Computation

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The first half of our course

The second half of our course

About

- This lecture note covers how to use statistical software, R, to deal with computations in linear algebra.
- Installation and basic guides for R can be found at: https://github.com/aceMKSim/teaching/blob/master/Data%20Visualization/Lecture%20Notes/L01_installR.pdf
- More interested in R and data visualization?
 - You can access my lecture note for a graduate course, 'Data Visualization' at https://github.com/aceMKSim/teaching/tree/master/Data%20Visualization
 - You can request an invitation link to a superb online course platform, www.datacamp.com, where courses for R and Python are available free through my invitation.

The first half of our course

Matrix Input

• Matrix input is done by 1) listing all elements by c() function, 2) telling the exact number of row by nrow=, and 3) whether you want to list elements by row or by column (byrow).

```
A1 <- matrix(c(1,-3,0,2,-1,5,1,2,3),
             nrow = 3, bvrow = FALSE)
b1 \leftarrow matrix(c(0,1,-1),
             nrow=3, byrow=FALSE)
A2 <- matrix(c(4,2,2,-1,1,-1,6,6,8),
             nrow=3, byrow=FALSE)
A3 <- matrix(c(1,0,-1,-7,0,7,0,1,4,6,-2,2),
             nrow=3, byrow=FALSE)
b3 \leftarrow matrix(c(5, -3, 7),
             nrow=3, byrow=FALSE)
# Following to check for correct input
Α1
##
        [,1] [,2] [,3]
## [1,]
## [2,]
          -3 -1
                5
## [3,]
```

```
b1
##
       [,1]
## [1,]
## [2,]
## [3,] -1
Δ2
##
       [,1] [,2] [,3]
## [1,]
## [2,] 2
## [3,] 2 -1
Α3
##
       [,1] [,2] [,3] [,4]
## [1,]
          1
## [2,]
                 1 -2
## [3,]
         -1
                        2
```

```
b3
##
        [,1]
## [1,]
## [2,]
        -3
## [3,]
```

 Concatenating function cbind() allows to construct a augumented matrix.

```
Aug3 <- cbind(A3, b3)
Aug3
       [,1] [,2] [,3] [,4] [,5]
## [1,]
                         5
## [2,]
            0 1 -2 -3
## [3,]
                       2
                           7
```

-1

Reduced Echelon Form

- Package (often called as library) provides additional functionality.
- In the first ever to use a package, it must be installed by following syntax.

```
install.packages("pracma")
```

 In your computation session, you must use following syntax to activate the package.

```
library(pracma)
```

 The package pracma offers rref() that gives the reduced echelon form.

```
library(pracma)
A1

## [,1] [,2] [,3]
## [1,] 1 2 1
## [2,] -3 -1 2
## [3,] 0 5 3

rref(A1)

## [,1] [,2] [,3]
## [1,] 1 0 0
## [2,] 0 1 0
## [3,] 0 0 1
```

A2								
##		[,1]	[,2]	[,3]				
##	[1,]	4	-1	6				
##	[2,]	2	1	6				
##	[3,]	2	-1	8				
rref(A2)								
##		[,1]	[,2]	[,3]				
##	[1,]	1	0	0				
##	[2,]	0	1	0				
##	[3,]	0	0	1				
А3								
##		[,1]	[,2]	[,3]	[,4]			
##	[1,]	1	-7	0	6			
##	[2,]	0	0	1	-2			
##	[3,]	-1	7	4	2			
rre	f (A3))						
##		[,1]	[,2]	[,3]	[,4]			
##	[1,]	1	-7	0	0			
##	[2,]	0	0	1	0			
##	[3,]	0	0	0	1			

```
Aug3
       [,1] [,2] [,3] [,4] [,5]
##
## [1,]
          1
                          5
## [2,]
            0 1 -2 -3
                      2
## [3,]
       -1
            7
                         7
rref(Aug3)
       [,1] [,2] [,3] [,4] [,5]
##
          1
             -7
                       0 -4.0
## [1,]
## [2,]
                       0.0
## [3,]
                       1 1.5
         0
```

