EXAM SL

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imports

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```
library(dplyr)
library(ggplot2)
library(purrr)
library(caret)
library(class)
library(MASS)
library(tidyverse)
library(nnet)
library(glmnet)
library(glmnet)
library(reticulate)
library(ncvreg)
library(e1071)
```

Q1

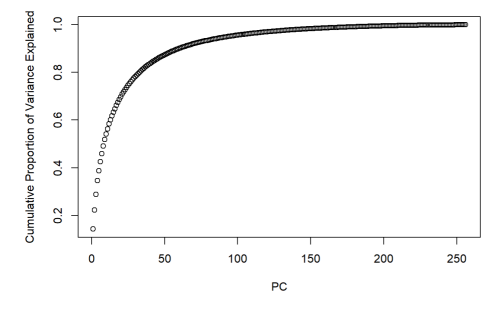
loading the data:

```
df_train <- read.table("zip.train") %>% rename("y" = "V1") %>%
    filter(y %in% c(2,3,5))
df_test <- read.table("zip.test") %>% rename("y" = "V1") %>%
    filter(y %in% c(2,3,5))
X_train = df_train[,2:257]
y_train = df_train[,1]
X_test = df_test[,2:257]
y_test = df_test[,1]
```

1.a

using PCA

```
pca <- prcomp(X_train)
plot(cumsum(pca$sdev^2/sum(pca$sdev^2)),
    xlab = "PC",
    ylab = "Cumulative Proportion of Variance Explained")</pre>
```



i will take 100 PCS which explain ~ 95% of the variance in the data:

```
X_pca_train_100 = as_tibble(pca$x[,1:100])
X_pca_test_100 = as_tibble(predict(pca,X_test)[,1:100])
```

building the models:

```
lda_full = lda(y~., df_train)
df_pca_train_100 = X_pca_train_100 %>% mutate(y=y_train)
lda_pca = lda(y~., df_pca_train_100)
```

getting results:

```
get_results = function(model, X, y_true, lda=TRUE, neural_model=FALSE){
 # this function returns the confusion matrix and miss classification error of a given model
 y_pred = predict(model, X)
 if (lda) {
    y_pred = y_pred$class
 if (neural_model){
   y_pred = apply(y_pred,1,which.max)
   y_pred[y_pred==1] = 5
  classification_err = mean(y_pred != y_true)
 conf_matrix = confusionMatrix(as.factor(y_true), as.factor(y_pred))
 return( list("classification_err" = classification_err,
              "conf_matrix" = conf_matrix$table))
}
lda_train_results = get_results(lda_full, X_train, y_train, TRUE)
lda_test_results = get_results(lda_full, X_test, y_test, TRUE)
lda_pca_train_results = get_results(lda_pca, X_pca_train_100, y_train, TRUE)
lda_pca_test_results = get_results(lda_pca, X_pca_test_100, y_test, TRUE)
```

we can see that both the model with the PCA and the model without the PCA preform rather similar on the test set (9.5% error) but the model on the full data gets a better result on the train set.

lets take a look at the confusion matrix of the full LDA

train:

```
## Reference
## Prediction 2 3 5
## 2 719 10 2
## 3 7 644 7
## 5 3 16 537
```

we can see that the model struggles more to distinguish between 5 and 3, after that it has a hard time with 3 and 2, and can separate well enough 5 and 2

test:

```
## Reference

## Prediction 2 3 5

## 2 184 10 4

## 3 6 148 12

## 5 0 18 142
```

we can see in general the same behavior as in the train set, but with larger percentage of mistake, notice that the test set is smaller then the train set and that we have higher values outside the diagonal (which indicate false prediction)

1.b

building the models:

```
multinom_full = multinom(y~., df_train)
```

```
## # weights: 774 (514 variable)
## initial value 2136.800901
## iter 10 value 400.032251
## iter 20 value 139.626891
## iter 30 value 56.619769
## iter 40 value 24.791592
## iter 50 value 15.917622
## iter 60 value 8.683908
## iter 70 value 3.882552
## iter 80 value 0.925665
## iter 90 value 0.066039
## final value 0.066039
## stopped after 100 iterations
```

```
multinom_pca = multinom(y~., df_pca_train_100)
```

```
## # weights: 306 (202 variable)

## initial value 2136.800901

## iter 10 value 192.871780

## iter 20 value 97.706870

## iter 30 value 54.966766

## iter 40 value 43.944227

## iter 50 value 41.985445

## iter 60 value 40.308383

## iter 70 value 37.790193

## iter 80 value 27.292217

## iter 90 value 0.983953

## iter 100 value 0.013846

## final value 0.013846

## stopped after 100 iterations
```

getting results:

```
multinom_train_results = get_results(multinom_full, X_train, y_train, FALSE)
multinom_test_results = get_results(multinom_full, X_test, y_test, FALSE)
multinom_pca_train_results = get_results(multinom_pca, X_pca_train_100, y_train, FALSE)
multinom_pca_test_results = get_results(multinom_pca, X_pca_test_100, y_test, FALSE)
```

clearly we overfit the training data, we can see a train error of 0, which means that in the train set we are able to classify each observation to the correct y value, but for the test set we have much larger errors 14.5% for the full multi-class logistic regression, and 9.35% for the multi-class logistic regression with PCA. we see that in this case that the PCA helps improve the results of the model, also the multi-class logistic regression with PCA has similar results to the LDA

lets take a look at the confusion matrix of the PCA multi-class logistic regression the train has perfect score thus the confusion matrix is diagonal and not that interesting, lets see the test confusion matrix:

```
## Reference
## Prediction 2 3 5
## 2 184 10 4
## 3 6 146 14
## 5 4 11 145
```

we can see a rather similar performance, the model struggles the most with 5 and 3, after that with 3 and 2, and is rather ok with 5 and 2.

1.c

we are using images, and as we know the SOTA for image predictive modeling are CNNS so lets make a CNN to predict the class first lets prepare the data: - the loss expects values to be between 0-2, so i will re encode $5 \Rightarrow 0$, $2 \Rightarrow 1$, $3 \Rightarrow 2$

```
X_train_net = array_reshape(as.matrix(X_train), c(nrow(X_train),16,16,1))
y_train_net = y_train-1
y_train_net[y_train_net == 4] = 0

X_test_net = array_reshape(as.matrix(X_test), c(nrow(X_test),16,16,1))
y_test_net = y_test-1
y_test_net[y_test_net == 4] = 0
```

now create the model

```
cnn_model <- keras_model_sequential() %>%
  layer_conv_2d(filters = 16, kernel_size = c(3, 3), activation = "relu", input_shape = c(16, 16, 1)) %>%
  layer_conv_2d(filters = 32, kernel_size = c(3, 3), activation = "relu") %>%
  layer_max_pooling_2d(pool_size = c(2, 2)) %>%
  layer_flatten() %>%
  layer_batch_normalization() %>%
  layer_dense(units = 32, activation = "relu") %>%
  layer_dropout(rate = 0.2) %>%
  layer_dense(units = 16, activation = "relu") %>%
  layer_dense(units = 16, activation = "relu") %>%
  layer_dropout(rate = 0.1) %>%
  layer_dense(units = 3, activation = "softmax")
```

compile and train the model

getting the results:

```
cnn_train_res = get_results(cnn_model, X_train_net, y_train, FALSE, TRUE)
cnn_test_res = get_results(cnn_model, X_test_net, y_test, FALSE, TRUE)
tibble(model_dataset = c("neural net train", "neural net test"),
    err = c(cnn_train_res$classification_err, cnn_test_res$classification_err))
```

we can see that this model is far better then the ones before, also it still has a much lower error in the train set, we can keep playing with the paramaters of the model (add some more regularization or use data augmentation) to get even better results

lets take a look at the test confusion matrix

```
## Reference
## Prediction 2 3 5
## 2 192 3 3
## 3 2 154 10
## 5 0 3 157
```

we can see that the results are much better, we still some difficulties with the 5's which is most challenging to classify.

lets improve the model by using some more modern techniques like data augmentation data augmentation "creates" new images by applying several transformations on the image and thus improves the generalization of the model by introducing new observations.

in this case i choose to apply:

rotation - rotates the images

width, height shifts - which applies translation on the image to left/right, up/down

zoom - which zooms in/ out on an image

all of this transformation are rather reasonable by the sense hat applying them wont change the image much but they are still valid drown numbers. it doesn't really matter if you draw a number in the middle of the box or a bit shifted up/down/left/right, also the scale does not matter, it the same if you draw a small number or a larger number, and also the same applies for rotation

```
datagen <- image_data_generator(rotation_range = 10,width_shift_range = 0.1,height_shift_range = 0.1,zoom_range = 0.1)

train_generator <- flow_images_from_data(X_train_net,y_train_net,datagen,32)

cnn_model_aug <- keras_model_sequential() %>%
    layer_conv_2d(filters = 16, kernel_size = c(3, 3), activation = "relu", input_shape = c(16, 16, 1)) %>%
    layer_conv_2d(filters = 16, kernel_size = c(3, 3), activation = "relu") %>%
    layer_max_pooling_2d(pool_size = c(2, 2)) %>%
    layer_flatten() %>%
    layer_flatten() %>%
    layer_dense(units = 32, activation = "relu") %>%
    layer_dense(units = 16, activation = "relu") %>%
    layer_dense(units = 16, activation = "relu") %>%
    layer_dense(units = 3, activation = "softmax")
```

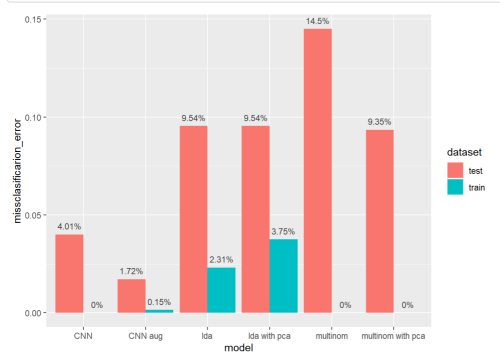
compile and train

get results

