## Machine Learning: Project

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### 1 Introduction

In the world today there are many methods that fall under the category of "Machine Learning", some being very similar and some very different. All of them share in common that they use data to produce a model that can be used to tell us something on previously unseen data. When successful this is very appealing in today's society where we got lots of data on situations where there is likely to be a underlying pattern, another requirement for learning. There is broad agreement that Machine Learning is a good way to make prediction and classification models, and often the only computational feasible way to do so. This makes it a task to decide which method in machine learning to choose for a given problem. The answer is not always the same and several methods can be good, but for different reasons. In this report I will utilize several of the most used machine learning algorithms and show how they perform on classifying images of handwritten digits.

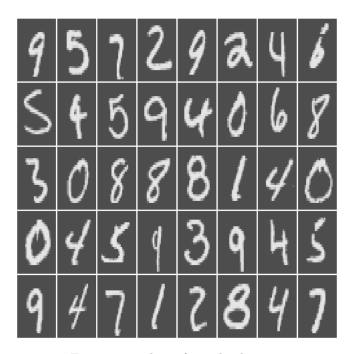


Figure 1: 40 digits from the data set

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### 2 Theory

### 2.1 Tree based methods

#### 2.1.1 Classification tree

Classification tree is maybe the method in Machine Learning which is easiest to interpret due to it's intuitive construction and clear visualization. The method uses a greedy approach using recursive binary splitting to structure a tree that can classify input based on it's variables. The goal of the classification tree is to classify a set of data as best as possible while have a low complexity to avoid overfitting. Below we see that one classification tree is not enough to make a great model for the problem, but combining many of them gives us a "Random Forest" which is discussed in subsection 2.1.2. Classification trees also gives a nice visual image of the classification.

#### 2.1.2 Random Forest

Random Forest is as mentioned (subsection 2.1.1) constructed of many classification tree. After such a construction by using a set of training data, new data can be ran through the "forest". The new data is classified with the label the majority of trees labelled it. The trees in other words "vote" for the best classification for the data. As in real life, voting makes little difference if all the votes are the same. To avoid making a forest out of n ( $n \in \mathbb{Z}_{>0}$ ) identical trees two aspects are changed when the constructing the classification trees. The first is that the data of size N used in training a tree is sampled from the whole set of N data with replacement. This will on average lead to about  $\frac{2}{3}$  data to be used in creating each tree leaving on average  $\frac{1}{3}$  Out-Of-Bag(OOB) which is used for validation. The OOB-data is critical in making insuring the Random Forest doesn't overfit. The other is that m randomly selected variables are used for each tree, where  $m \ll M$  and M is the whole set of variables in the problem. This gives a rich variety of trees where with enough trees a predictive model can be created.

#### 2.1.3 Bagging

Bagging is basically a Random Forest (see 2.1.2) where the number of variables considered for each split is the whole variable set for the problem. It shares the same behaviour as a Random Forest when sampling out the data and in classifying new data. The big difference is that the variables are not sampled in Bagging, this does that Bagging creates trees that are more correlated than a Random Forest making it more susceptible to dominating features. This will not directly lead to a bad classification model, but it does have the unwanted consequence of making semi-important features of the data to be underestimated in the contribution towards new predictions.

#### 2.1.4 Boosting

Boosting takes a different direction to find make a good predictive tree model than the two previous tree based methods. Instead of growing n trees independently of each other, boosting "grow" trees sequentially moving towards a good model. Boosting is a slowly improving method and often use a small amount of splits for each tree. It also have a shrinkage parameter  $\lambda$  that controls the rate of change in the model. Boosting learns slower, which means more trees are required compared to the random forest or bagging method in order to achieve the same predictive power.

### 2.2 Deep Learning

#### 2.2.1 Artificial Neural Network

Artificial Neural Networks got the name because it to some degree mimic the behaviour of neurons in the brain. It does so by connecting nodes (neurons) together with weights (synapses) connecting them. In the nodes that are not input nodes there is an activation function altering the values that are brought to the nodes through weights from other nodes. The activation is one of several factors

that can be tuned when creating a neural network. The relationship between the weights can be seen the equation for calculating the nodes in the hidden layers

$$x_j = \theta(s_j^{(l)}) = \theta(\sum_{i=0}^{d^{(l-1)}} w_{ij}^{(l)} x_i^{(l-1)})$$
(1)

, where l indicate the layer, i the input, j the output and  $\theta$  the activation function. Other factors that can be altered is the size of the hidden layers and the amount of nodes in each layer of the network. This method learn through updating the weights of using stochastic gradient descent(SGD). This is applied through the backpropagation algorithm. This method takes run data forward through the network and the backpropagates back updating the weights based on the error in the output using SGD. The backpropagation will penalize weights that contributed a lot to the error more than those less relevant. Neural networks are exposed to overfitting, but there are several techniques that can be used to decrease the chance of overfitting. Some of these are validation, using dropout in input and/or hidden layers and regularization.

#### 2.2.2 Convolutional Neural Network

Convolutional Neural Network(CNN) is in theory a great method to classify digits because of it's great reputation in performance for problems that can be represented as an "image". This because it CNN's works by recognizing patterns in problems that can be represented as by matrix or tensor(multidimensional matrix) in the computer. It does so by using convolutional, pooling and voting layers. Multiple of these can be stacked to create a very a precise model for recognising images.

### 2.3 Support Vector Machines

Support vector machines(SVM) is a powerful methods used either to classify data or in regression analysis. SVM tries as big a margin as possible between different categories. This is strived after to decrease the number of possible dichotomies. SVM's are powerful because of the ability to classify non-linearly separable data by transforming/mapping the data to higher dimensional space, using the inner product to classify the data. SVM's works well when using feature extraction like Principal Component Analysis(PCA) to reduce the amount of features to evaluate.

#### 2.4 K-Nearest Neighbours

K-Nearest Neighbours(KNN) is a easy-to interpret and maybe one of the simplest methods in Machine Learning. The method does exactly as the name suggest; find the k nearest neighbours, where k is a number and labels each data point based on those neighbours. KNN doesn't learn in the fashion random forest or neural net does, it rather just classifies data using the most similar known data. The method is attractive because of it's easy interpretability, fast execution time and often precise classification and regression power. There are many variants of the method where the distance measure and way to handle draws differ. KNN's with k=1 have a special property that the hyperplane of the data points will be partitioned into an equal amount of partitions as there are data points. Each partition consist of all point in the plane closer a specific data point than any other. The amount of neighbours k can have a great impact on the accuracy of the classification. In some cases a low k will be preferable with very separated data, in other case it can lead to overfitting due to higher influence from noise and outliers. Because a even number of categories is classified in this problem the concerns of draws is present when using a even k. This makes a odd k more attractive in this problem.

### 3 Methods

As mentioned in the introduction different methods in Machine Learning will be used to try to build a model that manage to correctly classify if pictures of handwritten numbers are odd or even. This

would be simple if the method should only classify previously seen data, the challenge arise when asked to make a model that is highly accurately in classifying new data. The goal is to make a a model that trains without overfitting on the in-sample data in order to well estimate the out-of-sample data. In the methods below this is done by using different techniques like cross-validation, validation-set and regularization. In all of the methods the data is classified into the numbers 0-9. Another approach would be to classify the number's directly into either a odd or even category. The reason for using all ten numbers is mainly because this makes the extension of the use purpose a lot smoother, f.ex. to also classify if the number is a prime number or not.

#### 3.1 Tree based methods

#### 3.1.1 Classification tree

The regression tree is implemented with the R-package "tree" which has a function with the same name. In this method the feature being classified is set and which variables it should consider in the splits. When classifying digits many of the pixels will have a low variance nearing the borders, these will most likely not be used in any of the splits. To reduce unnecessary computation, columns with variance close to zero are removed. Other parameters that can be altered is to control when the tree should stop splitting. Here the minimum development is set as stopping metric. This is measured by calculating the reduction in Residual Sum of Squared - RSS given by:

$$RSS = \sum_{i=1}^{n} (y_i - f(x_i))^2$$
 (2)

, where i is a data point in the set of n data points. After a tree has been fitted it might be that the number of terminal nodes are too high and a lower amount of nodes can classify the data as well meanwhile reducing overfitting.

#### 3.1.2 Random Forest

To create a random forest to classify digits the package "randomForest" is used. The function with the same name as the library is used both for creating the random forest and the bagging. These parameters was used when running the random forest:

ntree mtry
$$\frac{1}{3} \times 784 \approx 261$$
Table 1

, where ntree is the amount of trees and mtry the number of variables used in each tree. The fraction  $\frac{1}{3}$  is comes from the default size of the subset of variables in the "randomForest" package. The total number of variables is 784, equal to the amount of pixels in each image.

### 3.1.3 Bagging

The implementation of bagging or bootstrap aggregation is identical to random forest as described above except for mtry. Here the subset of variables is the whole set of variables. This is implemented with the parameters:

$$\begin{array}{c|c}
\text{ntree} & \text{mtry} \\
\hline
500 & 784
\end{array}$$

Table 2

. For both bagging and random forest there is no need to specify a form of validation due to the OOB-estimate for the ut-of-sample error as explain in the theory (2.1.2).

#### 3.1.4 Boosting

For the implementation of boosting, the library "gbm" is used. The function "gbm" is very similar in use compared to other tree-functions, but "gbm" has an advantage in being parallelized speeding up big runs. Boosting has the possibility to overfit and therefore the built-in validation metric is set to 10-fold cross-validation. When running boosting these parameter were set to:

n	itree	distribution	interaction.depth	${\tt shrinkage}$	cv.folds	n.cores
1	10000	"multinomial"	2	0.01	10	4

Table 3

. Here the distribution is set to be multinomial in order to make the boosting method handle the 10 categories for the digits. Here it is possible to note if one directly classify digits as odd or even a binomial distribution would possible to use. The interaction depth is set to two in order to keep the trees small with a  $\lambda$  (shrinkage) of 0.01 slowly improving the performance. The last two parameters gives the number of cross-validation folds and cores to use.

### 3.2 Deep Learning

#### 3.2.1 Artificial Neural Network

For the neural network a "h2o"-grid is run in order to find a good set up for the neural network. "h2o" is a 'R' package originally written in 'Java'. This is a powerful tool which has a deep learning function that suits very well for setting up a neural network. The function runs in parallel and can be used with the grid-function in "h2o". This makes it easier to build and compare several neural nets to find the best parameters for the network of a specific problem. To avoid overfitting 10-fold cross-validation is activated for all the models. The different parameters below were combined producing 40 neural networks:

Hyper parameter	value(s)
hidden	(100, 100), (150, 150), (540, 320), (100, 100, 100), (540, 320, 100)
<pre>input_dropout_ratio</pre>	0, 0.2
epochs	20
activation	"Rectifier", "RectifierWithDropout"
11	0, 1.4e - 5

Table 4

. The hidden layers are controlled by the "hidden"-parameter with the number of layers being the length of the vector and the size of each given by the number. "l1" regularization and drop-out in the input layers are given tried to deal with overfitting. To keep the neural network within feasible range 20 epochs were used for each model. The two rectifier activation functions for deep learning in "h2o" were in training the models.

### 3.2.2 Convolutional Neural Network

To implement a convolutional neural network the package "mxnet" was used. This is originally written in 'C++' and is available in several programming languages. The network is built by piecing together different layers and activation functions. A validation set were used when tweaking to find the best parameters. 15% of the training data was sampled out and used to estimate the out-of-sample error. Parameters for convolutional neural network with two convolutional layers:

Layer	parameters
convolution_1	$\mathtt{kernel} = 5 \times 5, \mathtt{num\_filter} = 30$
activation_1	"tanh"
pooling_1	$\texttt{pool\_type = "max", kernel} = 2 \times 2$
convolution_2	$\texttt{kernel} = 5 \times 5,  \texttt{num\_filter} = 50$
activation_2	"tanh"
pooling_2	$\texttt{pool\_type = "max", kernel} = 2 \times 2$
flatten	
fully_1	${\tt num\_hidden} = 500$
activation	"Rectifier Linear Unit - relu"
fully_2	${\tt num\_hidden} = 40$

Table 5: Parameters Covolutional Neural Network

. The full network is constructed of two convolution layers, two pooling layer, three activation function, one flatten layer and two fully connected layers. The network can be trained using either the CPU or a GPU, here the CPU is used due to lack of GPU.

