) Logistic regression
- (iv) Support Vector Machines

c) Multiple choice - 2P

We again look at the Covid-19 dataset from Problem 2 to study some properties of the bootstrap method. Below we estimated the standard errors of the regression coefficients in the logistic regression model with **sex**, **age** and **country** as predictors using 1000 bootstrap iterations (column **std.error**). These standard errors can be compared to those that we obtain by fitting a single logistic regression model using the **glm()** function. Look at the R output below and compare the standard errors that we obtain from these two approaches (note that the **t1*** to **t6*** variables are sorted in the same way as for the **glm()** output).

```
# library(boot)
# boot.fn <- function(data,index){
#   return(coefficients(glm(deceased ~ sex + age + country, family="binomial",data=data,subset=index)))
# }
# boot(d.corona,boot.fn,1000)

# Logistic regression
# r.glm <- glm(deceased ~ sex + age + country, d.corona,family="binomial")
# summary(r.glm)$coef
```

Which of the following statements are true?

- (i) There are large differences between the estimated standard errors, which indicates a problem with the bootstrap.
- (ii) The differences between the estimated standard errors indicate a problem with the assumptions taken about the distribution of the estimated parameters in logistic regression.
- (iii) The **glm** function leads to too small *p*-values for the differences between countries, in particular for the differences between Indonesia and France and between Japan and France.
- (iv) The bootstrap relies on random sampling the same data without replacement.

Problem 5 (Multiple and single choice questions; 11P)

a) Multiple choice - 2P

Which of the following are techniques for regularization?

- (i) Lasso
- (ii) Ridge regression
- (iii) Forward and backward selection
- (iv) Stochastic gradient descent

b) Multiple choice - 2P

Which of the following statements about principal component regression (PCR) and partial least squares (PLS) are correct?

- (i) PCR involves the first principal components that are most correlated with the response.
- (ii) PLS involves the first principal components that are most correlated with the response.
- (iii) The idea in PLS is that we choose the principal components that explain most variation among all covariates.
- (iv) The idea in PCR is that we choose the principal components that explain most variation among all covariates.

c) Single choice - 1P

In ridge regression, we estimate the regression coefficients in a linear regression model by minimizing

$$\sum_{i=1}^n \left(y_i - \beta_0 - \sum_{j=1}^p \beta_j x_{ij} \right)^2 + \lambda \sum_{j=1}^p \beta_j^2 .$$

What happens when we increase λ from 0? Choose the single correct statement:

- (i) The training RSS will steadily decrease.
- (ii) The test RSS will steadily decrease.
- (iii) The test RSS will steadily increase.
- (iv) The bias will steadily increase.
- (v) The variance of the estimator will steadily increase.

d) Single choice - 1P

Which statement about the *curse of dimensionality* is correct?

- (i) It means that we have a bias-variance tradeoff in K -nearest neighbor regression, where large K leads to more bias but less variance for the predictor function.
- (ii) It means that the performance of the K -nearest neighbor classifier gets worse when the number of predictor variables p is large.
- (iii) It means that the K -means clustering algorithm performs bad if the datapoints lie in a high-dimensional space.
- (iv) It means that support vector machines with radial kernel function should be avoided, because radial kernels correspond to infinite-dimensional polynomial boundaries.
- (v) It means that we should never measure too many covariates when we want to do classification.

e) Single choice - 1P

Now assume you have 10 covariates, X_1 to X_{10} , each of them uniformly distributed in the interval $[0, 1]$. To predict a new test observation $(X_1^{(0)}, \dots, X_{10}^{(0)})$ in a K -nearest neighbor (KNN) clustering approach, we use all observations within 20% of the range closest to each of the covariates (that is, in each dimension). Which proportion of available (training) observations can you expect to use for prediction?

- (i) $1.02 \cdot 10^{-7}$
- (ii) $2.0 \cdot 10^{-3}$
- (iii) 0.20
- (iv) 0.04
- (v) 10^{-10}

f) Multiple choice - 2P

This example is taken from a real clinical study by *Ikeda, Matsunaga, Irabu, et al. Using vital signs to diagnose impaired consciousness: cross sectional observational study. BMJ 2002;325:800*. Researchers investigated the use of vital signs as a screening test to identify brain lesions in patients with impaired consciousness. The setting was an emergency department in Japan. The study included 529 consecutive patients that arrived with consciousness. Patients were followed until discharge. The vital signs of systolic and diastolic blood pressure and pulse rate were recorded on arrival. The aim of this study was to find a quick test for assessing whether the newly arrived patient suffered from a brain lesion. While vital signs can be measured immediately, the actual diagnosis of a brain lesion can only be determined on the basis of brain imaging and neurological examination at a later stage, thus the quick measurements of blood pressure and heart rate are important to make a quick assessment. In total, 312 patients (59%) were diagnosed with a brain lesion.

The performance of each vital sign (systolic blood pressure, diastolic blood pressure and heart rate) was separately evaluated as a screening test to quickly diagnose brain lesions. To assess the quality of each of these vital signs, different thresholds were taken successively to discriminate between “negative” and “positive” screening test result. For each vital sign and each threshold the sensitivity and specificity were derived and used to plot a receiver operating characteristic (ROC) curve for the vital sign (Figure 1):

Which of the following statements are true?

- (i) The value of 1-specificity represents the proportion of patients without a diagnosed brain lesion identified as positive on screening.
- (ii) When we use different cut-offs, sensitivity increases at the cost of lower specificity, and vice versa.
- (iii) A perfect diagnostic test has an AUC of 0.5.
- (iv) The vital sign that is most suitable to distinguish between patients with and without brain lesion is systolic blood pressure.

g) Multiple choice - 2P

We study the `decathlon2` dataset from the `factoextra` package in R, where Athletes' performance during a sporting meeting was recorded. We look at 23 athletes and the results from the 10 disciplines in two competitions. Some rows of the dataset are displayed here:

```
# decathlon2.active[c(1,3,4),]
```

From a principal component analysis we obtain the biplot given in Figure 2.

Which of the following statements are true, which false?

- (i) The athlete named CLAY seems to be one of the fastest 1500m runners.
- (ii) Athletes that are good in 100m tend to be also good in long jump.
- (iii) The first principal component has the highest loadings for 100m and long jump.
- (iv) 110m hurdle has a very small loading for PC2.