```
cnn_aug_train_res = get_results(cnn_model_aug, X_train_net, y_train, FALSE, TRUE)
cnn_aug_test_res = get_results(cnn_model_aug, X_test_net, y_test, FALSE, TRUE)
tibble(model_dataset = c("CNN aug train", "CNN aug test"),
        err = c(cnn_aug_train_res$classification_err, cnn_aug_test_res$classification_err))
```

the over all results:

```
## # A tibble: 6 × 3
##
   model_dataset
                       err_train err_test
##
                          <dbl>
    <chr>
                                   <dbl>
## 1 neural net aug
                         0.00154
                                  0.0172
                                  0.0401
## 2 neural net
                        0
## 3 multinom with pca
                        0
                                  0.0935
## 4 lda
                         0.0231
                                  0.0954
## 5 lda with pca
                         0.0375
                                  0.0954
## 6 multinom
                                   0.145
```



we can see from the full results that the multinom and neural net both have perfect score on the train set, also the neural net with augmentation has the best results on the test set, and even better results on the test set then other models get on the train set!

Q2

2a.i

x has many categorical variables and missing values, accuracy is critical

good models can be - random forest and gradient boosting, both models have lower variance due to the fact that they average many predictors which will result in better accuracy, also both of them can handle missing values, they should work well for regression and classification.

bad models can be: Linear regression - LR will probably preform not that good as it can't handle missing values well. also K-nn wont work that well for both classification and regression as it cannot handle missing values, also the nature of the problem is in high dimension and we know that the meaning of distance in high dimension is impaired.

2a.ii

x has many categorical variables and missing values, speed of model building is critical

good: Two models that can be good given the proper processing of missing values can be Logistic regression for classification and Linear regression for regression problems. they are both fast to train and run inference, but both will have a hard time with the high dimension so we can use lasso as a feature selection technique and thus reduce the dimension. *a proper way for processing the missing values could be to set a new category "missing" for each categorical feature with missing values, after that we can use one hot encoding. it might even surprise us with very good predictions if the data is not missing at random and might have a connection to our target.

bad models: Random forest, Boosting, both these models can achieve high accuracy and tree based models can handle missing values as well, but they will take a larger time to train and also run inference, each observation must have prediction from all the trees and usually we want a lot of trees to be accurate so the total run time will be rather long. K-nn will also not be that good here as for each new observation we need to calculate the distance to all points in the train set in order to find the K closest ones, and also it will not be good from the reason listed above.

2a.iii

Large n, small p, you believe that there are complex dependencies between x and y (not only main effects / additive / low order)

good - gradient boosting, random forest, deep learning and kernel machines can model complex dependencies between x and y thus all of them will be good in this case, also because we have a large number of samples and not many variables they will likely perform well without any regularization. the tree based models can capture complex dependencies as they split the sub space, neural nets can capture complex dependencies by using nonlinear activation functions, and kernel machines can capture complex dependencies by the nature of increasing the dimension