### 3.3 Support vector machines/PCA

Before using support vector machines with the package "e1071" feature selection was performed on the data to reduce the number of features and thereby the complexity of the classification problem. To extract the features, principal component analysis was applied. PCA takes the original features and combines them making new features that tries to describe as much of the variance in the data as possible. After running PCA on the features the p most important PCA features is selected. PCA returns the most describing features in decreasing importance. Running the SVM with different amounts of PCA features yield different cross-validation error which is used to find the best number of principal components to use with SVM. The parameters used for support vector machines:

Parameter	<pre>value(s)</pre>
amount PCA	$p \in [1, 2,, 30]$
cost	$cost \in [0.5, 1, 1.5,, 10]$
	Table 6

### 3.4 K-Nearest Neighbours

The library used for the knn implementation is "Class". The only parameters one can really change in a k-nearest-neighbour model is the k number of neighbour and the distance metric used to obtain them. The standard euclidean distance is used, making it only a challenge to find the best k. This is done by performing cross-validation on the dataset for each k. This is implemented using a parallelized cross-validation function to speed it up. To experiment a bit with the effect of normalizing the data both a normalized and untouched dataset is run through KNN.

### 4 Results

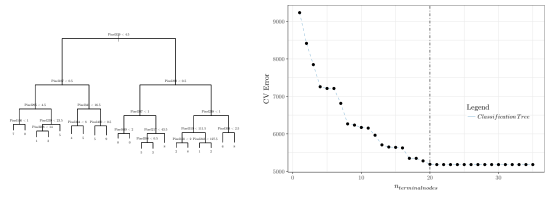
The result for each model are presented with a confusion matrix showing how well each model classified the different digits. With the figures are some plot justifying or illustrating the choices made when making the models. To decide which of the models, 20% of the training data is separated out and used to estimate the out-of-sample error when comparing the models. I will refer to this as the "test set" and the given test set of unlabelled data as the "unclassified set". The model with the lowest error is in used to train on the whole training data and used to predict on the unclassified data. This prediction is stored as .csv-file with the predicted digit for each image and if it is a even

or odd number. It is tempting to use the separated test set when training the individual model, but this would be snooping in the data, leading to contamination of the results.

### 4.1 Tree based methods

#### 4.1.1 Classification tree

The is very fast to make given that it is only one tree. As seen in the plot over the CV Error below there is little to gain in reduction of the error after 20 terminal nodes (NB! not the y-scale starts at 6000). The trees is pruned down to 20 terminal nodes before predicting on the created test set.



- (a)  $E_{CV}$  for different amounts of leaf nodes.
- (b) See that 20 terminal nodes is enough.

Figure 2: Classification tree

After the tree is pruned, the test set in run through the model and each data point is giving a predicted label. This is compared to the real value which gives an error of 37.1%. This is clearly the worst of all the models, but still not that bad thinking it is made using only one tree. The classification tree give a visual of how the tree models classify the data, but is to simple to make a good prediction.

	0	1	2	3	4	5	6	7	8	9	Error	Rate
				<u> </u>	4					9		
0	31	0	5	1	1	3	4	1	0	1	0.340	16/47
1	0	44	1	2	0	2	3	0	1	0	0.170	9/53
2	0	3	32	2	1	0	0	0	2	0	0.200	8/40
3	0	1	1	26	1	0	1	1	0	3	0.235	8/34
4	0	0	0	0	26	0	10	0	1	1	0.316	12/38
5	4	2	2	12	9	31	6	10	3	7	0.640	55/86
6	0	1	3	1	$^{2}$	1	23	0	0	2	0.303	10/33
7	1	0	1	2	0	0	2	34	0	5	0.244	11/45
8	0	2	5	3	6	5	6	0	39	3	0.435	30/69
9	3	0	2	2	7	3	2	1	6	27	0.491	26/53
Total	8	9	20	25	27	14	34	13	13	22	0.371	185/498

Table 7: Confusion Matrix: Classification tree

#### 4.1.2 Random Forest and Bagging

Since the random forest and bagging are very similar in how they are build up the results from these two models are presented in the same plot and confusion matrices together. As mentioned in the theory 2.1.2 we expect random forest to perform better than bagging due to it having less correlated trees. This is clearly the case when plotting the misclassification error function for each method. We see a very similar behaviour in each curve. Bagging starts with a lower error and keeps this up to around 25-30 trees when random forest passes it. This further indicates that the randomness when creating the subset of variables for the random forest in the start will produce worse results in the beginning, but quickly will be better when enough trees are created. This is because after some time there will be on average trees splitting on thee most important variables which creates more uncorrelated trees and a better performance than bagging. Another interesting feature of the models, is that after around 150 the improvement stalls and the solutions stabilizes. This is the limit of the models and increasing the number of trees only contributes to a longer wait when producing the models. This also show how the OOB-data for each tree keeps the models from overfitting.

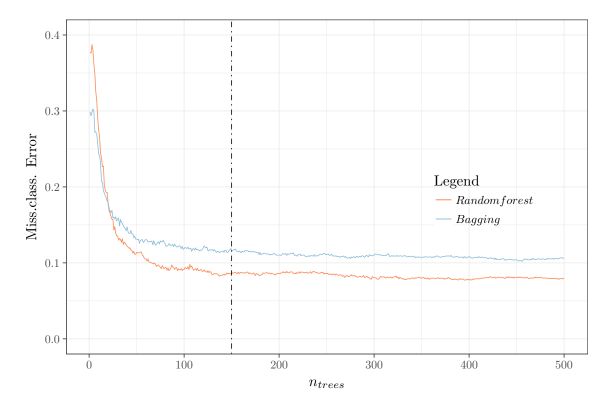


Figure 3: Bagging and Random Forest misclassification over 500 trees.

Below is the confusion matrices for bagging and random forest. As seen in the "Error" tab random forest yield an error of 8.1% while bagging give an error of 10.3%. This makes both of the models a clear improvement of the single classification tree with random forest being the superior over bagging.

	0	1	2	3	4	5	6	7	8	9	Error	Rate
0	46	0	0	1	0	0	0	0	0	0	0.021	1/47
1	0	42	0	0	0	1	0	1	0	0	0.045	2/44
2	0	2	48	0	0	0	0	1	0	0	0.059	3/51
3	0	0	0	41	0	1	0	0	3	0	0.089	4/45
4	0	1	0	0	52	1	1	4	0	1	0.133	8/60
5	0	0	0	4	0	42	1	0	1	0	0.125	6/48
6	0	0	0	0	1	0	47	0	2	0	0.060	3/50
7	0	1	0	0	0	0	0	48	0	2	0.059	3/51
8	1	2	0	1	0	0	0	0	42	1	0.106	5/47
9	0	0	0	1	2	1	0	0	1	48	0.094	5/53
Total	1	6	0	7	3	4	2	6	7	4	0.081	40/496

Table 8: Confusion matrix: Random Forest

	0	1	2	3	4	5	6	7	8	9	Error	Rate
0	47	0	0	2	0	3	0	0	0	0	0.096	5/52
1	0	42	0	0	0	0	0	0	0	0	0.000	0/42
2	0	2	46	0	0	0	0	1	0	0	0.061	3/49
3	0	1	0	41	0	2	0	1	2	0	0.128	6/47
4	0	1	0	0	49	1	1	2	0	2	0.125	7/56
5	0	0	0	3	0	38	1	0	1	0	0.116	5/43
6	0	0	1	0	1	1	47	0	2	1	0.113	6/53
7	0	0	0	0	0	0	0	47	0	3	0.060	3/50
8	0	2	1	1	1	0	0	2	43	1	0.157	8/51
9	0	0	0	1	4	1	0	1	1	45	0.151	8/53
Total	0	6	2	7	6	8	2	7	6	7	0.103	51/496

Table 9: Confusion matrix: Bagging

### 4.1.3 Boosting

From the plot below showing the cv-error over the number of trees made. We see that it decrease quickly in the start before flattening and after around 3000 trees start to increase. When the error starts increasing the boosting model begins to overfit. The best number of trees meaning the amount that gives the lowest cv-error is used to predict on the test set. This number is as pointed out in the plot  $\approx 2900$  trees.

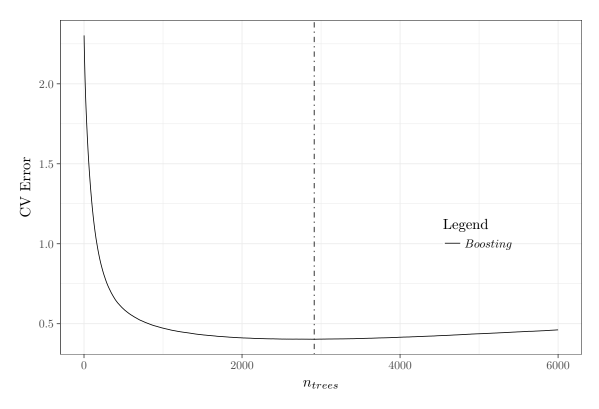


Figure 4: Boosting

Using the best number of trees to predict on the test set giving an error of 11.9%. This means it performes worse than both bagging and random forest, beating single classification tree.

	0	1	2	3	4	5	6	7	8	9	Error	Rate
0	45	0	0	1	0	1	2	0	2	0	0.118	6/51
1	0	41	0	0	0	1	0	1	0	0	0.047	2/43
2	1	1	43	1	0	1	0	2	1	0	0.140	7/50
3	0	1	2	41	0	0	0	1	1	0	0.109	5/46
4	0	1	0	0	46	1	0	4	0	2	0.148	8/54
5	0	0	0	3	0	40	1	0	1	0	0.111	5/45
6	0	0	1	1	2	1	45	0	0	0	0.100	5/50
7	1	0	0	0	0	0	0	43	0	1	0.044	2/45
8	0	3	2	1	2	0	1	2	44	0	0.200	11/55
9	0	1	0	0	5	1	0	1	0	49	0.140	8/57
Total	2	7	5	7	9	6	4	11	5	3	0.119	59/496

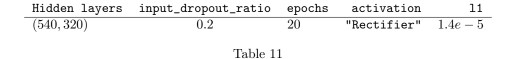
Table 10: Confusion matrix: Boosting

### 4.2 Deep Learning

Below the results for Artificial and Convolutional Neural Network showcasing the difference between them by pure performance. This clearly indicate that they are two distinct methods that share some features.

#### 4.2.1 Artificial Neural Network

The artificial neural network(ANN) was set up by running a grid with different networks as given by the parameter overview in 3.1.2. This is a way to find a good set-up for the neural net by basically just trial and error. Below is a plot where the misclassification error of the models is displayed. There are two lines where the slowly increasing one is the 10-fold cv-error and the oscillating one is the error on the test set. Here we come into the temptation of snooping as mentioned earlier. If I would use the results obtained using the test set I would have been snooping and increase the chance of underestimating the complexity of the problem. So the models are sorted by the cv-error and this is purely used to chose the best model. This even though we can see that there are models that gives lower error on the test set. Set-up for the best neural network based used to predict on the test set:



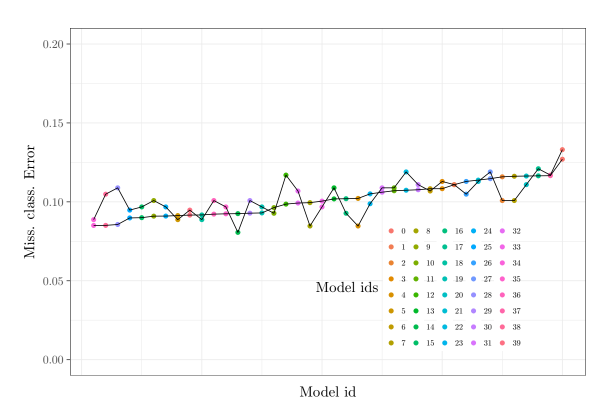


Figure 5: Misclassification error for each model in the grid sorted by the cv-error.

In the confusion matrix below we can see that the final ANN misclassified 9.1% of the digits in the test set. This places it between the bagging and random forest methods performance wise.

	0	1	2	3	4	5	6	7	8	9	Error	Rate
0	42	0	0	0	0	2	0	0	0	0	0.045	2/44
1	0	47	0	0	1	0	0	0	4	0	0.096	5/52
2	2	1	46	$^2$	0	0	1	0	0	0	0.115	6/52
3	0	0	0	42	1	1	0	1	0	0	0.067	3/45
4	0	0	0	0	50	0	0	$^2$	0	2	0.074	4/54
5	1	0	0	$^2$	1	43	0	1	6	1	0.218	12/55
6	1	0	0	1	1	0	48	0	0	0	0.059	3/51
7	1	0	2	0	1	0	0	49	1	3	0.140	8/57
8	0	0	0	1	0	0	0	0	38	0	0.026	1/39
9	0	0	0	0	0	0	0	1	0	46	0.021	1/47
Total	5	1	2	4	5	3	1	5	11	6	0.091	45/496

Table 12: Confusion matrix: Artificial Neural Network

#### 4.2.2 Convolutional Neural Networks

Using a convolutional neural network on this problem showed to be very successful. With a bit of tweaking in the layers show in 3.2.2, the error when validating(validation set 15% of training data) quickly went below 5%. In the plot below both the in-sample and validation error is plotted for the final set-up for the network as described 5. It shows the in-sample error quickly going to zero(overfitting) and the validation error converging to an accuracy around 95% after 10 rounds of training. The convergence of the error sped up when heavily when using "Xavier" distribution to initialize the "mxnet" network.