Solving Ax = b

```
Α1
       [,1] [,2] [,3]
##
## [1,]
## [2,] -3 -1 2
## [3,] 0 5
                   3
b1
##
       [,1]
## [1,]
## [2,] 1
## [3,] -1
x <- solve(A1, b1)
х
##
       [,1]
## [1,] 0.6
## [2,] -0.8
## [3,] 1.0
```

• %*% is an operator for matrix multiplication.

```
# Sanity check
A1 %*% x

## [,1]
## [1,] 0
## [2,] 1
## [3,] -1
b1

## [,1]
## [1,] 0
## [2,] 1
## [3,] -1
```

Inverting

```
Α1
       [,1] [,2] [,3]
##
## [1,]
        1
## [2,] -3 -1 2
## [3,]
            5
                   3
        0
inv A1 <- solve(A1)
inv_A1
##
       [,1] [,2] [,3]
## [1,] 1.3 0.1 -0.5
## [2,] -0.9 -0.3 0.5
## [3,] 1.5 0.5 -0.5
```

```
# Sanity check
A1 %*% inv A1
##
        [,1] [,2] [,3]
## [1,]
## [2,]
          0
                     0
## [3,]
                     1
inv A1 %*% A1
        [,1] [,2]
##
                          [,3]
## [1,]
          1
               0 0.000000e+00
## [2,]
          0
                1 2.220446e-16
## [3,]
                0 1.000000e+00
```

Rank

```
library(Matrix)
Α1
##
        [,1] [,2] [,3]
## [1,]
## [2,] -3 -1
## [3,]
                    3
rankMatrix(A1)
## [1] 3
## attr(,"method")
## [1] "tolNorm2"
## attr(,"useGrad")
## [1] FALSE
## attr(,"tol")
## [1] 6.661338e-16
```

LU decomposition

```
library(matrixcalc)
Α1
##
       [,1] [,2] [,3]
## [1,]
        1
## [2,] -3
              -1
                    2
## [3,]
          0
               5
                    3
lu.decomposition(A1)
## $L
##
       [,1] [,2] [,3]
## [1,]
## [2,] -3 1
                    0
## [3,]
          0
               1
                    1
##
## $U
##
       [,1] [,2] [,3]
  [1,]
## [2,]
## [3,]
          0
               0
                   -2
```

```
# Sanity check
A1 L <- lu.decomposition(A1)$L
A1 U <- lu.decomposition(A1)$U
A1 L %*% A1 U
##
        [,1] [,2] [,3]
   [1,]
## [2,]
        -3 -1
## [3,]
A1
##
        [,1] [,2] [,3]
## [1,]
          1
## [2,]
          -3
## [3,]
          0
                     3
```

Determinants

```
A1

## [,1] [,2] [,3]

## [1,] 1 2 1

## [2,] -3 -1 2

## [3,] 0 5 3

det(A1)

## [1] -10
```

```
## [,1] [,2] [,3]
## [1,] 4 -1 6
## [2,] 2 1 6
## [3,] 2 -1 8
det(A2)
## [1] 36
```

The second half of our course

Eigenvalue and Eigenvector

```
Α2
        [,1] [,2] [,3]
##
## [1,]
              -1
## [2,]
             1
## [3,]
          2 -1
eigen(A2)
## eigen() decomposition
## $values
## [1] 9 2 2
##
## $vectors
                      [,2]
##
             [,1]
                                 [,3]
  [1,] 0.5773503 0.2915782 0.3377822
## [2,] 0.5773503 0.8863713 0.0000000
## [3,] 0.5773503 0.0505358 -0.1125941
```

```
eigen(A2)$values
## [1] 9 2 2
eigen(A2)$vectors
##
             [,1]
                       [,2]
                                  [,3]
## [1,] 0.5773503 0.2915782 0.3377822
## [2,] 0.5773503 0.8863713 0.0000000
## [3,] 0.5773503 0.0505358 -0.1125941
```

```
lambda1 <- eigen(A2)$values[1]
lambda2 <- eigen(A2)$values[2]
lambda3 <- eigen(A2)$values[3]
v1 <- eigen(A2)$vectors[,1]
v2 <- eigen(A2)$vectors[,2]
v3 <- eigen(A2)$vectors[,3]</pre>
```

```
lambda1
## [1] 9
lambda2
## [1] 2
lambda3
## [1] 2
v1
## [1] 0.5773503 0.5773503 0.5773503
v2
  [1] 0.2915782 0.8863713 0.0505358
v3
## [1] 0.3377822 0.0000000 -0.1125941
```

