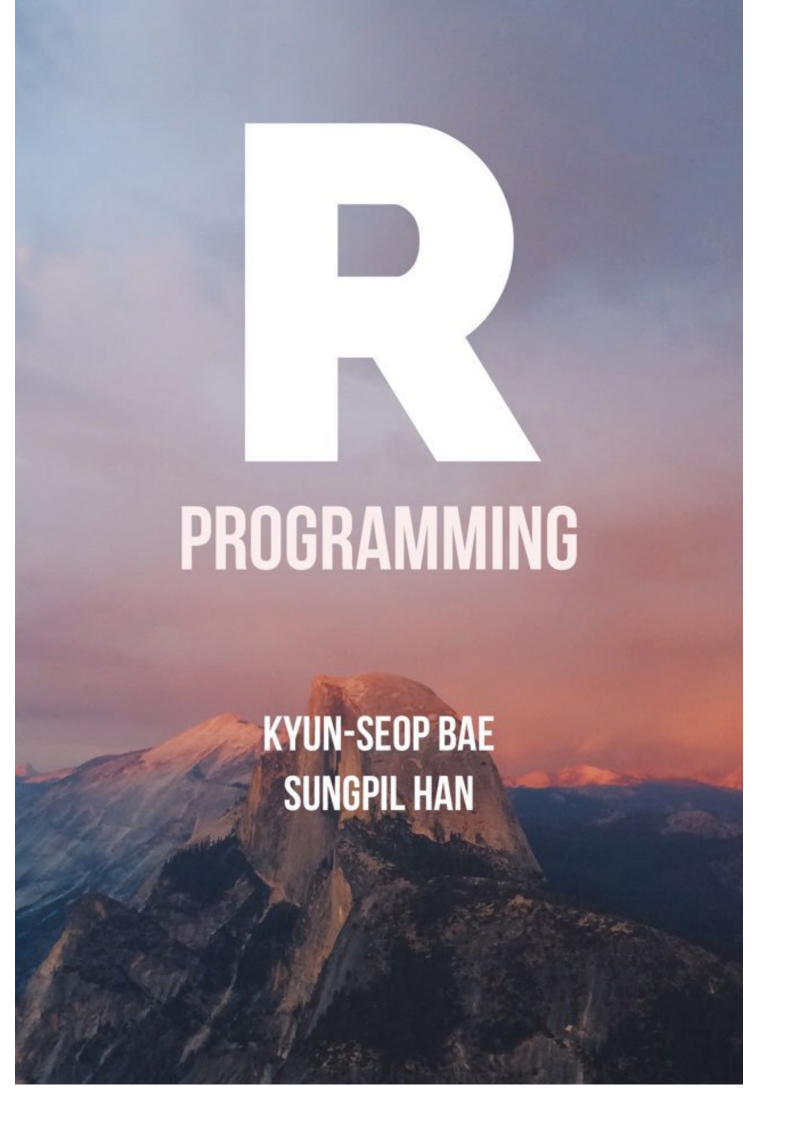
R Programming - Lecture Notes



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Preface

안녕하십니까?

2017년 1학기 울산대학교 의학과 대학원 수업 R Programming 과목 담당교수 배균섭입니다.

R은 http://cran.r-project.org 에서 다운로드받아 설치할 수 있습니다. 역시 같은 사이트에서 Manual이 나와 있으니 참고하시기 바랍니다. 구글에서 'R Programming pdf'와 같은 키워드로 검색하시면 많은 자료를 보실 수 있습니다.

첨부한 R.stx¹ 파일은 AcroEdit이라는 editor에서 사용할 syntax highlighting 용 구문 파일입니다. http://www.acrosoft.pe.kr 에서 다운로드 받아 설치하시기 바랍니다. AcroEdit 대신 notepad++를 선호하시는 분은 그대로 사용하셔도 됩니다.

저는 RStudio, tinnR 등을 이용해서 강의하지 않습니다만, 필요하신 분은 쓰셔도 괜찮습니다. 향후 R package 작성을 위해서는 MiKTeX와 Rtools를 설치하십시오.

추가로 말씀드리자면, http://www.coursera.org 에 많은 R 강좌가 개설되어 있습니다. Specialization course로 들어가면 유로이지만, (Specialization course는 여러 개의 과목이 합쳐져 있는 것입니다.) 개별 과목을 검색해서 들어가면, 무료로도 볼 수 있습니다. (대신시험을 칠 수 없거나, certificate를 받을 수 없습니다.)

좋은 강좌가 많으니 많이 활용하시기 바립니다.

강의 장소에 불편함이 많은 것으로 생각되어, 다음과 같이 Skype 모임을 개설하였습니다. 사정상 원거리에서 오시기 불편한 분들은 활용하시기 바랍니다. 출석은 화면을 캡쳐하거나 휴대폰으로 찍은 뒤 sec@acp.kr², shan@acp.kr□³ 보내주시면 출석으로 인정해드립니다.

Skype 모임 참가 https://meet.lync.com/uucp-acp/ksbae/SKGJ3BNQ

2017년 3월, 배균섭 배상

The online version of this book is licensed under the Creative Commons Attribution-NonCommercial-ShareAlike 4.0 International License⁴.

¹https://groups.google.com/a/acr.kr/group/r/attach/409db97bf453a/R.stx?part=0.1&authuser=0

²mailto:sec@acp.kr

³mailto:shan@acp.kr⊠

⁴http://creativecommons.org/licenses/by-nc-sa/4.0/

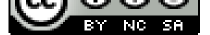


FIGURE 1: Creative Commons License

Teaching Assistant

안녕하십니까? 서울아산병원 임상약리학과 전공의 한성필입니다. 수업과 관련된 여러 제반 업무를 담당하고 있습니다. 언제든 의문사항 있으면 r@acr.kr⁵ 로 전체 메일 보내 시거나 교수님 k@acr.kr⁶ 혹은 제 개인 메일 shan@acp.kr⁰ 연락해 주십시오.

교수님께서 세우신 방침에 따라 수업시간에 출석을 부르지 않을 예정입니다. 수강하시는 화면(Skype)을 휴대폰으로 사진 찍으시거나 강의실의 스크린을 사진으로 촬영하셔서 sec@acp.kr⁸ / shan@acp.kr⁹ 로 동시에 보내주시면 됩니다. 가급적 "2017-03-31 한성필 출석" 과 같은 식의 제목을 유지해 주시면 처리하는데 큰 도움이 될 것 같습니다.