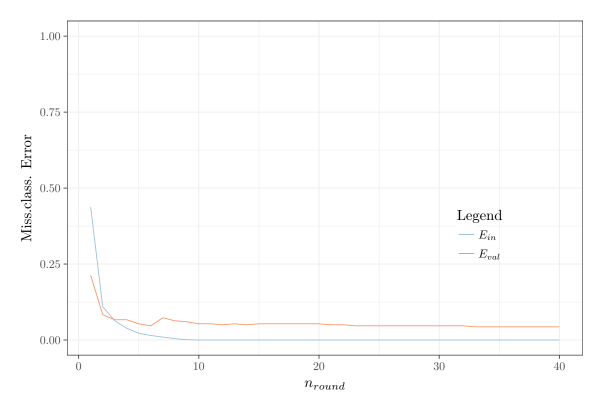


Figure 6:  $E_{in}$  and  $E_{val}$  for training the network over 40 rounds.

Below we see the predictions made on the test set(20%) of the training data) after using taking back the validation set into the training set. The predictions have a misclassification error of 3.6% clearly

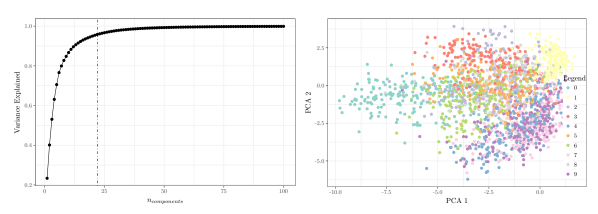
out concurring all the tree models and the ANN. This demonstrates how powerful CNN is when used on image classification.

	0	1	2	3	4	5	6	7	8	9	Error	Rate
0	47	0	0	0	0	0	1	0	0	0	0.021	1/48
1	0	43	0	0	0	0	0	0	0	0	0.000	0/43
2	0	1	47	0	1	0	0	1	1	0	0.078	4/51
3	0	1	0	48	0	0	0	0	0	0	0.020	1/49
4	0	1	0	0	51	1	0	1	0	0	0.056	3/54
5	0	0	0	0	0	44	1	0	0	0	0.022	1/45
6	0	1	0	0	1	0	47	0	0	0	0.041	2/49
7	0	0	1	0	0	0	0	52	0	1	0.037	2/54
8	0	1	0	0	0	1	0	0	48	0	0.040	2/50
9	0	0	0	0	2	0	0	0	0	51	0.038	2/53
Total	0	5	1	0	4	2	2	2	1	1	0.036	18/496

Table 13: Confusion Matrix: Convolutional Neural Network

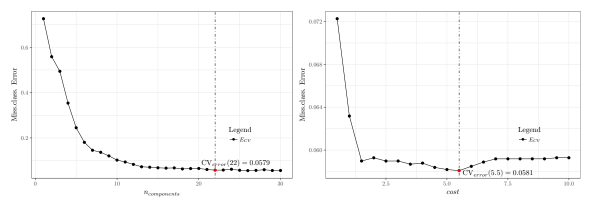
### 4.3 Support vector machines w/ PCA

Principal Components Analysis(PCA) showed quite powerful in combination with Support Vector Machines(SVM). First the principal components(PC) was calculated and a as low as possible while still describing the data well number of components were selected as seen in the first plot below. When plotting the first 100 PC 22 is a good number to satisfy these two objectives. On the right of this plot the training data is plotted defined by the two first principal components where the each digit is represented by a unique colour. In the third plot the error over each number of PC's used when using SVM. Here we see that 22 components give a small cv-error while not necessarily been the lowest. Since we want to keep the number of components as low as possible, picking more will lead to increased complexity without a justifiable increase in accuracy. A grid of cost parameters always returning the best were used to find optimal number of components using SVM. After having the PC's, the cost parameter is calculated the same way. This is shown in the last plot giving the best cost parameter to be 5.5. NB! Note that the last two plots have different scales in order to show the alteration in cv-error.



(a) Variance described by 100 first PC's

(b) Data by two first components from PCA



- (a) CV-error for SVM over number of PC's
- (b) CV-error for 22 PC's over cost parameter

Figure 8: Finding the best set up for PCA/SVM

With the set-up described above the PCA/SVM-model give a misclassification error of 5.6% meaning it predicts better than the other models with exception to the CNN still being the best model.

	0	1	2	3	4	5	6	7	8	9	Error	Rate
0	47	0	0	0	0	0	0	0	0	0	0.000	0/47
1	0	48	0	0	0	0	1	0	0	0	0.020	1/49
2	0	0	46	0	0	0	0	0	1	1	0.042	2/48
3	0	0	1	41	0	0	0	1	2	0	0.089	4/45
4	0	0	0	0	53	0	0	2	0	1	0.054	3/56
5	0	0	0	4	0	46	0	0	0	0	0.080	4/50
6	0	0	0	0	1	0	48	0	1	0	0.040	2/50
7	0	0	0	0	0	0	0	49	0	4	0.075	4/53
8	0	0	1	1	0	0	0	1	45	1	0.082	4/49
9	0	0	0	$^2$	1	0	0	1	0	45	0.082	4/49
Total	0	0	2	7	2	0	1	5	4	7	0.056	28/496

Table 14: Confusion Matrix: Support Vector Machine  $\mathbf{w}/$  PCA

### 4.4 K-Nearest Neighbours

The K-Nearest Neighbours algorithm is in this task used with the standard euclidean distance when calculating finding the nearest neighbours. This means the task is to find the optimal number of neighbours k to classify the data. In order to find a good measure for the out-of-sample using CV. This is sped up by parallelization of the CV. As seen below normalizing the data only slightly altered the cv-error and not in a clearly positive or negative way which makes sense since it here meant dividing all the pixels by maximum value for the greyscale colour. As seen in the plot below k=1 give the lowest cv-error meaning each new data point is classified by the single most similar neighbour. It is also worth noting the peak for k=2 which is the consequence of using a even number of neighbours for a problem with a even number of categories. This comes into greatest affect here when k=2, but also decreasingly visible for k=4,6,...

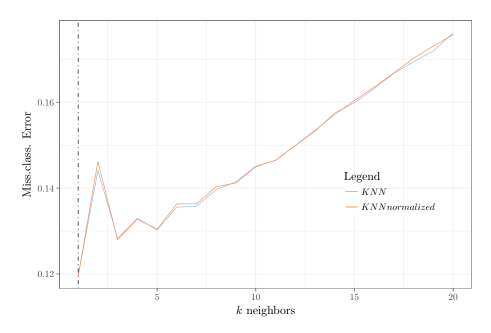


Figure 9

The test predicted using k=1 on the training set yield a misclassification error equal to 10.1% making it slightly better than bagging by 0.2%.

	0	1	2	3	4	5	6	7	8	9	Error	Rate
0	45	0	0	1	0	1	0	0	1	0	0.062	3/48
1	0	46	3	0	0	2	0	2	1	0	0.148	8/54
2	0	0	45	0	0	0	0	2	1	0	0.062	3/48
3	0	0	0	40	0	1	0	0	6	0	0.149	7/47
4	0	1	0	0	52	0	0	3	1	4	0.148	9/61
5	1	0	0	4	0	41	1	0	1	0	0.146	7/48
6	1	0	0	0	1	0	48	0	0	0	0.040	2/50
7	0	1	0	0	0	0	0	45	0	1	0.043	2/47
8	0	0	0	3	0	0	0	1	37	0	0.098	4/41
9	0	0	0	0	2	1	0	1	1	47	0.096	5/52
Total	2	2	3	8	3	5	1	9	12	5	0.101	50/496

Table 15: Confusion Matrix: K-Nearest Neighbours

### 4.5 Summary

Below is a table containing all the methods together with the digit classification error rate obtained on the test set sampled from the training set.

	Method	Digit Classification Error
1	Convolutional NN	0.036
2	Support Vector Machine	0.056
3	Random Forest	0.081
4	Artificial NN	0.091
5	K-Nearest Neighbours	0.101
6	Bagging	0.103
7	Boosting	0.119
8	Classification Tree	0.371

### 4.6 Conclusion

By the result obtained by testing each methods on the test set I conclude that Convolutional Neural Network produce the best method for classifying handwritten digits. Further I say that this means it also performs best on classifying if a digit is even or odd. The CNN as defined in 5 is used to train on the whole data set and make predictions for the unlabelled data. A .csv-file containing predicted digit and if it is odd or even in the columns Digit and isOdd(1 = odd).

### 5 Discussion

As viewed in the results 4 the different methods performed fairly well on classifying handwritten digits, with exception to the single classification tree where a higher error is expected. Four out of the eight models trained obtained an error below 10%. All these was obtained using around 2000 data points and with what we know from the theory the precision would most likely increase if given more data. We can clearly see that there are two models that outperform the other, these being convolutional neural network and support vector machines. This confirms the notion that these are very powerful methods within machine learning and why they are popular in use. Especially the convolutional neural network which was the clearly best model used in this report. Though some criteria are required to use this type of network, it is even more powerful where it shines, namely "image recognition". This being a field that is very popular today, makes machine learning a very interesting and exciting branch of applications to practise and study.

### 6 Appendices

### 6.1 R-Code: Help Functions

```
1 require (caret)
                             # useful library to split up data set
2 library(tikzDevice)
                             # library to export plots to .tex files
3 library(xtable)
                             # library to export data frames to tables in .tex
      files
4
  options(tikzMetricPackages = c("\\usepackage[utf8]{inputenc}\", "\\usepackage
5
       [T1] {fontenc}",
                                     "\\usetikzlibrary{calc}", "\\usepackage{
6
                                         amssymb}"))
8
  ggplot_to_latex <- function(</pre>
9
       ggplot,
10
       destination_path,
       width,
11
       height
12
  ) {
13
14
       tikz(file = paste0(destination_path, ".tex"), width = 6, height = 4)
15
       print(ggplot)
       dev.off()
16
17
  }
18
19
  create_confusion_matrix <- function(</pre>
20
       predicted_value,
21
       true_value,
       destination_path
22
  ) {
23
       conf <- confusionMatrix(predicted_value, true_value)</pre>
24
25
       conf_df <- as.data.frame.matrix(conf$table) # extract confusion matrix</pre>
26
       # add row for total error
27
       conf_df <- rbind(conf_df, Total = rep(0, ncol(conf_df)))</pre>
28
29
       rows_in_df <- nrow(conf_df)</pre>
30
       classification_frac <- rep("", rows_in_df)</pre>
31
       classification_float <- rep(0, rows_in_df)</pre>
32
33
       total_wrong <- 0
34
35
       total_classified <- 0
36
       # make columns that shows accuracy
37
38
       for(i in 1:(rows_in_df - 1)){
39
           correct_classified <- conf_df[i, i]</pre>
40
           amount_classified <- sum(conf_df[i, ])
41
           missclassified <- amount_classified - correct_classified
42
43
           classification_frac[i] <- paste0(missclassified, "/", amount_</pre>
               classified)
           classification_float[i] <- missclassified / amount_classified</pre>
44
45
46
           total_wrong <- total_wrong + missclassified
47
           total_classified <- total_classified + amount_classified
48
49
       classification_frac[rows_in_df] <- pasteO(total_wrong, "/", total_</pre>
50
           classified)
```

```
51
       classification_float[rows_in_df] <- total_wrong / total_classified</pre>
52
       conf_df <- cbind(temp = row.names(conf_df),</pre>
53
                                                                  # added extra
           column
                         conf_df,
                                                                  # to get predicted
54
                              classes
                         Error = classification_float,
55
                         Rate = classification_frac)
56
       names(conf_df) <- c("", names(conf_df)[-1]) # remove name of predicted
57
           classes
58
       write.csv(x = conf_df, file = paste0(destination_path, "_Confusion_
59
           Matrix.csv"))
       print(xtable(conf_df, display = c("s", rep("d", 11), "f", "s"),
60
                     digits = c(rep(0, 12), 3, 0)),
61
62
             #table.placement = "H",
             only.contents = TRUE,
63
             file = paste0(destination_path ,"_Confusion_Matrix.tex"),
64
             include.rownames = FALSE)
65
66
67
  }
68
69
  create_cv_indexes <- function(N, n_folds){</pre>
70
       indexes_per_fold <- floor(N/n_folds)</pre>
71
       index_matrix <- matrix(OL, nrow = n_folds, ncol = indexes_per_fold)</pre>
72
       index_available <- 1:N
73
       for(i in 1:n_folds){
           selected_indexes <- sample(index_available, indexes_per_fold)</pre>
74
           index_available <- index_available[! index_available %in% selected_</pre>
75
               indexesl
76
77
           index_matrix[i, ] <- selected_indexes</pre>
78
79
      return(index_matrix)
80 }
```