bad - Linear regression for regression, Logistic regression for classification, LDA for classification, all of these models assume some form of linearity thus will not be cable to capture complex dependencies between x and y and will probably preform poorly.

2a.iv

Small n, large p, you believe only a small number of variables are important

Good - Lasso: will help reduce the dimension of the problem by setting some of the coefficients to zero and keeping the important coefficients. We can use lasso as a model, but also as a feature selection technique and then use all of the other models with the selected features. We can also use PCA to reduce p and afterwards we can use all the models (if we can find a d dimensional subspace where d<<n)

Bad - linear regression will be very bad due to the fact that X^tX has no inverse. also without any regularization we can't expect any of the other models to perform well when p>>n

2b

LDA assumption: we assume that $X|Y_i \sim N(\mu_i, \Sigma)$

LR assumption: the model that we have for the linear regression is y=Xeta+arepsilon we assume that $arepsilon\sim N(0,\sigma^2)$

Ridge assumption: Ridge assumes a normal prior for eta meaning: $eta \sim N(0, au^2 I)$

Q3

3.a

the ridge regression model yields $\hat{y} = X \hat{eta}_{ridge} = X (X^T X + \lambda I)^{-1} X^T Y$

setting $S(X,\lambda)=X(X^X+\lambda I)^-1X^T$ we get that ridge regression follows the first condition

3.b

for clearer writing i will use the notation k instead of i_0

$$egin{aligned} ext{denote } \hat{eta} = argmin_{eta} \sum_i (y_i^{(-k)} - ((X^{(-k)})^t eta)_i)^2 + \lambda \sum_j eta_j^2 = argmin_{eta} L_k \ & \hat{ ilde{eta}} = argmin_{eta} \sum_i (ilde{y}_i - X_i^t eta)^2 + \lambda \sum_j eta_j^2 \end{aligned}$$

lets break down the expression above

$$\sum_{i} (ilde{y}_{i} - X_{i}^{t}eta)^{2} + \lambda \sum_{j} eta_{j}^{2} = \sum_{i
eq k} (y_{i} - X_{i}^{t}eta)^{2} + (\hat{y}_{k}^{(-k)} - X_{k}^{t}eta)^{2} + \lambda \sum_{j} eta_{j}^{2} = \sum_{i
eq k} (y_{i} - X_{i}^{t}eta)^{2} + (X_{k}^{T}\hat{eta} - X_{k}^{t}eta)^{2} + \lambda \sum_{j} eta_{j}^{2} = \sum_{i
eq k} (y_{i} - X_{i}^{t}eta)^{2} + (X_{k}^{T}(\hat{eta} - eta))^{2} + \lambda \sum_{j} eta_{j}^{2} = L_{k} + (X_{k}^{T}(\hat{eta} - eta))^{2}$$

thus
$$\hat{ ilde{eta}} = argmin_{eta} L_k + (X_k^T (\hat{eta} - eta))^2$$

the argmin of L_k is $\hat{\beta}$ and if we plug it in the formula above we get $L_k(\hat{\beta})$ which achives the minimum of the equation above

thus we can conclude that
$$\hat{ ilde{eta}}=\hat{eta}$$
 $\hat{\hat{y}}_k=S(ilde{y})_k=X_k ilde{eta}=X_k\hat{eta}=\hat{y}_k^{(-k)}$

3.c

we saw in calss that for a linear model (like ridge regression as we proved above) the optimism is: $\frac{2\sigma^2 tr(S)}{n}$

thus the smaller the diagonal elements are the smaller tr(S) becomes and the optimisim is smaller

$$S(X,\lambda)_{ii} = (X(X^TX + \lambda I)^{-1}X^T)_{ii}$$

using the SVD decomposition we can derive:

$$X=UDV^t$$
 $X^tX=VD^2V^t$
 $(X^tX)^{-1}=VD^{-2}V^t$
 $(X^tX+\lambda I_p)^{-1}=V(D^2+\lambda I_p)^{-1}V^t$
 $X(X^tX+\lambda I_p)^{-1}X^t=UD(D^2+\lambda I_p)^{-1}DU^t=D^2(D^2+\lambda I_p)^{-1}$
 $S(X,\lambda)_{ii}=D^2_{ii}(D^2_{ii}+\lambda I_p)^{-1}$
thus $S(X,\lambda)_{ii}$ is a decreaceing funtion of λ

3.d

i will show it formally, by differentiating the expression

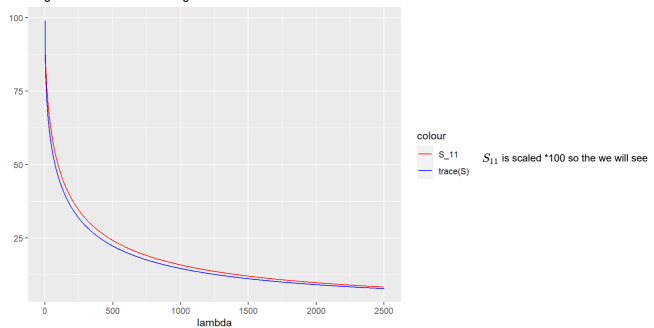
$$rac{\partial D_{ii}^2 (D_{ii}^2 + \lambda I_p)^{-1}}{\lambda} = -D_{ii}^2 (D_{ii}^2 + \lambda I_p)^{-2} < 0 \quad orall \lambda$$

thus we can conclude that $S(X,\lambda)_{ii}$ is indeed a decreaceing funtion of λ

and lets also show by simulation:

```
con <- url("http://www.tau.ac.il/~saharon/StatsLearn2022/train_ratings_all.dat")
X <- tibble(read.table(con))
con <- url("http://www.tau.ac.il/~saharon/StatsLearn2022/train_y_rating.dat")
y <- read.table(con)
X = as.matrix(X)[1:100,1:99]
S_lambda = function(X, lambda){
    return(X %*% solve(t(X) %*% X + lambda * diag(1,99,99)) %*% t(X))
}
lambdas = seq(0,2500,1)
s_11 = c()
trace_s = c()
for (lambda in lambdas){
    S_mat = S_lambda(X, lambda)
    s_11 = c(s_11, S_mat[1,1])
    trace_s = c(trace_s, sum(diag(S_mat)))
}</pre>
```

diagonal and trace behavior given lambda



the relation

3.e

we would not expect the lemma to hold for the lasso penelty the solution for the lasso problem does not hold for the first condition this is because we cannot derive a closed form for lasso regression, and thus cannot conclude it is a linear model

3.f

the lemma does not hold for the K-NN regression, the K-NN is a linear model but the second condition does not hold.

$$N_K(x_m):=\text{the K neighbours of }x_m$$

$$N_K'(x_m):=\text{the K-1 neighbours of }x_m(\text{that is }N_K(x_m)\text{whitout }x_m)$$

$$\begin{split} \hat{y_m} &= \sum_{x_j \in N_K(x_m)} \frac{1}{K} \tilde{y}_j = \frac{1}{K} [(\sum_{x_j \in N_K'(x_m)} y_j) + \hat{y}_m^{(-m)}] \\ \hat{y}_m^{(-m)} &= \sum_{x_j \in N_{K+1}'(x_m)} \frac{1}{K} y_j = \frac{\hat{y}_m^{(-m)} - \hat{y}_m^{(-m)}}{K} + \frac{1}{K} \sum_{x_j \in N_{K+1}'(x_m)} y_j = \frac{\hat{y}_m^{(-m)} - \hat{y}_m^{(-m)}}{K} + \frac{y_l}{K} + \frac{1}{K} \sum_{x_j \in N_K'(x_m)} y_j = \\ &= [(\sum_{x_j \in N_K'(x_m)} y_j) + \hat{y}_m^{(-m)}] + \frac{y_l - \hat{y}_m^{(-m)}}{K} = \hat{y_m} + \frac{y_l - \hat{y}_m^{(-m)}}{K} \end{split}$$

 y_l is the K+1 neighbour

thus we get that for all m: $\hat{y}_m^{(-m)} = \hat{y}_m$ $\iff y_l = \hat{y}_m^{(-m)} \iff y$ is a vector of the same values thus we can conclude that if y has more then one unique value the K-NN model does not supply the 2nd condition

Q4

4.a

X is of full rank, a rank, a matrix of p*n matrix is bounded by: $rank(X) \leq min(n,p)$ thus in our setup where $n \leq p, rank(X) = n$

from an algebraic view we know that for a system of linear equations with n equations and p variables where: p>n, we have an infinite number of solutions. thus clearly the solution is not unique. (in our case we have p+1 variables including the intercept)

this conclusion does not change with the loss. if $\sum_i (y_i - X_i^t \beta - \beta_0)^2 = 0$ then $y_i - X_i^t \beta - \beta_0 = 0$ $\forall i$ also if $\sum_i |y_i - X_i^t \beta - \beta_0| = 0$ then $y_i - X_i^t \beta - \beta_0 = 0 \ \forall i \Rightarrow$ we have an infinite number of solutions for the quantile loss:

$$\sum_{i} \tau(y_i - X_i^t \beta - \beta_0) I(y_i > X_i^t \beta - \beta_0) - (1 - \tau) (y_i - X_i^t \beta - \beta_0) I(y_i < X_i^t \beta - \beta_0) = 0$$
 notice the

 $\sum_{i}\tau(y_{i}-X_{i}^{t}\beta-\beta_{0})I(y_{i}>X_{i}^{t}\beta-\beta_{0})-(1-\tau)(y_{i}-X_{i}^{t}\beta-\beta_{0})I(y_{i}\leq X_{i}^{t}\beta-\beta_{0})=0 \text{ notice that:} \\ \tau(y_{i}-X_{i}^{t}\beta-\beta_{0})I(y_{i}>X_{i}^{t}\beta-\beta_{0})\geq0 \text{ , } -(1-\tau)(y_{i}-X_{i}^{t}\beta-\beta_{0})I(y_{i}\leq X_{i}^{t}\beta-\beta_{0})\geq0 \text{ thus we sum over non-negative number so in order for the sum to be zero each element need to be zero so we are left with: } y_{i}-X_{i}^{t}\beta-\beta_{0}=0 \text{ } \forall i \Rightarrow \text{ we have an infinite number of }$ solutions

to conclude all the above losses have an infinite number of solutions in the case were $p \geq n$

4.b

assume in contradiction that we do not have a unique solution, thus

$$egin{aligned} L &= \sum_{i=1}^n (y_i - X_i^t eta - eta_0)^2 + \lambda \sum_{j=1}^p eta_j^2 \ orall j &\geq 1: rac{\partial L}{\partial eta_j} = 2 \sum_i (y_i - X_i^t eta - eta_0) X_{ij}^t + 2 eta_j \end{aligned}$$