출석 체크를 위해 전체메일을 사용하지 말아주십시오!

아울러 수업 중에 사용한 코드/스크립트를 사용하여 R의 패키지인 bookdown을 사용해 웹북을 제작 중에 있습니다. (Xie, 2016) 여러분이 읽고 있는 이 책 자체가 R 코드의 일종인 Rmarkdown의 결과물이라고 보시면 됩니다. Github 저장소¹⁰가 있으니 소스 코드를 보실수 있습니다. 누구나 소스를 편집하여 Pull Request를 요청할 수 있으므로 혹시 Github를 사용하셔서 웹북의 질을 높이고자 하시는 수강생 선생님들께서는 도움을 주십시오.

감사합니다.

2017년 3월, 한성필 올림

FAQ

접속 관련

⁵mailto:r@acr.kr

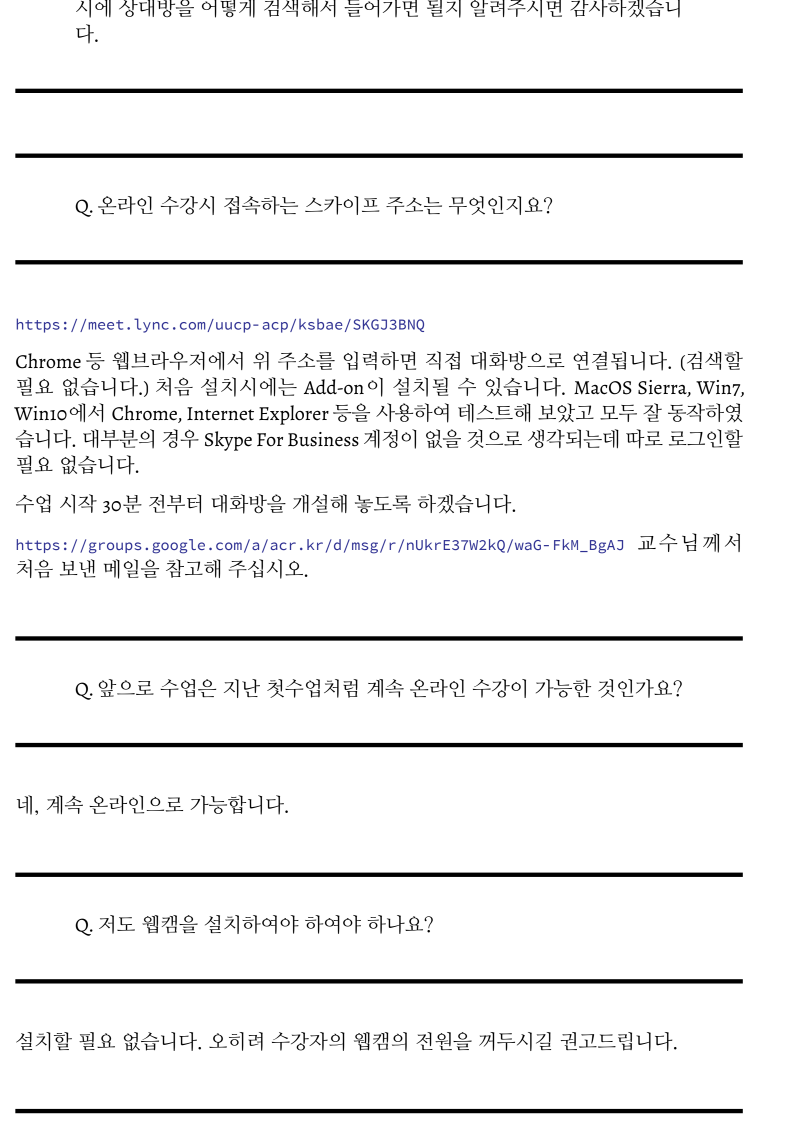
⁶mailto:k@acr.kr

⁷mailto:shan@acp.kr⊠

⁸mailto:sec@acp.kr

⁹mailto:shan@acp.kr

¹⁰https://github.com/asancpt/Rprogramming



수업 시작 30분 전부터 대화방을 개설하여 놓도록 하겠습니다.

출석관련

Q. 미국학회 참석으로 수업시간이 귀국행 비행기 기내에 있을거같아 출석이 안될것 같습니다. 방법이 있을지요?

결석 사유서를 제출해 주시면 출석 처리 하겠습니다. 대학원 홈페이지 참고 바랍니다."이 링크로 들어가시면 가장 위에 있습니다. (결석사유서.hwp) 참고로 수업 영상은 녹화하여 Youtube에 비공개 링크를 만들 예정이라서 추후에 관련 영상을 시청할 수 있을 것 같습니다. 결석사유서를 제출한다고 100% 출석이 인정되는 것은 아닙니다. 이것이 기본적으로는 offline 강의이기 때문에 강의시간에 강의실에 있든지, 또는 온라인으로 접속해 있어야 합니다. 출석사유서를 제출하거나, 추후 동영상 시청을 해서 그 증거(사진)을 제출하는 경우에 감점을 줄여드릴 수 있습니다. 예를 들어, 결석시에는 2점 감점인데, 결석사유서를 제출하면 1점만 감점한다는지, 동영상을 보면 0.5 점만 감점한다는지 하는 것입니다. 결석 사유서 제출 시 출석 처리 원칙에 대한 설명을 드리오니, 참고하시길 바랍니다.

과제 관련

Q. 과제물이 있다고 들었는데 언제 assign하게 되는지요?

과제물은 빨라야 5주차 이후에 나갑니다.

^{**}Inttp://www.medulsan.ac.kr/graduate/?mid=72&curpage=files

Q. 첫 수업 때, certification 관련 말씀을 하셨는데, 정확히 coursera 사이트에서 어떤 것을 듣고, 제출을 해야하는지 궁금합니다. (비슷한 내용이많아, 어떤것을 들어야하는지 헷갈립니다.)

Coursera는 꼭 어느 것을 들어야 하는 것은 아니고, R programming과 관련된 것이라면 자유로이 골라서 들으면 됩니다. 대표적인 두 가지만 들자면 다음과 같습니다.

- https://www.coursera.org/learn/r-programming
- https://www.coursera.org/learn/r-programming-environment

Q. Coursera 강의를 듣고 증명서를 내면 출석을 얼마나 커버할 수 있을런 지요?

Coursera는 출석 커버보다는 grade를 올려 주기 위한 것입니다. 출석은 Skype로 커버해야 합니다. 출석의 성적 반영비율은 25%이지만, 규정상 4회 이상 결석이면 성적이 나갈 수 없습니다.