../R scripts/Help Scripts/to latex functions.R

### 6.2 R-Code: Regression Tree

```
1 ## Libraries and seed
2 rm(list = ls())
3 library(caret)
4 library (readr)
5 library(tree)
                           # package used for regression tree
6 library(tikzDevice)
                           # library to export plots to .tex files
8
  options(tikzMetricPackages = c("\\usepackage[utf8]{inputenc}", "\\usepackage
      [T1]{fontenc}",
                                  "\\usetikzlibrary{calc}", "\\usepackage{
9
                                      amssymb}"))
10
  set.seed(420)
11
12
  if(!exists("create_confusion_matrix", mode = "function")){
13
      source("Help_Scripts/to_latex_functions.R")
14
15
16
17
        ----#
18
```

```
19 ## Data
20 path_data <- paste0(getwd(), "/data")</pre>
21 path_to_here <- paste0(getwd(), "/Tree_Based_Methods") # getwd give path
      to project
22
  # which is one folder over
2.3
  train_data <- read.csv(paste0(path_data, "/Train_Digits_20171108.csv"),</pre>
      header = TRUE)
  unclassified_data <- read.csv(paste0(path_data, "/Test_Digits_20171108.csv")
25
      , header = TRUE)
26
27 # Remove unnessesary varibles which have a low variance
28 split_train_test <- createDataPartition(train_data$Digit, p = 0.8, list =
      FALSE)
29 test_data <- train_data[-split_train_test, ]</pre>
30 train_data <- train_data[split_train_test, ]
32 # Remove variable with low variance which are near zero. Doing it after
33 # splitting in train/test set to avoid contaminating the data.
34 near_zero_variables <- nearZeroVar(train_data[,-1], saveMetrics = T, freqCut
       = 10000/1, uniqueCut = 1/7)
35 cut_variables <- rownames(near_zero_variables[near_zero_variables$nzv ==
      TRUE,])
36 variables <- setdiff(names(train_data), cut_variables)
38 train_data <- train_data[, variables]</pre>
39 test_data <- test_data[, variables]</pre>
40
41 # make the label Digit factor
42 train_data[, 1] <- as.factor(train_data[, 1])
43 test_data[, 1] <- as.factor(test_data[, 1])
44
45 unclassified_data[,1] <- as.factor(unclassified_data[,1])
46
47
48
49 ## REGRESSION - tree
50
51 regression <- function(
52
      minimum_development,
53
      train_data,
54
      test_data
      ) {
55
56
      # Chanage name of pixel columns to work with tikz library
57
       colnames(train_data)[ 2:length(train_data[1,])] <- c(pasteO("Pixel", 1:(</pre>
58
           length(train_data[1,]) - 1)))
       colnames(test_data)[ 2:length(train_data[1, ])] <- c(paste0("Pixel", 1:(</pre>
59
          length(train_data[1,]) - 1)))
60
      # Set minimum development required for making a new split
61
      minimum_development <- 0.005
62
      tree_model <- tree(Digit ~ ., data = train_data, mindev = minimum_</pre>
63
          development)
64
      plot(tree_model)
65
      text(tree_model, cex = .5)
      print(summary(tree_model))
66
67
68
       cross_validation <- cv.tree(tree_model, K = 10)</pre>
```

```
69
       cross_validation$k[1] <- 0</pre>
70
       alpha <- round <- round(cross_validation$k)</pre>
71
72
       plot(cross_validation$size, cross_validation$dev, type = "b",
             xlab = "Number of terminal nodes", ylab = "CV error")
73
74
75
       ggplot_df <- data.frame(size = cross_validation$size, dev = cross_
           validation $dev)
76
       destination_path <- paste0(path_to_here, "/Results_TBM/Regression_Tree")</pre>
77
78
79
       ggplot1 <- ggplot(data = ggplot_df, aes(x = size, y = dev)) +</pre>
                   geom_line(aes(colour = "$RegressionTree$"), linetype = "
80
                       dashed") +
                   geom_point() +
81
                   geom_vline(xintercept = 20, color = "black", linetype = "
82
                       dotdash") +
                   xlab("n\\_\{terminal nodes\}") +
83
                   ylab("CV Error") +
84
                   scale_colour_manual("Legend",
85
                                         breaks = c("$RegressionTree$"),
86
87
                                         values = c("#91bfdb"),
88
                                         guide = guide_legend(override.aes = list(
89
                                              linetype = c("solid"),
90
                                              shape = c(16)
91
                                         ))) +
92
                   theme_bw() +
                   theme(legend.position = c(0.8, 0.355),
93
                       legend.background = element_rect(fill=alpha('white', 0)))
94
       ggsave(paste0(destination_path, ".png"))
95
96
       ggplot_to_latex(ggplot1, destination_path, width = 5, height = 5)
97
98
       tree_prune <- prune.tree(tree_model, best = 20)</pre>
99
       summary(tree_prune)
100
       tikz(file = paste0(destination_path, "_Tree.tex"), width = 6, height =
101
           4)
102
       plot(tree_prune)
103
       text(tree_prune, cex = .5)
104
       dev.off()
105
       predicted <- predict(tree_prune, test_data, type = "class")</pre>
106
107
       create_confusion_matrix(predicted, test_data[,1], destination_path)
108
109
110 regression(0.05, train_data, test_data)
                     ../R scripts/Tree Based Methods/Regression Tree.R
```

### 6.3 R-Code: Random Forest

```
1 ## Libraries and seed
2 rm(list = ls())
3 library (randomForest)
                           # library giving a easy-to-use random forest method
                           # useful library to split up data set
4 library(caret)
5 library(tikzDevice)
                           # library to export plots to .tex files
6 library(xtable)
                           # library to export data frames to tables in .tex
      files
  set.seed(420)
                           # seed to replicate results and get consistent test
      and training set
8
  # Load help script with functions to export the results to latex
9
10 # These functions gathered to avoid duplicate code
11 if(!exists("create_confusion_matrix", mode = "function")){
12
      source("Help_Scripts/to_latex_functions.R")
13 }
14
15 #----#
16
17 ## Data
18 path_data <- paste0(getwd(), "/data")</pre>
19 path_to_here <- paste0(getwd(), "/Tree_Based_Methods")
                                                            # getwd give path
      to project
                                                             # which is one
20
                                                                 folder over
21
  train_data <- read.csv(paste0(path_data, "/Train_Digits_20171108.csv"),</pre>
22
      header = TRUE)
  unclassified_data <- read.csv(paste0(path_data, "/Test_Digits_20171108.csv")
      , header = TRUE)
24
  train_data[,1] <- as.factor(train_data[, 1])</pre>
25
26
  # split training set into training and test set
27
28
29 split_train_test <- createDataPartition(train_data$Digit, p = 0.8, list =
      FALSE)
30 test_data <- train_data[-split_train_test, ]
31 train_data <- train_data[split_train_test, ]</pre>
32
33
  #----#
34
35 ## Random forest
36
  # Train forest
  train_random_forest <- function(</pre>
37
38
      data,
39
      n_trees,
40
      minimum_development = 0.01
41
           random_forest <- randomForest(Digit ~ .,</pre>
42
43
                                          data = data,
44
                                          ntree = n_trees,
45
                                          #mindev = minimum_development,
                                          importance = TRUE,
46
                                          na.action = na.exclude)
47
          return(random_forest)
48
      }
49
50
```

```
51 # Plot error as the number of trees increase
52
  plot_error_development <- function(</pre>
53
       random_forest_data,
54
55
       destination_path
56
           error_data <- data.frame(n_trees = 1:nrow(random_forest_data$err.</pre>
57
               rate).
                                       error <- random_forest_data$err.rate[,"00B"</pre>
58
                                          ])
59
           write.csv(error_data, file = paste0(destination_path, ".csv"))
60
61
           ggplot1 <- ggplot(data = error_data, aes(x = n_trees)) +</pre>
62
                geom_line(aes(y = error, colour = "$Random forest")) +
63
64
               xlab("$n\\_{trees}$") +
               ylab("Miss. class. Error") +
65
               scale_colour_manual("Legend",
66
                                     breaks = c("$Random forest$"),
67
                                     values = c("black"),
68
                                     guide = guide_legend(override.aes = list(
69
                                          linetype = c("solid"),
70
71
                                          shape = c(16)
72
                                     ))) +
73
                theme(legend.position = c(0.9, 0.2))
74
           ggsave(paste0(destination_path, ".png"))
75
           ggplot_to_latex(ggplot1, destination_path, width = 6, height = 4)
76
77
78
  main <- function(){</pre>
79
      n_{trees} = 50
80
      random_forest <- train_random_forest(train_data, n_trees)</pre>
81
      plot_error_development(random_forest, paste0(path_to_here, "/Results_TBM
82
           /Random_Forest_",
                                                        n_trees, "trees_Error_plot"
83
                                                            ))
84
85
       prediction <- predict(random_forest, newdata = test_data)</pre>
86
       create_confusion_matrix(predicted_value = prediction, true_value = test_
           data$Digit,
                                 paste0(path_to_here, "/Results_TBM/Random_Forest
87
                                                                         n_trees, "
88
                                                                            trees"))
89
90
  main()
                    ../R scripts/Tree Based Methods/Random Forest.R
```

### 6.4 R-Code: Bagging

```
1 ## Libraries and seed
2 rm(list = ls())
3 library (randomForest)
                          # library giving a easy-to-use random forest method
4 library(caret)
                           # useful library to split up data set
5 library(tikzDevice)
                           # library to export plots to .tex files
6 library(xtable)
                           # library to export data frames to tables in .tex
      files
  set.seed(420)
                           # seed to replicate results and get consistent test
      and training set
8
  # Load help script with functions to export the results to latex
9
10 # These functions gathered to avoid duplicate code
11 if(!exists("create_confusion_matrix", mode = "function")){
12
      source("Help_Scripts/to_latex_functions.R")
13 }
14
15 #----#
16
17 ## Data
18
19 path_data <- paste0(getwd(), "/data")</pre>
20 path_to_here <- paste0(getwd(), "/Tree_Based_Methods")  # getwd give path
     to project
21 # which is one folder over
22
23 train_data <- read.csv(paste0(path_data, "/Train_Digits_20171108.csv"),
     header = TRUE)
24 unclassified_data <- read.csv(paste0(path_data, "/Test_Digits_20171108.csv")
      , header = TRUE)
25
  train_data[,1] <- as.factor(train_data[, 1])</pre>
26
27
28 # split training set into training and test set
29
30 split_train_test <- createDataPartition(train_data$Digit, p = 0.8, list =
      FALSE)
31 test_data <- train_data[-split_train_test, ]</pre>
32 train_data <- train_data[split_train_test, ]</pre>
33
  #----#
34
35
36 ## Random forest
37
  # Train forest
38 train_bagging <- function(
39
      data,
40
      n_trees
41
  ) {
42
      n_features <- ncol(data) - 1</pre>
      bagging <- randomForest(Digit ~ .,</pre>
43
44
                                      data = data,
45
                                      ntree = n_trees,
                                      mtry = n_features,
46
47
                                      importance = TRUE,
48
                                      na.action = na.exclude)
      return(bagging)
49
50 }
51
```

```
52 # Plot error as the number of trees increase
53
  plot_error_development <- function(</pre>
54
55
       random_forest_data,
56
       destination_path
57
  ) {
       error_data <- data.frame(n_trees = 1:nrow(random_forest_data$err.rate),</pre>
58
                                  error <- random_forest_data$err.rate[,"00B"])</pre>
59
60
       write.csv(error_data, file = paste0(destination_path ,".csv"))
61
       ggplot1 <- ggplot(data = error_data, aes(x = n_trees)) +</pre>
62
           geom_line(aes(y = error, colour = "$Bagging$")) +
63
           xlab("n\\\\) +
64
           ylab("Miss.class. Error") +
65
           scale_colour_manual("Legend",
66
                                 breaks = c("$Bagging$"),
67
                                 values = c("black"),
68
                                 guide = guide_legend(override.aes = list(
69
                                     linetype = c("solid"),
70
71
                                     shape = c(16)
72
                                 ))) +
73
           theme(legend.position = c(0.9, 0.2))
74
       ggsave(paste0(destination_path, ".png"))
75
76
       ggplot_to_latex(ggplot1, destination_path, width = 6, height = 4)
77
78
  main <- function(){</pre>
79
      n_trees <- 50
80
      bagging <- train_bagging(train_data, n_trees)</pre>
81
       plot_error_development(bagging, pasteO(path_to_here, "/Results_TBM/
82
           Bagging_",
83
                                                 n_trees, "trees_Error_plot"))
84
      prediction <- predict(bagging, newdata = test_data)</pre>
85
86
87
       create_confusion_matrix(predicted_value = prediction, true_value = test_
           data$Digit,
                                 pasteO(path_to_here, "/Results_TBM/Bagging_",
88
89
                                        n_trees, "trees"))
90
  }
91
92 main()
```