Diagonalization

$$A = PDP^{-1}$$

```
P <- eigen(A2)$vectors
D <- diag(eigen(A2)$values)
Ρ
##
             [,1]
                       [,2]
                                  [,3]
## [1,] 0.5773503 0.2915782 0.3377822
## [2,] 0.5773503 0.8863713 0.0000000
## [3,] 0.5773503 0.0505358 -0.1125941
D
        [,1] [,2] [,3]
##
## [1,]
## [2,]
## [3,]
                0
                     2
```

Dataset mtcars

- The rest of this lecture note will use the one of the popular built-in datasets in R, mtcats, which includes several cars and their fuel efficiency. To analyze the dataset.
- Typing help(mtcars) in your console will provide basic description on the dataset.

help(mtcars)

The data was extracted from the 1974 Motor Trend US magazine, and comprises fuel consumption and 10 aspects of automobile design and performance for 32 automobiles (1973–74 models)

The dataset can be explored by following syntax, str().

str(mtcars) 'data.frame': 32 obs. of 11 variables: 21 21 22.8 21.4 18.7 18.1 14.3 24.4 22.8 19.2 ... \$ mpg : num \$ cyl : num 6 6 4 6 8 6 8 4 4 6 ... \$ disp: num 160 160 108 258 360 ... \$ hp : num 110 110 93 110 175 105 245 62 95 123 ... 3.9 3.9 3.85 3.08 3.15 2.76 3.21 3.69 3.92 3.92 ... ## \$ drat: num \$ wt : num 2.62 2.88 2.32 3.21 3.44 ... \$ qsec: num 16.5 17 18.6 19.4 17 ... \$ vs : num 0011010111... \$ am : num 1110000000... ## \$ gear: num 4 4 4 3 3 3 3 4 4 4 ... \$ carb: num 4 4 1 1 2 1 4 2 2 4 ...

Each varible's description is also provided from the help page as follows.

- [, 1] mpg, Miles/(US) gallon
- [, 2] cy1, Number of cylinders
- [, 3] disp, Displacement (cu.in.)
- [, 4] hp, Gross horsepower
- [, 5] drat, Rear axle ratio
- [, 6] wt, Weight (1000 lbs)
- [, 7] qsec, 1/4 mile time
- [, 8] vs, Engine (0 = V-shaped, 1 = straight)
- [, 9] am, Transmission (0 = automatic, 1 = manual)
- [,10] gear, Number of forward gears
- [,11] carb, Number of carburetors
- The first variable mpg refers to the fuel efficiency, since it measures how
 many miles the car can move with a gallon of its fuel. The other 10
 variables should affect the mpg. That is, we call mpg as a dependent
 variable and the other variables as independent variables.

Linear regression

- For simplicity, the rest of this will use only the first six independent variables from now on: cyl, disp, hp, drat, wt, qsec
- We will use

Hornet 4 Drive

Valiant

- disp as an independent variable (i.e. 'X' variable)
- mpg as dependent variable (i.e. 'Y' variable).
- Following codes will reveal a few observed pairs of (X, Y).
- i.e. the function head() reveals the first six observations of the dataset.