Syllabus

2017년 4월 10일 개정된 수업계획서입니다.

2017-1학기 수업계획서(Course Outline)

년도-학기 (year-semester)	2017-1	과= (course		R 프로그래밍 R Programming	g		
과목번호-분반 (courseNo-classNo	WA5493 - 01	학점(강의 (cre		3학점(3-0))	ABEEK(설계학점) (Abeek credit)	
담당교수 (professor)	배균섭 Bae, Kyun Seop		! 번호 phone)	02-3010-46	11	학부(과)사무실 (Dept. office phone	02-3010-4217
개설학과-학년 (department-year)	의학과					이수구분 (type of course requirement)	공통 common
E-MAIL	ksbae@amc.seoul.kr (t			좌구분 f lecture)		일반2 general l	t좌 ecture
홈페이지 (Homepage)				법(method of valuation)		절대평가 absolute ev	/ 등급 aluation

1.교과목 개요(course description)

Data Science의 가장 기본적인 tool인 R로 어떻게 프로그래밍을 하는지에 기술을 익힌다. 자신의 세부 전공분야에 상관없이 적용할 수 있는 공통적인 부분을 중심으로 학습할 것이다. Students will learn how to program using R which is the basic tool for the data science. The subjects will be common ones regardless of their specific majors.

2.교수목표(goal of instruction)

- 1.R을 이용하여 자신의 문제를 해결할 수 있다.
- 2.R package를 개발하여 다른 사람의 문제 해결을 돕는다.
- 3.정답이 알려져 있지 않는 문제에 대하여, 체계적인 시행착오로 최적의 해답을 찾는 습관을 들인다.
- 4.자신이 가진 Tool들의 한계와 장단점을 이해하고 자신의 문제에 적용한다.

3.주요 학습내용 및 수업진행방법(main contents & methods of teaching)

강의, 동영상 시청

4.학습 성과 평가방법(evaluation criteria)

과제 (중간, 기말고사 대신 과제를 제출해야 하며, 중간, 기말고사 기간에도 강의가 있습니다.)

	평가항목(eva luation	출석 (attendance)	중간고사(mid term exam)	기말고사 (final exam)	· ·	발표(present ation)	퀴즈 (quiz)	Term Project	기타 (etc.)
t	성적반영 비율(percentage)	25.00	25.00	25.00	25.00	0.00	0.00	0.00	0.00

5.교재 및 참고 문헌(textbook & reference books)

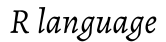
- 1. [부교재] [도서관] (데이터 고급 분석과 통계 프로그래밍을 위한) 빅데이터 분석 도구 R 프로그래밍 노만 매트로프 지수 옮김 의왕 : 에이콘 9788960773332; 9788960772793(set)
- 2. [부교재] [도서관] R Cookbook : 데이터 분석과 통계, 그래픽스를 위한 실전 예제 폴 티터 지음:이제원 옮김 서울: 인시 9788966260379
- 3. [부교재] [수기입력] Software for Data Analysis Chambers JM Springer
- 4. [부교재] [수기입력] Advanced R Wickham H CRC Press
- 5. [부교제] [도서관] [ebook]The Basics of S-PLUS [electronic resource] Krause A, Olson M New York, NY : S₁ New York 9780387283906
- 6. [부교제] [도서관] [ebook]Introduction to Scientific Programming and Simulation Using R, Second Edition [electronic resource] Jones O, Maillardet R. Hoboken: CRC Press 9781466570016
- 7. [부교재] [수기입력] R Programming for Data Science Peng R. Iulu.com
- 8. [주교재] [수기입력] The R Manuals R Core Team http://cran.r-project.org

6.주별 진도계획 및 학습자료(weekly plan & study materials)							
제1주(week 1)	[주별진도(topic)] Course introduction: textbooks, resources, installation, …	[학습자료(materials)] R-admin					
제2주(week 2)	[주별진도(topic)] Data objects: basic types, vector, matrix, list, data frame	[학습자료(materials)] R-intro					
제3주(week 3)	[주별진도(topic)] Plotting and graphics	[학습자료(materials)] Handout					
제4주(week 4)	[주별진도(topic)] Data manipulation: loading, subsetting, merging, saving	[학습자료(materials)] R-data					

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제5주(week 5)	[주별진도(topic)] Basic commands and functions	[학습자료(materials)] R-lang
제6주(week 6)	[주별진도(topic)] Elements of programming style	[학습자료(materials)]
제7주(week 7)	[주별진도(topic)] Developing R package without Rstudio	[학습자료(materials)] R-exts
제8주(week 8)	[주별진도(topic)] Some useful packages 1 - RODBC, rtf	[학습자료(materials)]
제9주(week 9)	[주별진도(topic)] (석가탄신일 휴무)	[학습자료(materials)]
제10주(week 10)	[주별진도(topic)] Rstudio and some useful packages 2 - ggplot2	[학습자료(materials)]
제11주(week 11)	[주별진도(topic)] Some useful packages 3 - dplyr, tidyr	[학습자료(materials)]
제12주(week 12)	[주별진도(topic)] Some useful packages 4 - lubridate, stringr	[학습자료(materials)]
제13주(week 13)	[주별진도(topic)] Handling date, time, and string	[학습자료(materials)]
	[주별진도(topic)] Functional and object-oriented programming in R	[학습자료(materials)]
	[주별진도(topic)] 과제 solution 예시 & feedback	[학습자료(materials)]
제16주(week 16)	[주별진도(topic)] Pitfalls and limitations of R	[학습자료(materials)]

울산대학교 *U*WIN 2/2



2017-03-15 배균섭 교수님 강의

R Language Definition¹의 초반 내용에 대해 설명하였습니다.

Graphics

2017-03-22 임형석 교수님 강의

R을 사용해 그림 그리는 방법에 대해 알아보겠습니다.

2.1 Introduction

- 상위수준 그림 함수는 그림을 생성한다.
- 하위수준 그림 함수는 기존의 그림에 그림을 추가한다.