### 6.5 R-Code: Boosting

```
1 ## Libraries and seed
2 rm(list = ls())
3 library(caret)
                           # useful library to split up data set
4 library (gbm)
                           # library with powerful boosting method
                           # seed to replicate results and get consistent test
5 set.seed (420)
      and training set
6
7
  # Load help script with functions to export the results to latex
8
  # These functions gathered to avoid duplicate code
  if(!exists("create_confusion_matrix", mode = "function")){
9
      source("Help_Scripts/to_latex_functions.R")
10
11 }
12
  #----#
1.3
14
15 ## Data
17 path_data <- paste0(getwd(), "/data")</pre>
18 path_to_here <- paste0(getwd(), "/Tree_Based_Methods") # getwd give path
     to project
19
20 train_data <- read.csv(paste0(path_data, "/Train_Digits_20171108.csv"),
     header = TRUE)
21 unclassified_data <- read.csv(paste0(path_data, "/Test_Digits_20171108.csv")
      , header = TRUE)
22
23 train_data[,1] <- as.factor(train_data[, 1])
24
25 # split training set into training and test set
26
27 split_train_test <- createDataPartition(train_data$Digit, p = 0.8, list =
      FALSE)
28 test_data <- train_data[-split_train_test, ]</pre>
29 train_data <- train_data[split_train_test, ]</pre>
30
31 # Remove variable with low variance which are near zero. Doing it after
32 # splitting in train/test set to avoid contaminating the data.
33 near_zero_variables <- nearZeroVar(train_data[,-1], saveMetrics = T, freqCut
      = 10000/1, uniqueCut = 1/7)
34 cut_variables <- rownames(near_zero_variables[near_zero_variables$nzv ==
      TRUE,])
35 variables <- setdiff(names(train_data), cut_variables)
36
37 train_data <- train_data[, variables]
38 test_data <- test_data[, variables]</pre>
39
40
  #----#
41
  ## Boosting
42
  # Train booster
43
44 boosting <- function(
45
      data,
46
      n_trees,
47
      interaction_depth = 2,
48
      shrinkage = 0.001
49 ) {
      boosting <- gbm(Digit ~ .,</pre>
50
```

```
51
                         data = data,
                         distribution = "multinomial",
52
                         n.trees = n_trees,
53
54
                         interaction.depth = interaction_depth,
55
                         shrinkage = shrinkage,
                         cv.folds = 10,
56
                         n.cores = 4)
57
       return(boosting)
58
   }
59
60
61
62
   # Plot error as the number of trees increase
63
   plot_error_development <- function(</pre>
64
65
       boosting_data,
66
       best_n_trees,
67
       destination_path
   ) {
68
       error_data <- data.frame(n_trees = 1:length(boosting_data$cv.error),</pre>
69
                                   error <- boosting_data$cv.error)</pre>
70
       write.csv(error_data, file = paste0(destination_path ,".csv"))
71
72
73
       ggplot1 <- ggplot(data = error_data, aes(x = n_trees)) +</pre>
74
            geom_line(aes(y = error, colour = "$Boosting$")) +
75
            xlab("$n_{trees}) +
76
            ylab("Miss.class. Error") +
77
            geom_vline(xintercept = best_n_trees, color = "black", linetype = "
                dotdash") +
            scale_colour_manual("Legend",
78
                                  breaks = c("$Boosting$"),
79
                                  values = c("black"),
80
                                  guide = guide_legend(override.aes = list(
81
                                      linetype = c("solid"),
82
83
                                      shape = c(16)
                                  ))) +
84
            theme(legend.position = c(0.9, 0.2)) +
85
86
            theme_bw() +
87
            theme(legend.position = c(0.8, 0.355),
                  legend.background = element_rect(fill=alpha('white', 0)))
88
29
       ggsave(paste0(destination_path, ".png"))
90
       ggplot_to_latex(ggplot1, destination_path, width = 6, height = 4)
91
92
   }
93
   predict_data <- function(</pre>
94
95
       boosting_train,
96
       best_n_trees,
97
       test_data
98
   ) {
       predicted <- predict(boosting_train, newdata = test_data, n.trees = best</pre>
99
           _n_trees, type = "response")
100
       predicted <- apply(predicted, 1, function(x) which.max(x) - 1)</pre>
101
       return(predicted)
102
103 }
104
105 main <- function(){
       n_{trees} = 10000
107
       boosting_train <- boosting(train_data,n_trees, shrinkage = 0.01)</pre>
```

```
108
109
       \# Get amount of trees with best performance based on 10 fold cv
110
       best_n_trees <- gbm.perf(boosting_train, method = "cv", plot.it = FALSE)</pre>
111
       # Plot error
112
       plot_error_development(boosting_train, best_n_trees, paste0(path_to_here
113
                                                         "/Results_TBM/Boosting_",
114
115
                                                         n_trees,
116
                                                         "trees_Error_plot"))
117
       # Perdict test set labels
118
119
       predicted <- predict_data(boosting_train, best_n_trees, test_data)</pre>
120
121
       # Create confusion matrix for the result
       create_confusion_matrix(predicted, test_data$Digit, paste0(path_to_here,
122
                                                                      "/Results_TBM/
123
                                                                         Boosting_"
124
                                                                      n_trees))
125 }
126
127 main()
```

### 6.6 R-Code: Plot Random Forest w/ Bagging

```
1 rm(list = ls())
2 library(ggplot2)
3 library(tikzDevice)
                            # library to export plots to .tex files
  path_data <- paste0(getwd(), "/data")</pre>
5
  path_to_here <- pasteO(getwd(), "/Tree_Based_Methods")</pre>
6
                                                               # getwd give path
      to project
                                                               # which is one
                                                                   folder over
8
  plot_random_forest_bagging <- function(</pre>
9
      n_trees,
10
      path,
11
12
      destination_path
13
      ) {
14
           rf_path <- paste0(path,
                               "Random_Forest_",
15
16
                               n_trees,
17
                               "trees_Error_plot_",
18
                               n_trees,
                               "trees.csv")
19
           bagging_path <- paste0(path,
20
21
                               "Bagging_",
22
                               n trees.
23
                               "trees_Error_plot_",
24
                               n trees.
25
                               "trees.csv")
26
27
           random_forest_error <- read.csv(rf_path)
28
           bagging_error <- read.csv(bagging_path)</pre>
29
           ggplot_df <- data.frame(n_trees = 1:nrow(bagging_error),</pre>
30
31
                                     rf = random_forest_error[3],
                                     bag = bagging_error[3])
32
           names(ggplot_df) <- c("n_trees", "rf", "bag")</pre>
33
           print(str(ggplot_df))
34
35
           tikz(file = paste0(destination_path, ".tex"), width = 6, height = 4)
36
           ggplot1 <- ggplot(data = ggplot_df, aes(x = n_trees)) +</pre>
37
38
               geom_line(aes(y = rf, colour = "$Random forest$")) +
39
               geom_line(aes(y = bag, colour = "$Bagging$")) +
40
               geom_vline(xintercept = 150, color = "black", linetype = "
                   dotdash") +
               xlab("$n_{trees}) +
41
               ylab("Miss.class. Error") +
42
               scale_colour_manual("Legend",
43
                                     breaks = c("$Random forest$", "$Bagging$"),
44
                                     values = c("#91bfdb", "#fc8d59"),
45
                                     guide = guide_legend(override.aes = list(
46
47
                                          linetype = c("solid", "solid"),
                                          shape = c(16, 16)
48
                                     ))) +
49
               scale_y_continuous(limits = c(0, 0.4)) +
50
               theme_bw() +
51
52
               theme(legend.position = c(0.8, 0.455),
                      legend.background = element_rect(fill=alpha('white', 0)))
53
           ggsave(paste0(destination_path, ".png"))
54
```

```
55
           print(ggplot1)
56
           dev.off()
57
58
  main <- function(
59
60
      ) {
61
           n_{trees} = 500
62
           path = paste0(path_to_here, "/Results_TBM/")
63
           destination_path = paste0(path, "/Random_Forest_Bagging_",
64
                                       n_trees, "trees")
65
66
           plot_random_forest_bagging(n_trees, path, destination_path)
67
68
      }
69
70
71 main()
```