Hornet Sportabout 360 18.7

258 21.4

225 18.1

- Based on the textbook (Ch.6, Example 4),
 - we shall construct a A matrix where
 - the first column is one-vector (generated by rep (1, length(mtcars\$disp)))
 - the second column is the observed quantity of disp (generated by mtcars\$disp).
 - The matrix (or, a vector) b is nothing but the quantities of <code>mtcars\$mpg</code>.
- This can be done with following codes. (The function matrix() is another way to create matrix.)

```
A <- matrix(cbind(rep(1,length(mtcars$disp)), mtcars$disp), ncol = 2)
b <- matrix(mtcars$mpg, ncol = 1)
```

 To make sure A and b are correctly assigned, following codes reveal its beginning part.

```
head(A)
                                                  head(b)
        [,1] [,2]
                                                          [,1]
##
                                                  ##
## [1,]
           1 160
                                                  ## [1,] 21.0
## [2,]
           1 160
                                                  ## [2,] 21.0
## [3,]
           1 108
                                                  ## [3,] 22.8
## [4,]
           1 258
                                                  ## [4,] 21.4
## [5,]
                                                  ## [5,] 18.7
           1 360
              225
                                                  ## [6,] 18.1
## [6,]
```

• Using *A* and *b*, the least-squares solution is obtained by solving

$$A^t A x = A^t b$$

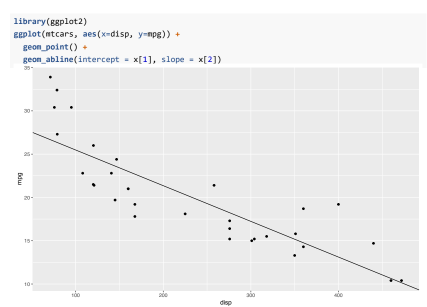
• Assign $(A^tA)^{-1}A^tb$ to the variable x

```
x <- solve(t(A) %*% A) %*% t(A) %*% b
x
## [1,] 29.59985476
## [2,] -0.04121512
x[1] # alpha
## [1] 29.59985
x[2] # beta
## [1] -0.04121512</pre>
```

- As a statistical software, R offers linear regression function called lm().
- Basic usage is given below. Confirm that the results match the numbers in the previous page.

```
lm(mtcars$mpg ~ mtcars$disp)
##
## Call:
## lm(formula = mtcars$mpg ~ mtcars$disp)
##
## Coefficients:
## (Intercept) mtcars$disp
## 29.59985 -0.04122
```

• Following is visualizion of original data and regression line.



Principal Component Analysis

Preparation

- PCA (Principal Component Analysis) is to identify common underlying components in dataset.
- In our set of six independent variables, i.e. mtcars[,c("cyl", "disp", "hp", "drat", "wt", "qsec")], the six variables are naturally highly correlated to each other.
- For example, larger engine size (disp) is likely to be positively correlated to its horse power (hp) and its weight (wt).
- Let X be the dataset of the independent variables.

```
X <- as.matrix(mtcars[,c("cyl", "disp", "hp", "drat", "wt", "qsec")])
head(X)</pre>
```

Correlation matrix

 To explore its correlation structure, the following code generates its correlation matrix.

```
## cyl disp hp drat wt qsec
## cyl 1.0000000 0.9020329 0.8324475 -0.69993811 0.7824958 -0.59124207
## disp 0.9020329 1.0000000 0.7909486 -0.71021393 0.8879799 -0.43369788
## hp 0.8324475 0.7909486 1.000000 -0.44875912 0.6587479 -0.70822339
## drat -0.6999381 -0.7102139 -0.4487591 1.0000000 -0.7124406 0.09120476
## wt 0.7824958 0.8879799 0.6587479 -0.71244065 1.0000000 -0.17471588
## qsec -0.5912421 -0.4336979 -0.7082234 0.09120476 -0.1747159 1.0000000
```

Assumption of PCA

- PCA assumes that, each observation can be tersely expressed without using all of the six variables.
- For example, since cyl and disp are highly correlated, and one can find a nearly linear combination between the two.
- Or, it would mean that each observation may be characterized by a single variable, which is a linear combination of cyl and disp.

Dimension reduction

- Since the original two variables, cyl and disp, can be used to generate a single variable through their linear combination without necessarily hurting represent-ability, now the originally two variables are reduced to a single variable.
- Thus, PCA is one of the dimension reduction techniques, and maybe useful especially if
 - i) variables are highly correlated to each other and/or
 - ii) there are too many variables to consider.