2.2 상위수준 그림 함수

2.2.1 상위수준 그림 함수의 주요 인자 (arguments)

• main : 제목

• xlab/ylab:x축 및 y축 레이블

• xlim/ylim:x축 및 y축 범위

• col: 색깔

• lty:선 모양

• pch : 점 모양

• cex: 그림 성분의 크기

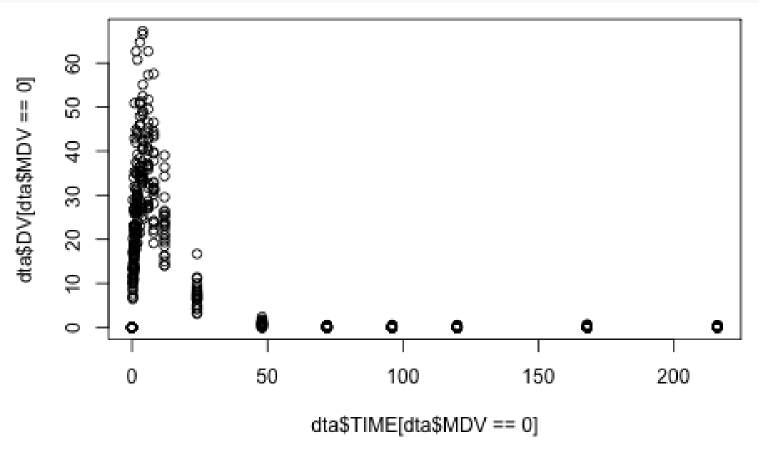
• lwd : 선 굵기

• tvpe : 그림 타입

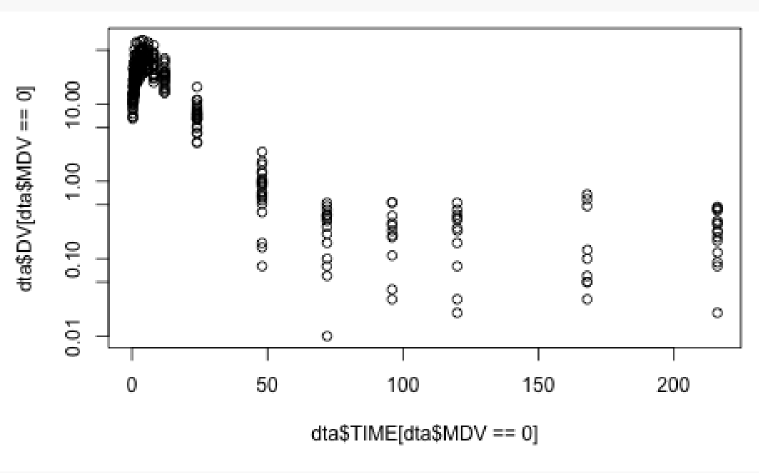
```
head(dta)
##
     ID TIME AMT
                     DV MDV
      1 0.00
                   0.00
##
               0
                          0
## 2
      1 0.00
               4 0.00
                          1
      1 0.33
## 3
                  9.40
                          0
               0
## 4
      1 0.66
               0 13.71
                          0
## 5
      1 1.00
               0 16.52
## 6
      1 1.50
               0 29.36
                          0
str(dta)
## 'data.frame':
                    456 obs. of 5 variables:
    $ ID : num
##
                  1 1 1 1 1 1 1 1 1 1 ...
    $ TIME: num
                  0 0 0.33 0.66 1 1.5 2 3 4 6 ...
##
    $ AMT : num
##
                  0 4 0 0 0 0 0 0 0 0 ...
    $ DV
                  0 0 9.4 13.7 16.5 ...
##
         : num
##
    $ MDV : num
                  0 1 0 0 0 0 0 0 0 0 ...
```

2.2.2 scatter plot

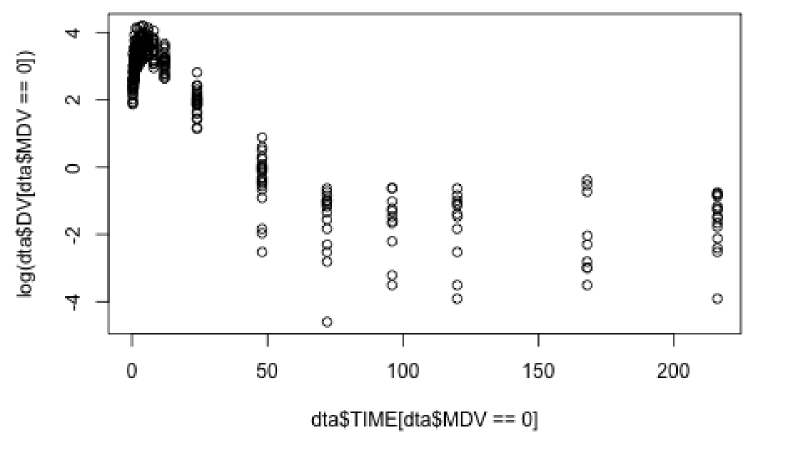
```
plot(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0])
```



```
## values <= 0 omitted from logarithmic plot
```



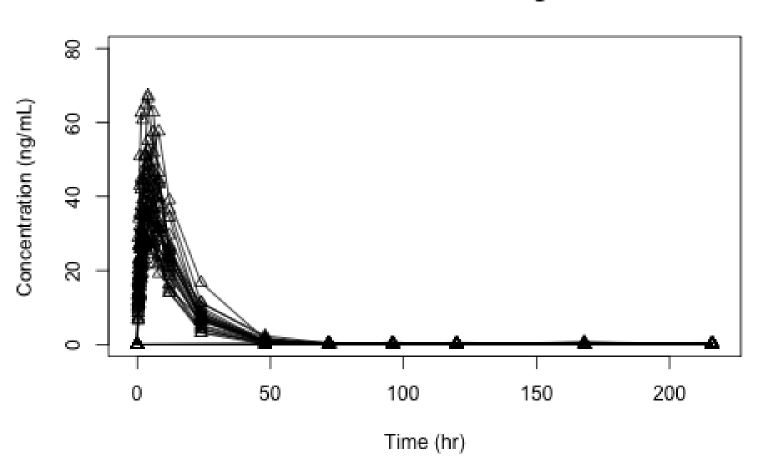
```
plot(dta$TIME[dta$MDV==0], log(dta$DV[dta$MDV==0]))
```