 $../R\_scripts/Tree\_Based\_Methods/Plot\_Random\_Forest\_Bagging.R$ 

### 6.7 R-Code: Neural Network

```
1 ## Libraries and seed
2 rm(list = ls())
3 library(h2o)
4 library(caret)
5 library (reshape2)
  set.seed(420)
9
  # Load help script with functions to export the results to latex
10
  # These functions gathered to avoid duplicate code
  if(!exists("create_confusion_matrix", mode = "function")){
11
       source("Help_Scripts/to_latex_functions.R")
12
13 }
14
  #----#
15
  ## Data
16
17
18 path_data <- getwd()
19 path_to_here <- pasteO(getwd(), "/Neural_Networks")</pre>
21 train_data <- read.csv(paste0(path_data, "/data/Train_Digits_20171108.csv"))
22 unclassified_data <- read.csv(pasteO(path_data, "/data/Test_Digits_20171108.
      csv"))
23
24 local.h2o <- h2o.init(ip = "localhost", port = 54321, startH2O = TRUE,max_
      mem_size = "7G", nthreads = -1)
25
26
  train_data[,1] <- as.factor(train_data[, 1])</pre>
  split_train_test <- createDataPartition(train_data$Digit, p = 0.8, list =</pre>
      FALSE)
28 test_data <- train_data[-split_train_test, ]
  train_data <- train_data[split_train_test, ]</pre>
29
30
31 train_data <- as.h2o(train_data)</pre>
32 unclassified_data <- as.h2o(unclassified_data)
33 test_data <- as.h2o(test_data)
34
35
36
  ## Getting useful data from grid run of neural networkss
38
39
  get_data_in_df <- function(</pre>
40
      data
41 )
42 {
43
      n <- length(data@model_ids)</pre>
      mse_errors <- rep(0,n)</pre>
44
45
      mean_per_class_errors <- rep(0,n)</pre>
      hidden <- rep("", n)
46
47
      str(hidden)
48
      rate <- rep(0,n)
49
      11 < - rep(0,n)
       epochs <- rep(0,n)
50
      model_numbers <- rep(0,n)
51
      train_error <- rep(0,n)</pre>
52
      train_mse <- rep(0,n)</pre>
53
      test_error <- rep(0,n)</pre>
54
```

```
55
        test_mse <- rep(0,n)</pre>
56
        activation <- rep("",n)
        input_dropout_ratio <- rep(0,n)
57
       nesterov_accelerated_gradient <- rep("", n)</pre>
58
59
60
       model_df <- data.frame(model_numbers = mse_errors, hidden, rate, 11,</pre>
            epochs,
                                 train_error, test_error, train_mse, test_mse,
61
                                      activation, input_dropout_ratio,
                                      stringsAsFactors = FALSE)
        str(model df)
62
63
       for(i in 1:n){
64
            model <- h2o.getModel(data@model_ids[[i]])</pre>
65
            model_df$mse_errors[i] <- h2o.mse(model)</pre>
66
            #model_df$mean_per_class_error[i] <- model@model$cross_validation_
67
                metrics@metrics$mean_per_class_error
            model_df$mean_per_class_error[i] <- h2o.performance(model, xval = T)</pre>
68
                @metrics$mean_per_class_error
69
70
            model_paramaters <- model@allparameters</pre>
            model_name <- model@model_id
71
            model_number <- sub(".*model_(.*)$", "\\1", model_name)</pre>
72
73
            model_df$model_numbers[i] <- as.integer(model_number)</pre>
74
            model_df$hidden[i] <- paste(as.character(model_paramaters$hidden),</pre>
                sep = " ", collapse = ", ")
75
            model_df$rate[i] <- model_paramaters$rate</pre>
            model_df$11[i] <- model_paramaters$11</pre>
76
            model_df$epochs[i] <- model_paramaters$epochs
77
            model_df$activation[i] <- model_paramaters$activation</pre>
78
            model_df$input_dropout_ratio[i] <- model_paramaters$input_dropout_
79
                ratio
            model_df$nesterov_accelerated_gradient[i] <- model_paramaters$
80
                nesterov_accelerated_gradient
81
            train_performance <- h2o.performance(model, train_data)@metrics</pre>
82
            train_performance_error <- train_performance$mean_per_class_error</pre>
83
84
            train_performance_mse <- train_performance$MSE</pre>
85
86
            model_df$train_error[i] <- train_performance_error</pre>
87
            model_df$train_mse[i] <- train_performance_mse</pre>
88
            test_performance <- h2o.performance(model, test_data)@metrics</pre>
89
90
            test_predictions <- h2o.predict(model, test_data)</pre>
91
            test_accuracy <- test_predictions$predict == test_data$Digit</pre>
            test_performance_error <- 1 - mean(test_accuracy)</pre>
92
93
            test_performance_mse <- test_performance$MSE</pre>
94
            model_df$test_error[i] <- test_performance_error</pre>
95
            model_df$test_mse[i] <- test_performance_mse</pre>
96
97
98
       model_df <- model_df[with(model_df, order(model_numbers)),]</pre>
99
        model_df
100
101 }
102
103 # Run and plot grid of n neural nets
104 run_neural_network_grid <- function(){</pre>
       activation <- list("Rectifier", "RectifierWithDropOut")# "Tanh")</pre>
```

```
hidden <- list(c(100,100), c(150, 150), c(540, 320), c(100, 100, 100),
106
           c(540, 320, 100))
107
       input_dropout_ratio <- list(0, 0.2)
       nesterov_accelerated_gradient <- list( TRUE)</pre>
108
109
       epochs <- list(20)
       11 = list(0, 1.4e-5)
110
       hyper_params <- list(activation = activation, hidden = hidden, input_
111
           dropout_ratio = input_dropout_ratio, nesterov_accelerated_gradient =
            nesterov_accelerated_gradient, epochs = epochs, l1 = l1)
112
       grid_deep_learning <- h2o.grid(algorithm = "deeplearning",</pre>
113
114
                                x = 2:785,
115
                                y = 1,
116
                                training_frame = train_data,
117
                                nfolds = 10,
                                stopping_metric = "MSE",
118
119
                                stopping_tolerance = 0.0025,
120
                                hyper_params = hyper_params)
       save_results <- function(results){</pre>
121
122
       write.csv(results, file = paste0(path_to_here, "/Neural_Networks/results
           _NN/grid_run_20.csv"))
123
124
125
       df <- get_data_in_df(grid_deep_learning)</pre>
126
       save_results(df)
127
128
       #results_df <- df
129
       results_df <- read.csv(pasteO(path_to_here, "/Neural_Networks/results_NN
130
           /grid_run_40.csv"))
131
       results_df <- results_df[with(results_df, order(mean_per_class_error)),]
132
133
       results_df$row_names <- 1:length(results_df[,1])</pre>
134
       melt_datas <- melt(results_df[c("test_error","mean_per_class_error", "</pre>
135
           row_names",
                                 "model_numbers")], id = c("row_names", "model_
136
                                     numbers"))
137
       plot_list <- list()</pre>
138
       # Plot classification error
139
       plot_list[[1]] <- ggplot(data=melt_datas,</pre>
                          aes(x=row_names, y=value)) +
140
       geom_point(aes(colour = as.factor(model_numbers), group = as.factor(
141
           model_numbers)), size = 1.25) +
       geom_line(aes(group = variable)) +
142
       xlab("Model id") +
143
       ylab("Miss. class. Error") +
144
145
       scale_y_continuous(limits = c(0, 0.2)) +
146
       theme_bw() +
       theme(legend.position = c(0.675, 0.255),
147
         legend.background = element_rect(fill=alpha('white', 0)),
148
         legend.direction = "horizontal",
149
150
         legend.text = element_text(size=6),
         legend.key = element_rect(size = 3),
151
152
         legend.key.size = unit(1.0, 'lines'),
153
         axis.text.x=element_blank(),
154
         axis.ticks.x=element_blank()) +
       scale_colour_discrete(name = "Model ids") +
155
156
       guides(fill = guide_legend(title = "Legend"))
```

```
157
        ggplot_to_latex(plot_list[[1]],
158
159
                paste0(path_to_here, "/results_NN/per_class_error"), width = 6,
                    height = 4)
        ggsave(paste0(path_to_here, "/results_NN/per_class_error3.png"))
160
161
162
   # Run final model based on results from grid run and make confusion matrix
163
       for prediction on test set
   run_final_model <- function(){</pre>
164
       deep_learning_results3<- h2o.deeplearning(x = 2:785,</pre>
165
                                                     y = 1,
166
                                                     training_frame = train_data,
167
                                                     activation = "Rectifier",
168
                                                     input_dropout_ratio = 0.2,
169
170
                                                     nfolds = 10,
                                                    balance_classes = TRUE,
171
                                                    hidden = c(540, 320),
172
                                                    11 = 1.4e-5,
173
                                                     stopping_metric = "MSE",
174
                                                     stopping_tolerance = 0.0025,
175
176
                                                     nesterov_accelerated_gradient =
                                                         TRUE,
                                                     epochs = 20)
177
178
179
       h2o.performance(deep_learning_results3, test_data)
180
181
       predicted <- predict(deep_learning_results3, test_data, type = "response</pre>
182
           ")
183
       predicted_confusion_matrix <- as.factor(as.vector(predicted$predict))</pre>
184
       test_data_confusion_matrix <- as.data.frame(test_data)</pre>
185
        create_confusion_matrix(predicted_confusion_matrix,
186
                                  test_data_confusion_matrix[, "Digit"],
187
                                  paste0(path_to_here, "/results_NN/540_320_neural
188
189 }
190
191 run_final_model
192
193 h2o.shutdown()
```

../R scripts/Neural Networks/neural network.R

### 6.8 R-Code: Convolutional Neural Network

```
1 ## Libraries and seed
2 rm(list = ls())
3 library(mxnet)
                       #library for running convolutional neural network
4 library(caret)
  set.seed(420)
6
  # Load help script with functions to export the results to latex
8
9
  # These functions gathered to avoid duplicate code
  if(!exists("create_confusion_matrix", mode = "function")){
10
      source("Help_Scripts/to_latex_functions.R")
11
12 }
  #----#
1.3
14
15 ## Data
16
17 path_data <- getwd()
path_to_here <- pasteO(getwd(), "/Neural_Networks")</pre>
20 train_data <- read.csv(paste0(path_data, "/data/Train_Digits_20171108.csv"))
21 unclassified_data <- read.csv(paste0(path_data, "/data/Test_Digits_20171108.
      csv"))
22
23 train_data[, 1] <- as.factor(train_data[, 1])
24 train_data_full <- train_data
25
26
  # Split the data into training, test and validation set
  split_train_test <- createDataPartition(train_data$Digit, p = 0.8, list =</pre>
      FALSE)
28
29
  test_data <- train_data[-split_train_test, ]</pre>
30 train_data <- train_data[split_train_test, ]
31
32 | split_train_validation <- createDataPartition(train_data$Digit, p = 0.85,
      list = FALSE)
33
34 validation_data <- train_data[-split_train_validation, ]</pre>
35 train_data_val <- train_data[split_train_validation, ]
36
37 # Setting up datasets as matrices in order to get correct input for mxnet
38
39 # Training after tuning
40 train <- train_data
41 train_x <- t(train[, -1])
42 train_y <- train[, 1]
43 train_array <- train_x
44 dim(train_array) <- c(28, 28, 1, ncol(train_x))
45
  # Training under tuning
46
47
  train_val <- train_data_val</pre>
48 train_x_val <- t(train_val[, -1])
49 train_y_val <- train_val[, 1]
50 train_array_val <- train_x_val
51 dim(train_array_val) <- c(28, 28, 1, ncol(train_x_val))
52
53 # Test set to compare with other methods
54 test_x <- t(test_data[, -1])
```

```
55 test_y <- test_data[, 1]
56 test_array <- test_x
   dim(test_array) <- c(28, 28, 1, ncol(test_x))</pre>
57
59 # Validation set used to validate under tuning
60 validation_x <- t(validation_data[, -1])
61 validation_y <- validation_data[, 1]
62 validation_array <- validation_x
63 dim(validation_array) <- c(28, 28, 1, ncol(validation_x))
64
65 # Get right format on the unclassified set
66 unclassified_x <- t(unclassified_data[, -1])
67 unclassified_array <- unclassified_x
68 dim(unclassified_array) <- c(28, 28, 1, ncol(unclassified_x))
70 # Full training data when making prediction for unclassified data
71 train_full <- train_data_full
72 train_x_full <- t(train_full[, -1])
73 train_y_full <- train_full[, 1]
74 train_array_full <- train_x_full
75 dim(train_array_full) <- c(28, 28, 1, ncol(train_x_full))
76
77
78
79
   data <- mx.symbol.Variable("data")</pre>
80
81
   two_layer_concolutional_network <- function(){</pre>
82
       # Setting up first convolutional layer
83
       convolution_1 <- mx.symbol.Convolution(data = data, kernel = c(5,5), num</pre>
84
           _filter = 30)
       activation_1 <- mx.symbol.Activation(data = convolution_1, act_type = "
85
           tanh")
       pooling_1 <- mx.symbol.Pooling(data = activation_1, pool_type = "max",</pre>
86
           kernel = c(2, 2), stride = c(2,2))
87
88
       # Setting up second convolutional layer
89
       convolution_2 <- mx.symbol.Convolution(data = pooling_1, kernel = c(5,5)</pre>
90
           , num_filter = 50)
       activation_2 <- mx.symbol.Activation(data = convolution_2, act_type = "
91
           tanh")
       pooling_2 <- mx.symbol.Pooling(data = activation_2, pool_type = "max",</pre>
92
           kernel = c(2, 2), stride = c(2,2)
93
       # Setting up first fully connected layer
94
95
96
       flatten <- mx.symbol.Flatten(data = pooling_2)</pre>
       fully_1 <- mx.symbol.FullyConnected(data = flatten, num_hidden = 500)
97
       activation_3 <- mx.symbol.Activation(data = fully_1, act_type = "tanh")
98
99
100
       # Setting up second fully connected layer
101
       fully_2 <- mx.symbol.FullyConnected(data = activation_3, num_hidden =
102
           10)
103
       return(fully_2)
104
105
106 }
```

```
107
108
   # three layer convolutional neural network to experiment, turned out to
       perform worse
   three_layer_concolutional_network <- function(){</pre>
109
       # Setting up first convolutional layers
110
111
       convolution_1 <- mx.symbol.Convolution(data = data, kernel = c(5,5), num</pre>
112
           _filter = 30)
       activation_1 <- mx.symbol.Activation(data = convolution_1, act_type = "
113
           tanh")
       pooling_1 <- mx.symbol.Pooling(data = activation_1, pool_type = "max",</pre>
114
           kernel = c(2, 2), stride = c(2,2)
115
       # Setting up second convolutional layers
116
117
       convolution_2 <- mx.symbol.Convolution(data = pooling_1, kernel = c(5,5)</pre>
118
           , num_filter = 50)
       activation_2 <- mx.symbol.Activation(data = convolution_2, act_type = "
119
           tanh")
       pooling_2 <- mx.symbol.Pooling(data = activation_2, pool_type = "max",</pre>
120
           kernel = c(2, 2), stride = c(2,2)
121
       # Setting up thrid convolutional layers
122
123
       convolution_3 <- mx.symbol.Convolution(data = pooling_2, kernel = c(3,3)</pre>
           , num_filter = 50)
124
       activation_3 <- mx.symbol.Activation(data = convolution_3, act_type = "
           tanh")
       pooling_3 <- mx.symbol.Pooling(data = activation_3, pool_type = "max",</pre>
125
           kernel = c(2, 2), stride = c(2,2))
126
       # Setting up first fully connected layer
127
128
129
       flatten <- mx.symbol.Flatten(data = pooling_3)</pre>
       fully_1 <- mx.symbol.FullyConnected(data = flatten, num_hidden = 500)
130
       activation_3 <- mx.symbol.Activation(data = fully_1, act_type = "tanh")
131
132
133
       # Setting up second fully connected layer
134
135
       fully_3 <- mx.symbol.FullyConnected(data = activation_3, num_hidden =</pre>
           40)
136
       return(fully_3)
137
138
139 }
140
   # Output layer, softmax gives probabilies for the output
141
142
   run_convolutional_neural_network <- function(</pre>
143
144
       neural_net_model,
145
       train_array,
146
       train_y,
       test_array,
147
       validation_array = NULL,
148
       validation_y = NULL
149
150 ) {
       mx.set.seed(100)
151
152
       cpu_used <- mx.cpu()</pre>
153
154
```

```
155
        train_y <- as.factor(as.integer(train_y) %% 2)</pre>
156
        validation_y <- as.factor(as.integer(validation_y) %% 2)</pre>
157
158
        logger_mx <- mx.metric.logger$new()</pre>
159
        train_model <- mx.model.FeedForward.create(neural_net_model,</pre>
160
                                                        X = train_array,
                                                         y = train_y,
161
                                                         eval.data = list(data =
162
                                                            validation_array, label =
                                                             {\tt validation\_y)}, {\tt \#must} be
                                                            set when validating
163
                                                         ctx = cpu_used,
                                                        num.round = 40,
164
                                                         array.batch.size = 50,
165
166
                                                         learning.rate = 0.01,
167
                                                        momentum = 0.9,
168
                                                         initializer = mx.init.Xavier
                                                            (),
                                                        eval.metric = mx.metric.
169
                                                            accuracy,
                                                         epoch.end.callback = mx.
170
                                                            callback.log.train.metric
                                                            (100, logger_mx)
171
172
173
        predicted <- predict(train_model, test_array)</pre>
        \#mxnet gives wrong labels by 1, that is 1-10 instead of 0-9
174
        #fix by subtracting 2
175
        predicted_labels <- max.col(t(predicted)) - 2</pre>
176
177
        return(list(logger_mx, predicted_labels))
178
179 }
180
   plot_error_development <- function(</pre>
181
       neural_net_model,
182
        train_array_val,
183
184
        train_y_val,
185
        test_array,
186
        test_y,
187
        validation_array,
        validation_y
188
189 ) {
190
        # NB! remember to remove "#" from "eval.data" in "run_convolutional_
           neural_network"
        # run cnn to find error in and val
191
        error <- run_convolutional_neural_network(neural_net_model,
192
193
                                                        train_array_val,
194
                                                       train_y_val,
195
                                                       test_array,
                                                       validation_array = validation_
196
                                                           array,
197
                                                       validation_y = validation_y)
198
        # extract error from results
199
        error_t <- error[[1]]
200
201
        error_in_sample <- (1 - error_t$train)</pre>
202
203
        error_val <- (1 - error_t$eval)</pre>
204
```

```
205
        ggplot_df <- data.frame(round = 1:length(error_in_sample),</pre>
206
                                  error_in_sample = error_in_sample,
207
                                  error_val = error_val)
208
209
        # plot error in and validation
210
        ggplot1 <- ggplot(data = ggplot_df, aes(x = round)) +</pre>
            geom_line(aes(y = error_in_sample, colour = "$E_{in}$")) +
211
            geom_line(aes(y = error_val, colour = "$E_{val}$")) +
212
            xlab("$n_{round})$") +
213
            ylab("Miss.class. Error") +
214
            scale_y_continuous(limits = c(0, 1.0)) +
215
216
            scale_colour_manual("Legend",
                                  breaks = c("$E_{in}$", "$E_{val}$"),
217
                                  values = c("#91bfdb", "#fc8d59"),
218
                                  guide = guide_legend(override.aes = list(
219
220
                                       linetype = c("solid", "solid"),
221
                                       shape = c(16, 16)
                                  ))) +
222
223
            theme_bw() +
            theme(legend.position = c(0.8, 0.355),
224
                   legend.background = element_rect(fill=alpha('white', 0)))
225
226
        ggplot_to_latex(ggplot1, paste0(path_to_here, "/results_NN/98_attempt_
            convolutional_neural_network_40_rounds"),
227
                         width = 6, height = 4)
228 }
229
230
   # Predict on test set splitted from the training data to compare with other
       methods
   predict_on_test_set <- function(</pre>
231
232
       neural_net_model,
233
       train_array,
       train_y,
234
235
       test_array,
236
        test_y
237 ) {
       predicted <- run_convolutional_neural_network(neural_net_model,</pre>
238
239
                                                           train_array,
240
                                                           train_y,
241
                                                           test_array)
242
       predicted <- predicted[[2]]</pre>
243
        \verb|create_confusion_matrix(as.factor(predicted), test_y, \\
244
                                  pasteO(path_to_here, "/results_NN/convolutional_
245
                                      neural_network_40_rounds_test"))
246 }
   # Give predictions on unclassified data
   predict_on_unclassified_data <- function(</pre>
249
250
       neural_net_model,
251
        train_array_full,
252
        train_y_full,
253
        unclassified_array
   } {
254
255
       predicted <- run_convolutional_neural_network(neural_net_model,</pre>
256
                                                           train_array_full,
257
                                                           train_y_full,
258
                                                           unclassified_array)
       predicted <- predicted[[2]]</pre>
259
260
```

```
261
        predicted_digit <- predicted</pre>
        predicted_odd_even <- as.factor(as.integer(predicted) %% 2)</pre>
262
263
        prediction_csv <- data.frame(number = 1:length(predicted_digit),</pre>
264
                                       digits = predicted_digit,
265
                                       odd_even = predicted_odd_even)
        write.csv(prediction_csv, file = paste0(path_data, "/data/predictions_
266
            STNKAR012.csv"))
267
268 }
269
270 main <- function()
271
        {
        # Validation used under tuning, set to false so it does run after tuning
272
        validation_boolean <- TRUE</pre>
273
        # Test used under testing of the final tuned model to compare with other
            methods
275
        train_boolean <- FALSE
276
        fully_2 <- two_layer_concolutional_network()</pre>
277
       neural_net_model <- mx.symbol.SoftmaxOutput(data = fully_2)</pre>
278
279
280
        if(validation_boolean){
281
            plot_error_development(neural_net_model,
282
                                      train_array_val,
283
                                      train_y_val,
284
                                      test_array,
285
                                      test_y,
286
                                      validation_array,
287
                                      validation_y)
       }
288
289
        if(train_boolean){
290
291
            predict_on_test_set(neural_net_model,
                                  train_array,
292
293
                                  train_y,
294
                                  test_array,
295
                                  test_y)
       }
296
297
298
        predict_on_unclassified_data(neural_net_model,
299
                                        train_array_full,
300
                                        train_y_full,
301
                                        unclassified_array)
302
303 }
304 main()
```