now let $\tilde{\beta}$ be an interpulation point and plug it in the derevetive above we get:

$$rac{\partial L}{\partial eta_j}(ilde{eta}) = 0 + ilde{eta}_j = 0 \iff ilde{eta}_j = 0$$

thus we have that for any interpolation point $\tilde{eta}_j=0\ orall j\geq 1,\ ext{thus}\ y_i-X_i^t ilde{eta}- ilde{eta}_0=y_i- ilde{eta}_0$

but this is an interpolation point thus $y_i = \tilde{\beta}_0 \forall i \Rightarrow y$ is a vector of constants thus if y is not a vector of constants we can see that there are no interpolation points thus the optimal solution is not an interpolation point

4.c

let
$$ilde{eta}$$
 be an optimal solution to $min\sum_{j=1}^p eta_j^2 \ s. \ t \sum_{i=1}^n (y_i - X_i^t eta - eta_0)^2 = 0$

the objective of this minimization problem is the L2 norm squared $\sum_{j=1}^p \beta_j^2 = ||\beta||_2^2$, so the β that minimized it also minimize the L2 norm squared β

$$L(\tilde{\beta}) = \sum_{i=1}^n (y_i - X_i^t \tilde{\beta} - \tilde{\beta}_0)^2 + \lambda \sum_{j=1}^p \tilde{\beta}_j^2 = 0 + \lambda ||\tilde{\beta}||_2^2 \leq \lambda ||\beta||_2^2 \quad \forall \beta \ s. \ t \sum_{i=1}^n (y_i - X_i^t \beta - \beta_0)^2 = 0$$

thus $\tilde{\beta}$ is the optimal solution to the ridge problem over all interpolation points

4.d

$$orall \lambda > 0 ext{ let } \hat{eta}(\lambda) = argmin_{eta} \sum_{i=1}^n (y_i - X_i^t eta - eta_0)^2 + \lambda \sum_{j=1}^p eta_j^2$$

from the previous results we know:

1.
$$\hat{\beta}(\lambda)$$
 is unique

2. $\hat{\boldsymbol{\beta}}^{(l2)}$ has the minimal norm over all the interpolation points and also has the smalleset ridge penelty

as
$$\lambda$$
 converge to 0: $\hat{\beta}(\lambda)$ converges to some $argmin_{\beta}\sum_{i=1}^{n}(y_{i}-X_{i}^{t}\beta-\beta_{0})^{2}$

meaning that $\hat{\beta}(0)$ is an interpolation point

so exsists ε close to zero such that for each $\lambda < \varepsilon$ exsits an interpolation solution

thus for such λ , $\hat{\beta}(\lambda)$ is an interpolation point, and due to the optimality it also has the minimal norm $\Rightarrow ||\hat{\beta}^{(l2)}||_2 = ||\hat{\beta}(\lambda)||_2 \, \forall \lambda < \text{recall that } \hat{\beta}(\lambda)$ is a unique solution thus there is only one vector that is an interpolation point and satisfies: $||\hat{\beta}^{(l2)}||_2 = ||\hat{\beta}(\lambda)||_2$

thus we get
$$\hat{eta}(\lambda) \stackrel{\lambda o 0}{\longrightarrow} \hat{eta}^{(l2)}$$

4.e

general idea: Lagrange multipliers

$$\begin{split} & minimize\beta^t\beta\\ & s.\,t:X\beta=y\\ & \text{using lagrange multiplers we get: } L=\beta^t\beta+\lambda^T(X\beta-y)\\ & \frac{\partial L}{\partial\beta}=2\beta+X^T\lambda=0 \iff \beta=-\frac{1}{2}X^T\lambda\\ & \frac{\partial L}{\partial\lambda}=X\beta-y=-X\frac{1}{2}X^T\lambda-y=0 \iff \lambda=-2(XX^t)^{-1}y\\ & \text{plug it back in to }\beta\text{ and we get: } \hat{\beta}=X^T(XX^t)^{-1}y \end{split}$$

4.f

$$\begin{array}{l} \text{in this case: } \hat{\beta}(\lambda) = argmin_{\beta} \sum_{i=1}^{n} (y_i - X_i^t \beta - \beta_0)^2 + \lambda \sum_{j=1}^{p} |\beta_j| \\ \\ \text{and } \hat{\beta}^{(l1)} = argmin \sum_{i=1}^{p} |\beta_j| \ s. \ t \sum_{i=1}^{n} (y_i - X_i^t \beta - \beta_0)^2 = 0 \end{array}$$

in this case the result will be that $\hat{\beta}(\lambda) \stackrel{\lambda \to 0}{\longrightarrow} \hat{\beta}^{(l1)}$

and we would expect the results to not change according to the loss by the same arguments in section 4.a in general for L1 loss, L2 loss and quantile loss the constraints are equivelent to $y_i - X_i^t \beta - \beta_0 = 0 \quad \forall i$

4.g

4.g.i

for some reason i cant load the I1ce package so i will use glmnet.