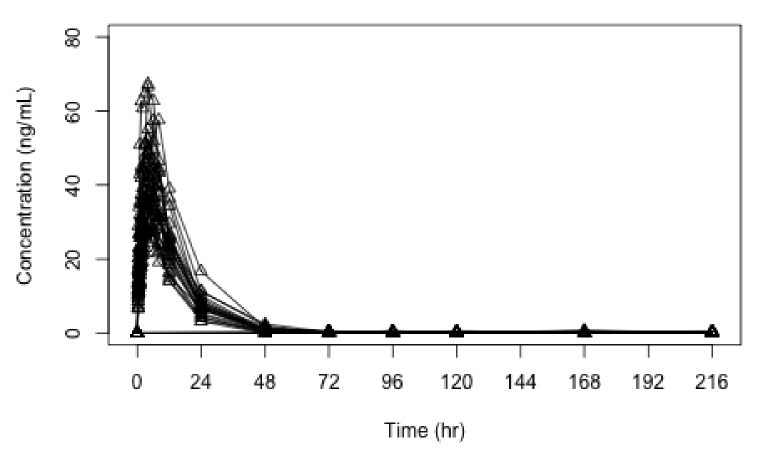


```
plot(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0]
, xlab="Time (hr)", ylab="Concentration (ng/mL)"
```

, xlim =c(-2,218), ylim=c(0,80))

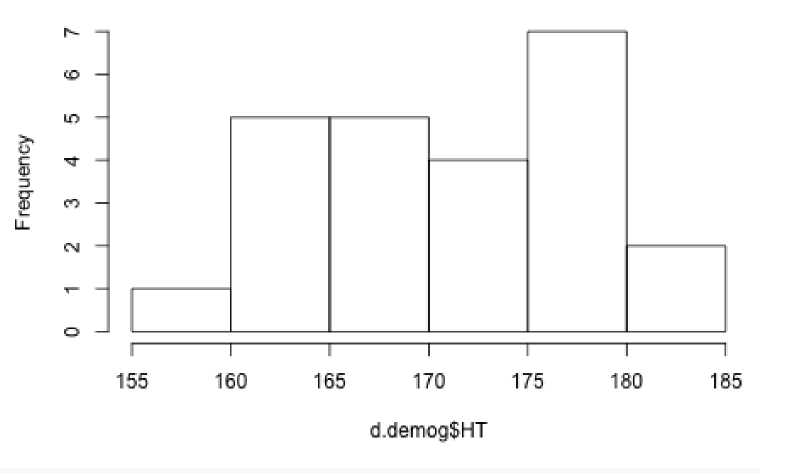
PK time-course of Drug X





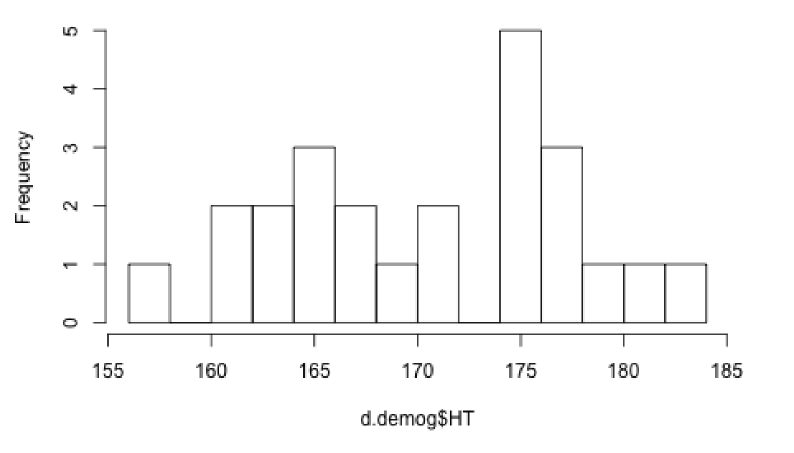
2.2.3 Histogram

```
d.demog <- read.csv("DEMOG.csv")
hist(d.demog$HT)</pre>
```



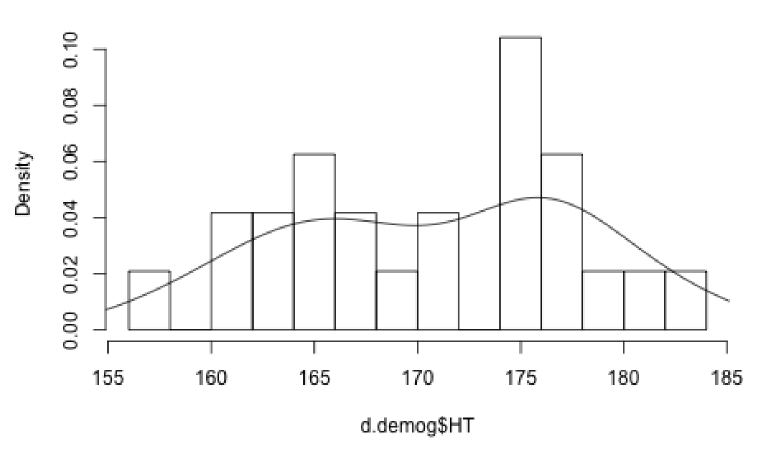
```
hist(d.demog$HT, breaks=10)
hist(d.demog$HT, nclass=10)
```

Histogram of d.demog\$HT

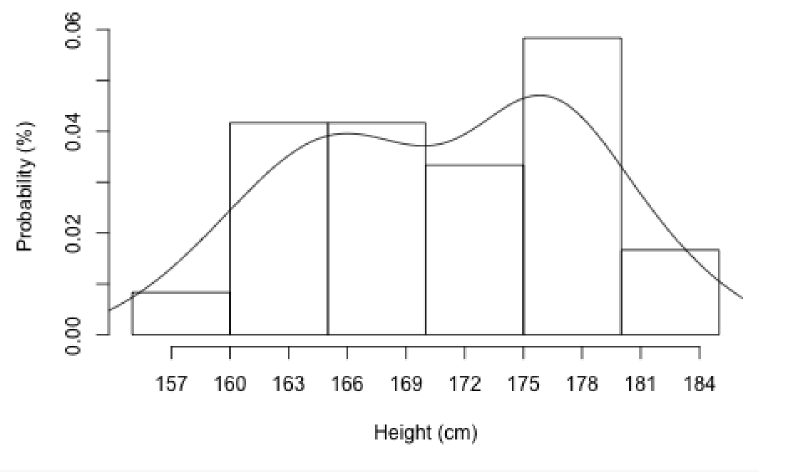


```
hist (d.demog$HT, probability=TRUE, breaks=10)
lines(density(d.demog$HT))
```