../R scripts/Neural Networks/convolutional neural network.R

## 6.9 R-Code: K-nearest Neighbours

```
1 ## Libraries and seed
2 rm(list = ls())
3
                        # library for running loop in parallel
4 library (foreach)
5 library(doParallel)
                           # library for running loop in parallel
                           # useful library to split up data set
6 library(caret)
7
  library(tikzDevice)
                           # library to export plots to .tex files
8 library(xtable)
                           # library to export data frames to tables in .tex
      files
9 library(class)
                          # library for knn method
10 set.seed(420)
                           # seed to replicate results and get consistent test
      and training set
11
12 # Load help script with functions to export the results to latex
13 # These functions gathered to avoid duplicate code
14 if (!exists("create_confusion_matrix", mode = "function")) {
      source("Help_Scripts/to_latex_functions.R")
16 }
17
18 #----#
19
20 ## Data
21 path_data <- pasteO(getwd(), "/data")</pre>
22 path_to_here <- paste0(getwd(), "/Tree_Based_Methods")  # getwd give path
     to project
23 # which is one folder over
24
25
26 train_data <- read.csv(paste0(path_data, "/Train_Digits_20171108.csv"),
      header = TRUE)
  unclassified_data <- read.csv(paste0(path_data, "/Test_Digits_20171108.csv")
      , header = TRUE)
28
29 train_data[,1] <- as.factor(train_data[, 1] )
30
31 # split training set into training and test set
32
33 split_train_test <- createDataPartition(train_data$Digit, p = 0.8, list =
34 test_data <- train_data[-split_train_test, ]</pre>
35 train_data <- train_data[split_train_test, ]
36
37
38 k_nearest_neighbors <- function(
      train_data,
39
40
      train_data_norm,
      k_folds,
41
42
      k_neighbors = 20
  ) {
43
      # First find number of k(neighbors) using crossvalidation
44
45
      cv_indexes <- create_cv_indexes(nrow(train_data), k_folds)</pre>
46
      cores=detectCores()
47
      cl <- makeCluster(cores[1]-1) #not to overload your computer</pre>
48
      registerDoParallel(cl)
49
50
      cv_error <- c()</pre>
51
```

```
52
        indexes <- 1:nrow(train_data)</pre>
53
        final_df <- foreach(i = 1:k_neighbors,</pre>
54
55
                              .combine = "rbind"
                              .packages = "class") %dopar%{
56
57
                 error_kfolds <- 0
58
                 error_kfolds_norm <- 0
                 for(k in 1:k_folds){
59
                     train_val <- train_data[!indexes %in% cv_indexes[k, ], ]</pre>
60
                     validation_val <- train_data[cv_indexes[k, ],]</pre>
61
62
63
                     train_val_norm <- train_data_norm[!indexes %in% cv_indexes[k</pre>
                         , ], ]
                     validation_val_norm <- train_data_norm[cv_indexes[k, ],]</pre>
64
65
66
                     # without normalization
67
                     knn_pred_class <- knn(train_val[-1],</pre>
                                              validation_val[-1],
68
                                              train_val[, 1],
69
70
                                              k = i
71
                     # with normalization
72
                     knn_pred_class_norm <- knn(train_val_norm[-1],</pre>
73
                                                    validation_val_norm[-1],
                                                    train_val_norm[, 1],
74
75
                                                    k = i)
76
77
                     error <- 1 - mean(validation_val[, 1] == knn_pred_class)
                     error_norm <- 1 - mean(validation_val_norm[, 1] == knn_pred_
78
                         class_norm)
                     error_kfolds <- error_kfolds + error
79
                     error_kfolds_norm <- error_kfolds_norm + error_norm</pre>
80
81
                 temp_df <- data.frame(k_neighbors = i,</pre>
82
                                          error_kfolds = error_kfolds,
83
84
                                          error_kfolds_norm = error_kfolds_norm)
                 temp_df
85
86
87
        final_df[, 2:3] <- final_df[, 2:3] / k_folds</pre>
88
        stopCluster(cl)
89
        return(final_df)
90
   }
91
   average_cv_error <- function(</pre>
92
93
        train_data,
94
        test_data,
95
        train_data_norm,
96
        test_data_norm,
97
       n_avg
   ) {
98
       k_neighbors <- 20
99
100
       # set up error data frame
101
        error_df <- data.frame(k_neighbors = 1:k_neighbors,</pre>
102
                                  error_kfolds = rep(0, k_neighbors),
103
                                  error_kfolds_norm = rep(0, k_neighbors))
104
105
       # progress bar
106
107
       pb <- txtProgressBar(min = 0, max = n_avg, style = 3)</pre>
108
```

```
109
       # average cv-error over several runs to get more precise measure
110
       for(i in 1:n_avg){
111
            setTxtProgressBar(pb, i)
            error_df[-1] <- error_df[-1] + k_nearest_neighbors(train_data,</pre>
112
113
                                 train_data_norm,
114
                                 2,
115
                                 k_neighbors)[-1]
116
117
       close(pb)
       error_df[-1] <- error_df[-1]/n_avg
118
       ggplot1 <- ggplot(data = error_df, aes(x = k_neighbors)) +</pre>
119
120
            geom_line(aes(y = error_kfolds, colour = "$KNN$")) +
            geom_line(aes(y = error_kfolds_norm, colour = "$KNN normalized$")) +
121
            geom_vline(xintercept = 1, color = "black", linetype = "dotdash") +
122
            xlab("$k$ neighbors") +
123
124
            ylab("Miss.class. Error") +
125
            scale_colour_manual("Legend",
                                 breaks = c("$KNN$", "$KNN normalized$"),
126
                                 values = c("#91bfdb", "#fc8d59"),
127
                                  guide = guide_legend(override.aes = list(
128
                                      linetype = c("solid", "solid"),
129
130
                                      shape = c(16, 16)
                                 ))) +
131
132
            theme_bw() +
133
            theme(legend.position = c(0.8, 0.355),
134
                  legend.background = element_rect(fill=alpha('white', 0)))
135
       ggplot_to_latex(ggplot1,
                         paste0("K_Nearest_Neighbors/Results_KNN/knn_error_
136
                             compare", n_avg),
                         width = 5, height = 5)
137
       return(error_df)
138
139 }
140
   predict_on_test <- function(</pre>
141
142
       best_k,
       train_data,
143
144
       test_data
145 ) {
146
       knn_pred_class <- knn(train_data[-1],</pre>
147
                                     test_data[-1],
148
                                     train_data[, 1],
                                     k = best_k)
149
150
       create_confusion_matrix(predicted_value = knn_pred_class,
151
                                 true_value = test_data[, 1],
152
                                  destination_path = "K_Nearest_Neighbors/Results_
                                     KNN/")
153 }
   #cv_final <- k_nearest_neighbors(train_data, 10)</pre>
155 train_data_norm <- train_data
156 train_data_norm[-1] <- train_data_norm[-1]/255
157 test_data_norm <- test_data
158 test_data_norm[-1] <- test_data_norm[-1]/255
159 #cv_norm <- k_nearest_neighbors(train_data, train_data_norm, 10)
160
161 average_error <- average_cv_error(train_data = train_data,
162
                     test_data = test_data,
163
                     train_data_norm = train_data_norm,
                     test_data_norm = test_data_norm,
165
                     n_avg = 30
```