```
con <- url("http://www.tau.ac.il/~saharon/StatsLearn2022/train_ratings_all.dat")
X <- tibble(read.table(con))
con <- url("http://www.tau.ac.il/~saharon/StatsLearn2022/train_y_rating.dat")
y <- read.table(con)

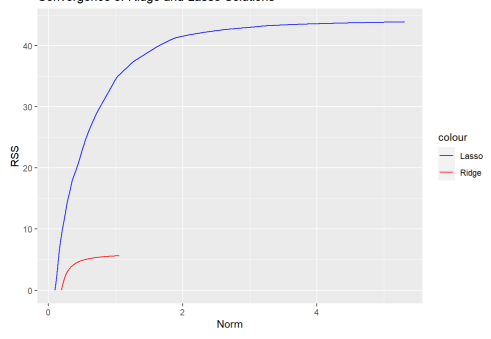
X_small = as.matrix(X[1:50,])
y_small = y[1:50,]</pre>
```

```
ridge_fit = glmnet(X_small, y_small, alpha = 0, lambda = seq(0, 0.5, by = 0.0001))
ridge_rss = colSums((y_small - predict(ridge_fit, s = seq(0, 0.5, by = 0.0001), newx = X_small))^2)
ridge_norm = colSums(coef(ridge_fit)[2:100,]^2)

lasso_fit = glmnet(X_small, y_small, alpha = 1, lambda = seq(0, 0.5, by = 0.0001))
lasso_rss = colSums((y_small - predict(lasso_fit, s = seq(0, 0.5, by = 0.0001), newx = X_small))^2)
lasso_norm = colSums(abs(coef(lasso_fit)[2:100,]))

ggplot() +
    geom_line(aes(x = ridge_norm, y = ridge_rss, color = "Ridge")) +
    geom_line(aes(x = lasso_norm, y = lasso_rss, color = "Lasso")) +
    scale_color_manual(values = c("Ridge" = "red", "Lasso" = "blue")) +
    xlab("Norm") +
    ylab("RSS") +
    ggtitle("Convergence of Ridge and Lasso Solutions")
```

Convergence of Ridge and Lasso Solutions



4.g.i

```
ridge_fit = glmnet(X_small, y_small, alpha = 0, lambda = exp(-15))
ridge_rss = colSums((y_small - predict(ridge_fit, s = exp(-15), newx = X_small))^2)
ridge_norm = sum(coef(ridge_fit)[2:100]^2)

lasso_fit = glmnet(X_small, y_small, alpha = 1, lambda = exp(-15))
lasso_rss = colSums((y_small - predict(lasso_fit, s = exp(-15), newx = X_small))^2)
lasso_norm = sum(abs(coef(lasso_fit)[2:100]))
```

```
## [1] "lasso norm is: 8.693 ridge norm is: 2.4802"
```

```
## [1] "lasso rss is: 0 ridge rss is: 0"
```

we can see that indeed both models end up in an interpolation point. and also that the ridge norm is alot lower than the lasso norm. on one hand it makes sense because if a lot of coefficients are small then the L2 loss should be lower (if they achieve a similar solution), but this got me to think and along with the recommendation to not use glmnet i took a closer look at the coefficient of the lasso, and saw non of them are 0!

```
sum(abs(coef(lasso_fit))>0)
```

```
## [1] 100
```

i dont know for sure why we get such a solution, i could not find a proper explanation in help(glmnet), but it might be that for very small values of λ the glmnet treats it as 0 and thus yield a linear regression solution.

if i set lambda = 0.01 we get that some coefficients do become 0

```
lasso_fit = glmnet(X_small, y_small, alpha = 1, lambda = 0.01)
lasso_rss = colSums((y_small - predict(lasso_fit, s = 0.01, newx = X_small))^2)
lasso_norm = sum(abs(coef(lasso_fit)[2:100]))
sum(abs(coef(lasso_fit))>0)
```

```
## [1] 41
```

4.h

in regression we said that the model interpolate the data if it can fit each value of y correctly, in the classification scenario it is the same, we mange to classify each of the train observation to the appropriate class.

in the multi class case we classify an observation to class i if $P(y_j=i|x_j,\beta)$ is the maximum over the set $\{p(y_j=k|x_j,\beta)\}_{k=1}^K$, where K is the number of classes

```
denote \hat{y}_k(\beta) = argmax_k \{ p(y_i = k | x_i, \beta) \}_{k=1}^K
```

the corresponding minimum norm interpolation would be: $argmin\sum_{j=1}^p \beta_j^2 \ s. \ t\sum_{i=1}^n (1-I(y_i=\hat{y}_k(\beta)))=0$

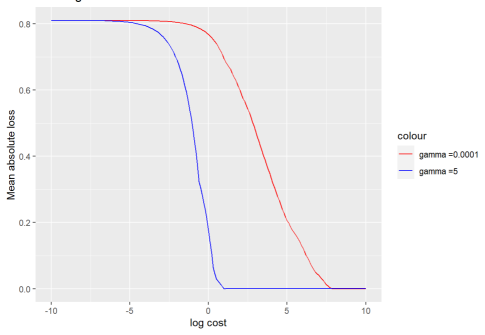
4.i

4.i.

we saw that the SVM loss is $L(y,\hat{y})=(|y-\hat{y}|-\varepsilon)_+$ so in the case that $\varepsilon=0$ the loss becomes: $(|y-\hat{y}|)_+=|y-\hat{y}|$ which is the absolute loss