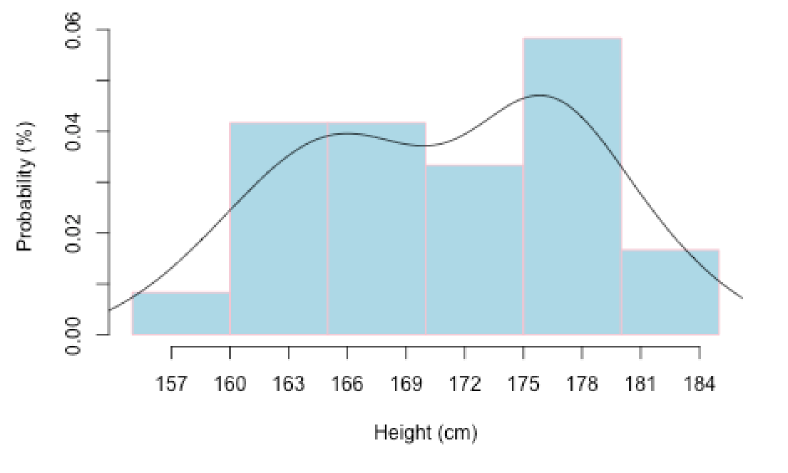
Histogram of d.demog\$HT



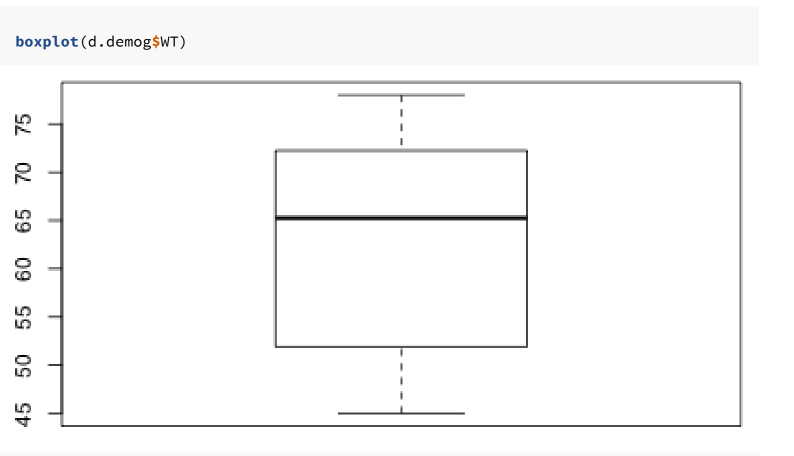
```
hist (d.demog$HT, probability=TRUE, breaks=9, xaxt="n"
    , main="Histogram for Height", xlab="Height (cm)", ylab="Probability (%)")
axis(1, at=seq(min(d.demog$HT), max(d.demog$HT), 3))
lines(density(d.demog$HT))
```



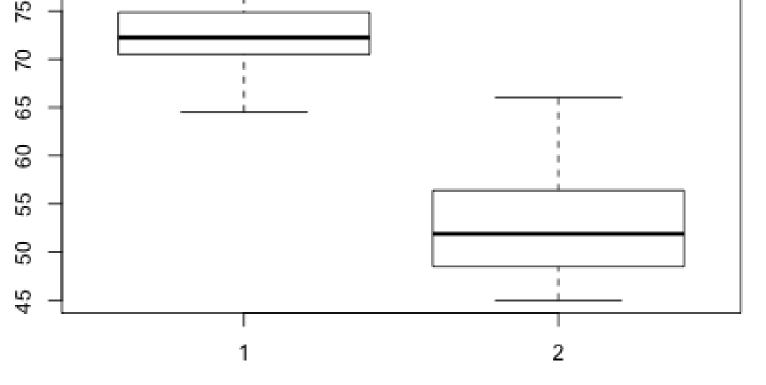
```
hist (d.demog$HT, probability=TRUE, breaks=9, xaxt="n"
    , main="Histogram for Height", xlab="Height (cm)", ylab="Probability (%)"
    , col = "lightblue", border = "pink")
axis(1, at=seq(min(d.demog$HT), max(d.demog$HT), 3))
lines(density(d.demog$HT))
```

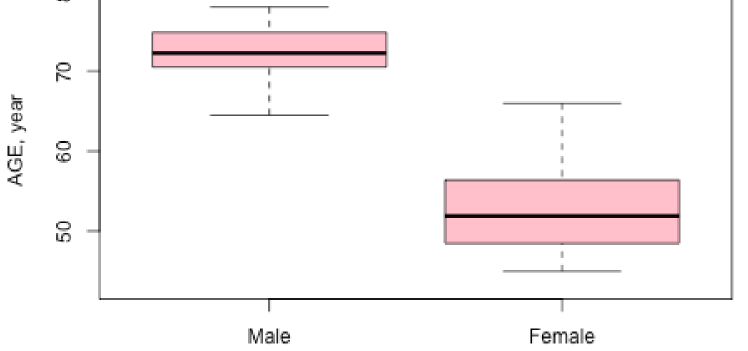


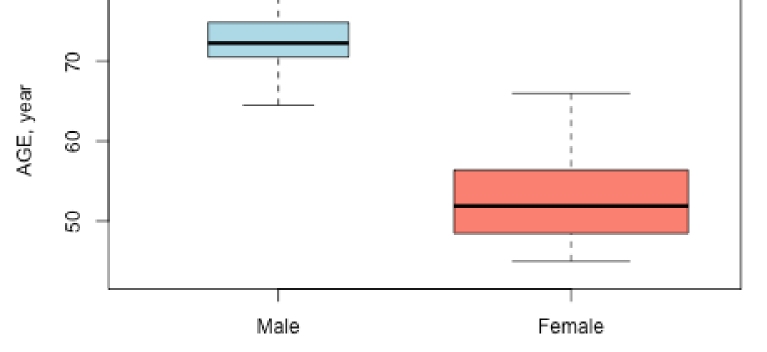
2.2.4 Box-Whisker Plot



```
boxplot(d.demog$WT ~ d.demog$SEX)
boxplot(split(d.demog$WT, d.demog$SEX))
```