Preparing covariance matrix

- Remind that, from your elementary statistics course, one variable's sample variance calculated as $\frac{\sum_{i=1}^{n}(X_{i}-\overline{X})^{2}}{n-1}$.
- In order words, from original observations, i) subtract its sample mean (a.k.a. *de-mean* process), then ii) take squared value, and then divide it by n-1.
- Preparing sample covariance matrix is analgous to that as follows.
- Please attemp to understand the following code.

```
# de-meaning process of $X$
# (for each column, x-mean(x) is applied)
X_demeaned <- apply(X, 2, function(x) x-mean(x))</pre>
```

• Remind that, for a matrix A, t(X)%*%X is equivalent to squaring it.

```
Cov <- (t(X_demeaned) %*% X_demeaned)/(nrow(X)-1)
```

 \bullet Covariance matrix Cov is a 6×6 matrix, because six variables are involved here.

Cov	,							
##		cyl	disp	hp	drat	wt	qsec	
##	cyl	3.1895161	199.66028	101.93145	-0.66836694	1.3673710	-1.88685484	
##	disp	199.6602823	15360.79983	6721.15867	-47.06401915	107.6842040	-96.05168145	
##	hp	101.9314516	6721.15867	4700.86694	-16.45110887	44.1926613	-86.77008065	
##	drat	-0.6683669	-47.06402	-16.45111	0.28588135	-0.3727207	0.08714073	
##	wt	1.3673710	107.68420	44.19266	-0.37272073	0.9573790	-0.30548161	
##	qsec	-1.8868548	-96.05168	-86.77008	0.08714073	-0.3054816	3.19316613	

Positive semi-definite (psd)

- A proper covariance matrix is positive semi-definite, meaning that its eigenvalues are all non-negative. (Just like your familiar variance is always non-negative)
- Find its eigenvalues and see if they are all non-negative numbers.

```
eigen(Cov)$values
```

```
## [1] 1.861324e+04 1.453791e+03 1.586537e+00 4.584231e-01 1.248216e-01 ## [6] 8.738486e-02
```

• All looks good, so we can proceed.

Principal components

• Remind that the six pricinpal components correspond to $v_1,v_2,...,v_6$. It can be retrieved as following.

```
eigen(Cov)$vectors
##
                [,1]
                            [,2]
                                         [,3]
                                                      [,4]
                                                                   [,5]
## [1.] -0.012042822 -0.003376004 0.225464493 0.913153920 0.116301042
## [2.] -0.900263018  0.435130058  0.004337984 -0.012130860  0.004694063
## [3,] -0.435075671 -0.899991800 -0.026532118 -0.002512279 0.003469759
## [4,] 0.002661502 -0.003901248 0.057414058 -0.323952312 -0.131060362
## [5,] -0.006242691
                     0.004868276 -0.203604150 0.191625494 -0.958484375
## [6,] 0.006676402
                     0.025025499 -0.950627144 0.155984033 0.224879700
               [,6]
##
## [1,] 0.318799814
## [2,]
        0.000566658
## [3,] -0.001271341
## [4,] 0.935178808
## [5,] -0.055408055
## [6,] 0.143997516
```

Explanation power of each principal component.

- Let the eigenvalues of the matrix Cov be $\lambda_1,\lambda_2,...,\lambda_6$ in a descending order.
- The amount of overall variation that the first principal component can explain is $\frac{\lambda_1}{\sum_{i=1}^{6} \lambda_i}$. (Check this number is about 92.7%)
- The amount of overall variation that the second principal component can explain is $\frac{\lambda_2}{\sum_{i=1}^6 \lambda_i}$, and so on.
- In sum, the following is the explanation power of each principal components.

```
exp_power <- eigen(Cov)$values/sum(eigen(Cov)$values)
exp_power
## [1] 9.274490e-01 7.243857e-02 7.905297e-05 2.284202e-05 6.219529e-06
## [6] 4.354157e-06
```

ullet Cumulatively, the first n principal components' explanation power is:

```
cumsum(exp_power) # cumulative sum
## [1] 0.9274490 0.9998875 0.9999666 0.9999894 0.9999956 1.0000000
```

"Optimism is the faith that leads to achievement - Hellen Keller"