## 6.10 R-Code: Support Vector Machines

```
1 ## Libraries and seed
2 rm(list = ls())
3 library(e1071)
                           # library to make margins for svm
4 library(caret)
                           # useful library to split up data set
5 library (readr)
6
  set.seed(420)
                           # seed to replicate results and get consistent test
      and training set
8
9
  # Load help script with functions to export the results to latex
10 # These functions gathered to avoid duplicate code
11 if(!exists("create_confusion_matrix", mode = "function")){
      source("Help_Scripts/to_latex_functions.R")
12
13 }
14
  #----#
15
16
17 ## Data
18 path_data <- paste0(getwd(), "/data")</pre>
19 path_to_here <- paste0(getwd(), "/Support_Vector_Machines") # getwd give
      path to project
20 # which is one folder over
21
22 train_data <- read.csv(paste0(path_data, "/Train_Digits_20171108.csv"),
      header = TRUE)
23 unclassified_data <- read.csv(paste0(path_data, "/Test_Digits_20171108.csv")
      , header = TRUE)
24
25
  train_data[,1] <- as.factor(train_data[, 1])</pre>
26
  nzr <- nearZeroVar(train_data[,-1], saveMetrics = TRUE, freqCut = 10000/1,</pre>
27
      uniqueCut = 1/7)
28 sum(nzr$zeroVar)
29
30 sum(nzr$nzv)
31
32 cut_variables <- rownames(nzr[nzr$nzv == TRUE, ])
33 variable <- setdiff(names(train_data), cut_variables)</pre>
34 train_data <- train_data[, variable]
36 # split data into test and training set
37 split_train_test <- createDataPartition(train_data$Digit, p = 0.8, list =
      FALSE)
38 test_data <- train_data[-split_train_test, ]</pre>
39 train_data <- train_data[split_train_test, ]
40
  # remove label temporarely by storeing as it's own vector
41
42 digit <- train_data[1]
43 train_data$Digit <- NULL
  train_data <- train_data/255</pre>
45 cov_train <- cov(train_data)
46
47 digit_test <- test_data[1]
48 test_data$Digit <- NULL
49 test_data <- test_data/255
50
51 # use prcomp to do PCA on the covariance matrix
```

```
52 train_pc <- prcomp(cov_train)
53
   # get information of about the variance
   var_explained <- train_pc$sdev^2/sum(train_pc$sdev^2)</pre>
   var_explained_cumsum <- cumsum(var_explained)</pre>
57
   var_explained_df <- data.frame(number = 1:length(train_pc$sdev),</pre>
58
                                      variance_explained = var_explained,
                                      cumsum = var_explained_cumsum)
59
   # see plot in end of script
60
61
62
63 # find optimal number of components
64 find_number_of_components <- function(
65
        components_range,
66
        increase_by
   ) {
67
        components <- seq(from = components_range[1],</pre>
68
69
                            to = components_range[2],
                            by = increase_by)
70
        missclassification_error <- rep(0, length(components))</pre>
71
        cv_error <- rep(0, length(components))</pre>
72
73
74
        components_error_df <- data.frame(components,</pre>
75
                                             missclassification_error,
76
                                             cv_error)
77
78
        for(i in 1:length(components)){
79
            # traing the svm for the components given
80
            train_score <- as.matrix(train_data) %*% train_pc$rotation[,1:</pre>
81
                components[i]]
            train_data_temp <- cbind(Digit = digit, as.data.frame(train_score))</pre>
82
83
            tune_svm <- tune.svm(Digit ~ ., data = train_data_temp, cost = 1:10,</pre>
84
                 kernal = "radial")
            fit_svm <- tune_svm$best.model
85
86
87
            # review the performance
88
29
            predicted <- predict(fit_svm, train_score, type = "response")</pre>
            missclassification_error <- 1 - mean(train_data_temp[, "Digit"] ==
90
                predicted)
91
            components_error_df[i, "cv_error"] <- tune_svm$best.performance
components_error_df[i, "missclassification_error"] <-</pre>
92
93
                missclassification_error
94
95
        return(components_error_df)
96
97
   # make ggplot of cv error for different amount of components
98
   # also marks selected number of components manually (22 in this task)
99
   plot_components_error <- function(</pre>
100
101
        components_error_df
102 ) {
        components_chosen <- components_error_df[22, ]</pre>
103
104
        ggplot1 <- ggplot(data = components_error_df, aes(x = components)) +</pre>
106
            geom_line(aes(y = cv_error, colour = "$E_{CV}$")) +
```

```
107
            geom_point(aes(y = cv_error, colour = "$E_{CV}$")) +
108
            geom_vline(xintercept = 22, color = "black", linetype = "dotdash") +
            xlab("$n_{components}$") +
109
            ylab("Miss.class. Error") +
110
111
            geom_point(data = components_chosen,
112
                        aes(x = components, y = cv_error),
                        color = "red") +
113
            geom_text(data = components_chosen,
114
                       aes(y = cv_error,
115
                           label = paste0("\mbox{"}\mbox{mathrm}{CV}_{error}(22) = ",
116
117
                                            round(cv_error,4), "$")),
118
                       hjust = 0.2, vjust = -1.0) +
            scale_colour_manual("Legend",
119
                                  breaks = c("$E_{CV}$"),
120
                                  values = c("black"),
121
122
                                  guide = guide_legend(override.aes = list(
                                      linetype = c("solid"),
123
124
                                      shape = c(16)
                                  ))) +
125
            theme_bw() +
126
            theme(legend.position = c(0.8, 0.255),
127
                  legend.background = element_rect(fill=alpha('white', 0)))
128
       ggplot_to_latex(ggplot1,
129
130
                         paste0(path_to_here,
131
                                 '/Results_SVM/number_of_components_cv_error"),
132
                         width = 5, height = 5)
133
134
   # Calculate best cost parameter for svm
135
136 | find_optimal_cost_for_components <- function(
       selected_components,
137
       cost_range,
138
       cost_increase,
139
140
       n_avg
141 ) {
       train_score <- as.matrix(train_data) %*% train_pc$rotation[,1:selected_</pre>
142
           componentsl
143
       train_data_temp <- cbind(Digit = digit, as.data.frame(train_score))</pre>
144
145
        cost <- seq(from = cost_range[1], to = cost_range[2], by = cost_increase
           )
146
       cost_error <- data.frame(cost = cost, cv_error = rep(0, length(cost)))</pre>
147
148
       for(i in 1:n_avg){
            tune_svm <- tune.svm(Digit ~ ., data = train_data_temp, cost = cost,</pre>
149
                 kernal = "radial")
150
            cv_error <- tune_svm$performances$error</pre>
151
            cost_error[, "cv_error"] <- cost_error[, "cv_error"] + cv_error</pre>
152
153
154
       cost_error[, "cv_error"] <- cost_error[, "cv_error"]/n_avg</pre>
155
156
       best_cost <- cost_error[which(cost_error[, "cv_error"] == min(cost_error</pre>
157
           [, "cv_error"])),]
158
159
       ggplot1 <- ggplot(data = cost_error, aes(x = cost)) +</pre>
            geom_line(aes(y = cv_error, colour = "$E_{CV}$")) +
160
            geom_point(aes(y = cv_error, colour = "$E_{CV}$")) +
161
```

```
162
            geom_vline(xintercept = best_cost[, "cost"],
                        color = "black", linetype = "dotdash") +
163
164
            xlab("$n_{components}$") +
165
            ylab("Miss.class. Error") +
166
            geom_point(data = best_cost,
167
                        aes(x = cost, y = cv_error),
                        color = "red") +
168
            geom_text(data = best_cost,
169
                       aes(y = cv_error,
170
                           label = paste0("$\\mathrm{CV}_{error}(",
171
                                            cost,
172
                                            ") = ",
173
                                            round(cv_error,4), "$")),
174
175
                       hjust = -0.05, vjust = 0.9) +
            scale_colour_manual("Legend",
176
                                  breaks = c("$E_{CV}$"),
177
178
                                  values = c("black"),
                                  guide = guide_legend(override.aes = list(
179
                                      linetype = c("solid"),
180
181
                                      shape = c(16)
                                  ))) +
182
183
            theme_bw() +
184
            theme(legend.position = c(0.8, 0.255),
                  legend.background = element_rect(fill=alpha('white', 0)))
185
186
        ggplot_to_latex(ggplot1,
187
                         paste0(path_to_here,
188
                                 "/Results_SVM/cost_cv_error"),
                         width = 5, height = 5)
189
190
        return(best_cost)
191 }
192
193 # Make prediction on the test set and make confusion matrix
194 predict_on_test <- function(
195
        selected_components,
196
        cost
   ) {
197
198
        train_score <- as.matrix(train_data) %*% train_pc$rotation[,1:selected_
           components]
199
        train_data_temp <- cbind(Digit = digit, as.data.frame(train_score))</pre>
200
        test_score <- as.matrix(test_data) %*% train_pc$rotation[,1:selected_</pre>
201
           components
        test_data_temp <- cbind(Digit = digit_test, as.data.frame(test_score))</pre>
202
203
       svm_predict <- svm(Digit ~ ., data = train_data_temp, cost = cost,</pre>
204
           kernal = "radial")
205
206
       predicted <- predict(svm_predict, test_data_temp)</pre>
207
        create_confusion_matrix(predicted_value = predicted,
208
                                  true_value = test_data_temp[, "Digit"],
209
210
                                  destination_path = paste0(path_to_here,
                                                               "/Results_SVM/
211
                                                                  confusion_matrix")
                                                                  )
212
213 }
214
215 # some help-plots for pca
```

```
216 plot_variance_explained <- function(
217
       var_explained_df,
218
       chosen_number_components
219
   ) {
       # plot amount of varibles and how much of the variance this describe
220
       ggplot1 <- ggplot(data = var_explained_df[1:100,], aes(x = number, y =</pre>
221
           cumsum)) +
           geom_line() +
222
           geom_point() +
223
           geom_vline(xintercept = chosen_number_components,
224
225
                       color = "black", linetype = "dotdash") +
226
           xlab("$n_{components}$") +
           ylab("Variance Explained") +
227
228
           theme_bw() +
           theme(legend.position = c(0.8, 0.255),
229
230
                  legend.background = element_rect(fill=alpha('white', 0)))
231
       # plot the data as described by the two first principle components
232
233
       train_score <- as.matrix(train_data) %*% train_pc$rotation[,1:chosen_</pre>
234
           number_components]
235
       train_data_temp <- cbind(Digit = digit, as.data.frame(train_score))</pre>
236
237
       ggplot2 <- ggplot(data = train_data_temp, aes(x = PC1, y = PC2)) +</pre>
238
           geom_point(aes(colour = Digit)) +
           xlab("PCA 1") +
239
           ylab("PCA 2") +
240
           241
242
                                    fb8072',
                                             '#80b1d3','#fdb462','#b3de69','#
243
                                                fccde5'.
                                             '#d9d9d9','#bc80bd')
244
                                 ) +
245
246
           theme_bw() +
           theme(legend.position = c(0.9545, 0.355),
247
248
                  legend.background = element_rect(fill=alpha('white', 0)))
249
250
       ggplot_to_latex(ggplot1,
251
                        paste0(path_to_here,
                                "/Results_SVM/variance_explained_pca"),
252
253
                        width = 5, height = 5)
254
       ggplot_to_latex(ggplot2,
255
256
                        paste0(path_to_here,
                                "/Results_SVM/map_pca1_pca2"),
257
258
                        width = 5, height = 5)
259
260
   main <- function(){</pre>
261
       # Find best number of components to use with svm
262
       components_range <- c(1, 30)</pre>
263
264
       increase_by <- 1
       components_error_df <- find_number_of_components(components_range =</pre>
265
           components_range,
266
                                                           increase_by = increase_
                                                               by)
       # Plot best cv error over number of components
267
268
       plot_components_error(components_error_df)
```

```
269
270
       selected_components <- 22
271
272
       # Find optimal cost parameter for the svm function
273
       best_cost <- find_optimal_cost_for_components(cost_range = c(0.5, 10),</pre>
274
275
                                                         cost_increase = 0.5,
276
                                                         selected_components =
                                                             selected_compnents,
                                                         n_avgs = 5)
277
       # Plot pca variance and to principle components
278
279
       plot_variance_explained(var_explained_df, selected_components)
280 }
281
282 main()
```

## 6.11 R-Code: Visualize data examples

```
## Data
2
3
  path_data <- paste0(getwd(), "/data")</pre>
4 path_to_here <- paste0(getwd(), "/Support_Vector_Machines") # getwd give
      path to project
5
  train_data <- read.csv(paste0(path_data, "/Train_Digits_20171108.csv"),</pre>
      header = TRUE)
  train_images <- train_data[, -1]</pre>
10 rotate <- function(x) t(apply(x, 2, rev))
11
12 # plot 5 x 8 images from the data set
png(paste0(path_data,"/images_digits.png"))
14 opar <- par(no.readonly = T)
16 par(mfrow = c(5, 8), mar = c(.1, .1, .1, .1))
17 for(i in 1:40){
       im_matrix <- matrix(train_images[i, ], nrow = 28, ncol = 28)</pre>
18
      im_matrix <- apply(im_matrix, c(1, 2), function(x) as.numeric(x))</pre>
19
20
      im_matrix <- rotate(im_matrix)</pre>
      image(im_matrix, col = grey.colors(255), axes = F)
21
22 }
23 par(opar)
24 dev.off()
                          ../R\_scripts/Help\_Scripts/print\_images.R
```