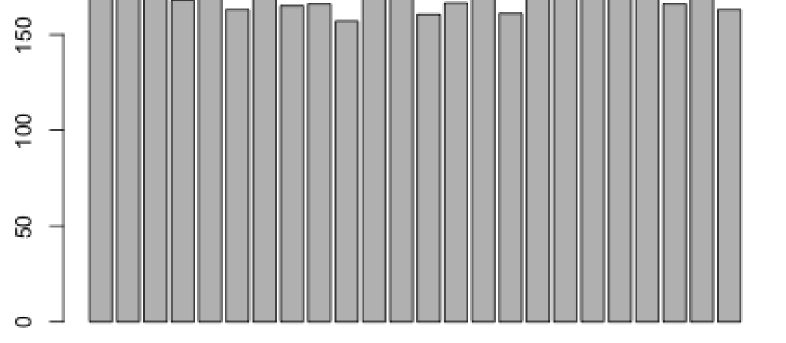
-varwidth: if varwidth is TRUE, the boxes are drawn with widths proportional to the square-roots of the number of observations in the groups.

```
boxplot(d.demog$WT ~ d.demog$SEX
, names=c("Male","Female"), ylab="AGE, year", ylim=c(min(d.demog$WT)-2, max(d., col=c("lightblue", "salmon")
, varwidth=TRUE)

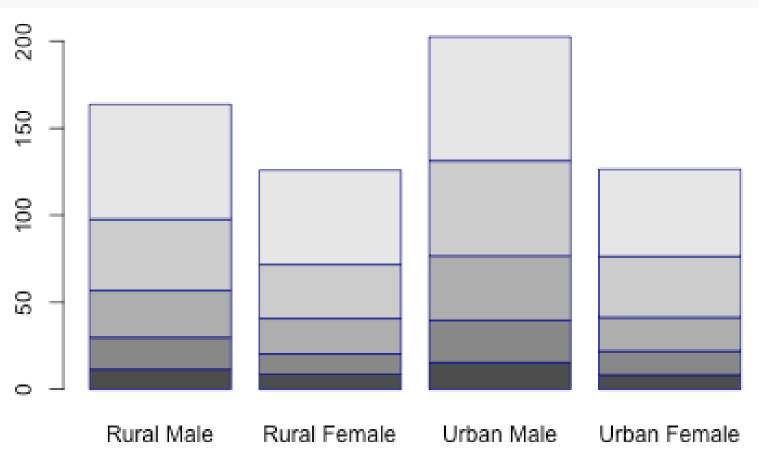
Male

Female
```

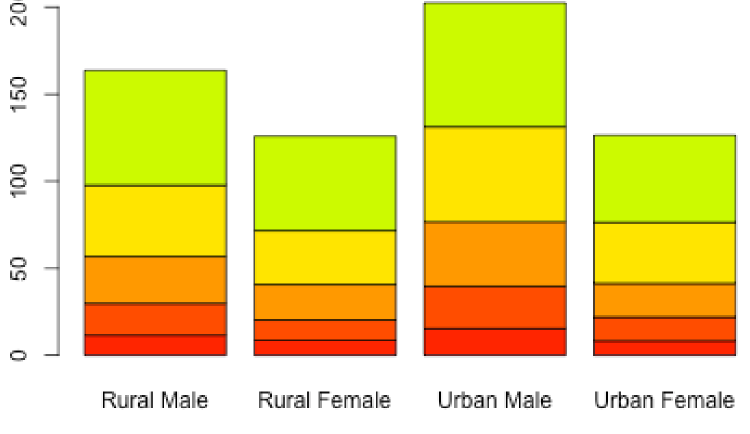
2.2.5 Bar Plot



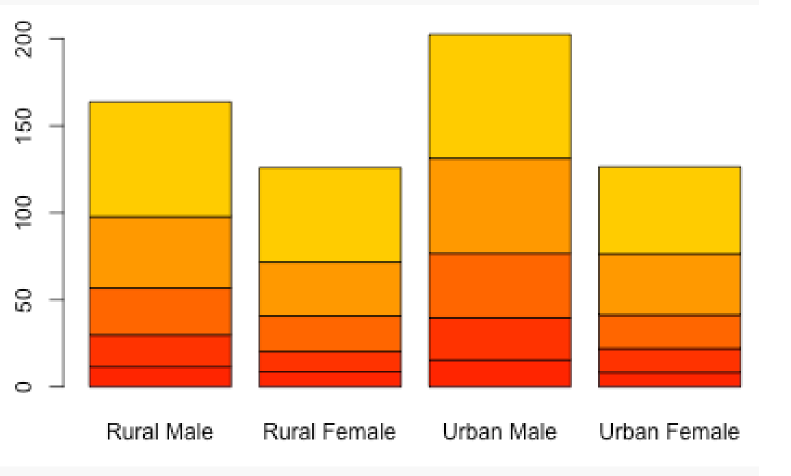
VADeaths ## Rural Male Rural Female Urban Male Urban Female 8.7 15.4 8.4 ## 50-54 11.7 ## 55-59 18.1 11.7 24.3 13.6 ## 60-64 26.9 20.3 37.0 19.3 ## 65-69 41.0 30.9 54.6 35.1 ## 70-74 66.0 54.3 71.1 50.0 barplot(VADeaths, border = "dark blue")



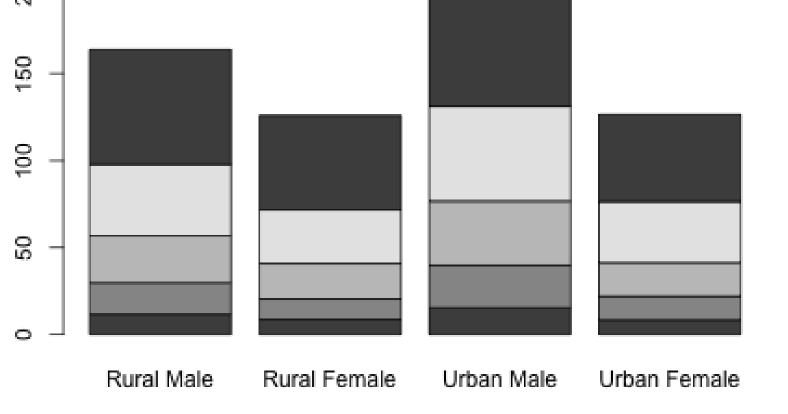




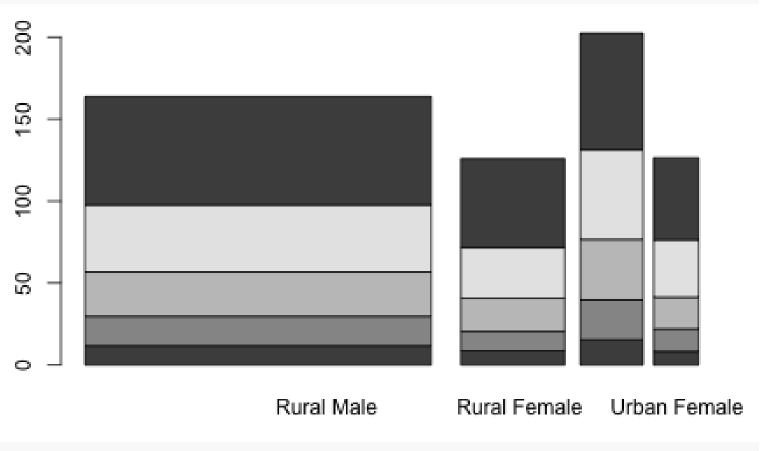




barplot(VADeaths, col = gray.colors(4))



barplot(VADeaths, col = gray.colors(4), log="x")



barplot(VADeaths, col = gray.colors(4), log="y")