```
X_{100} = as.matrix(X[1:100,])
y_100 = y[1:100,]
svm_small_gamma = svm(X_100,y_100, type="eps-regression", epsilon = 0, gamma =0.0001)
svm_large_gamma = svm(X_100,y_100, type="eps-regression", epsilon = 0, gamma =5)
cost_vals = exp(seq(-10,10,0.1))
err_small = c()
err_large = c()
for (cost in cost vals){
  svm_small_gamma = svm(X_100,y_100, type="eps-regression", epsilon = 0, gamma =0.0001, cost=cost)
  svm\_large\_gamma = svm(X\_100,y\_100, type="eps-regression", epsilon = 0, gamma = 5, cost=cost)
 err_small = c(err_small,mean(abs(y_100 - predict(svm_small_gamma,X_100))))
 err_large = c(err_large,mean(abs(y_100 - predict(svm_large_gamma,X_100))))
}
ggplot() +
 geom_line(aes(x = log(cost_vals), y = err_small, color = "gamma =0.0001")) +
  geom\_line(aes(x = log(cost\_vals), y = err\_large, color = "gamma = 5")) \ +
  scale_color_manual(values = c("gamma =0.0001" = "red", "gamma =5" = "blue")) +
  xlab("log cost") +
  ylab("Mean absolute loss") +
  ggtitle("Convergence of SVM solutions")
```

Convergence of SVM solutions



4.i.ii

for a kernel $K(x,y)=< h(x), h(y)>=\sum_{j=1}^q h_j(x)h_j(y)$ we defined $f=\sum_j \beta_j h_j$, also we know that: $\left|\left|f\right|\right|_{HK}^2=\sum_j \beta_j^2$

the kernel is a function of γ , in our case $K_{\gamma}(x,y)=exp(-||x-y||^2/(2\gamma^2))$, thus for different values of γ we have different kernels, different h functions and different β_j for each h_j . so if $\gamma\neq\gamma'$ we have $||f_{\gamma}||^2_{HK}\neq f_{\gamma'}||^2_{HK}$, and this norm is the regularization term

4.i.iii

```
fake_x = matrix(rep(5,99),1,99)
colnames(fake_x) = colnames(X_100)
svm_small_gamma = svm(X_100,y_100, type="eps-regression", epsilon = 0, gamma =0.0001, cost=exp(15))
svm_large_gamma = svm(X_100,y_100, type="eps-regression", epsilon = 0, gamma =5, cost=exp(15))
```

[1] "small gamma prediction: 3.36514 the training mean error is: 2.37040608507377e-05"

[1] "large gamma prediction: 3.58998 the training mean error is: 2.46564716309861e-05"

```
## [1] "the mean value of y is : 3.59"
```

4.i.iv

we can see that the results are rather close, but a person that rates everything 5 would probably rate the target movie 5 as well, so we can see that both models are interpolating

we see that for large γ we get a prediction very close to the mean of our target vector, this is due to the fact that for large gammas the observations are considered closer together and our K matrix becomes almost $\underline{1}^T\underline{1}$ (a matrix full of ones) so each new prediction will get a constant value of the mean of the target

we can easily demonstrate it using a fake observation with all ones:

```
fake_x = matrix(rep(1,99),1,99)
colnames(fake_x) = colnames(X_100)
svm_large_gamma = svm(X_100,y_100, type="eps-regression", epsilon = 0, gamma =5, cost=exp(15))
```

[1] "large gamma prediction: 3.58998"

```
## [1] "the mean value of y is : 3.59"
```

for small values of γ our model acts more or less like 1-NN. for very small values of γ we have $K_{\gamma}(x,y) = exp(-||x-y||^2/(2\gamma^2)) \approx exp(-\infty)I(x \neq y) + exp(0)I(x = y) = I(x = y)$ so for each observation on the train set we predict the value of the observation itself because all other observation are given a weight close to zero.

this is in the case that γ is very very small, in our case γ is small but not that close to zero so we have a larger neighborhood

lets demonstrate by taking the 5 closest neighbors of our movie lover fake person

```
dist_mat = (X_100 - matrix(rep(5,99*100),100,99))^2
norms = apply(dist_mat,1,sum)
nn_5 = order(norms)[1:5]
mean(y_100[nn_5])
```

```
## [1] 3.4
```

as we can see the result is very close to the prediction of our small γ

4.i.v

as we saw in the case where $\gamma=5$ the fit at this point (or in general at any other point) is the mean of y (our target vector).

we have a large cost, thus for minimizing the loss function we need to minimize the errors term because the penalty for large coefficient becomes neglect able.

thus for each interior point we have an interpolation, and each exterior point is a weighted sum over our observation, the weights are given by the kernel, and because of the large γ our distances increase and lose their meaning. so for a new observation we have similar weighted distance to all other observation, thus our new observation is assigned with the value of mean(y)