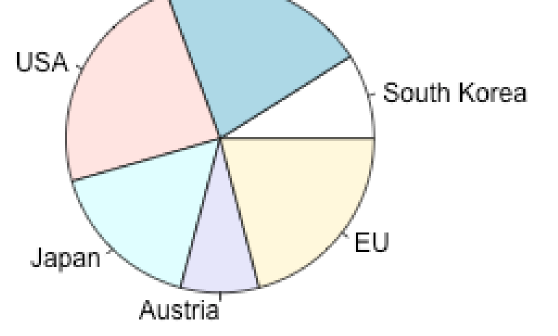


barplot(VADeaths, col = gray.colors(4), log="xy")



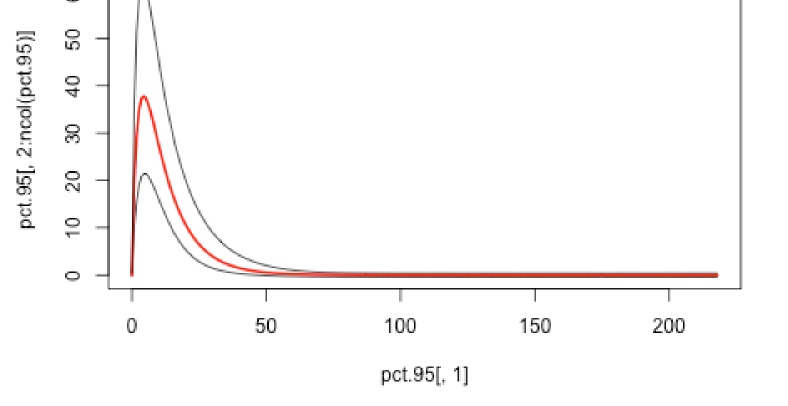
2.2.6 pie chart

```
drug.X.market <- c(0.12, 0.29, 0.32, 0.22, 0.11, 0.28)
names(drug.X.market) <- c("South Korea", "China", "USA", "Japan", "Austria", "EU")
pie(drug.X.market)</pre>
```



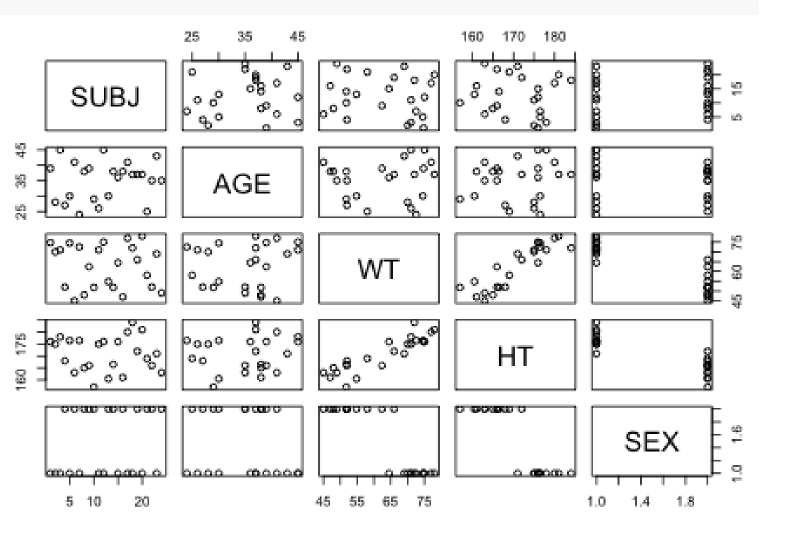
2.2.7 matplot 함수

2.2.7.1 matrix와 column 사이의 그림



2.2.8 Scatter plot matrices (pairs plots)

pairs(d.demog)



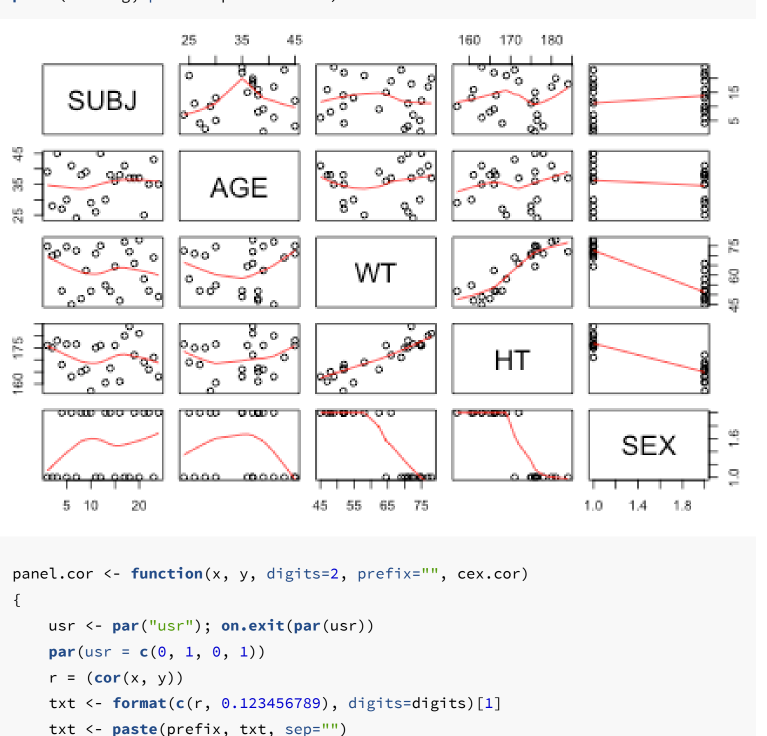
```
pairs(d.demog, panel = panel.smooth)
```

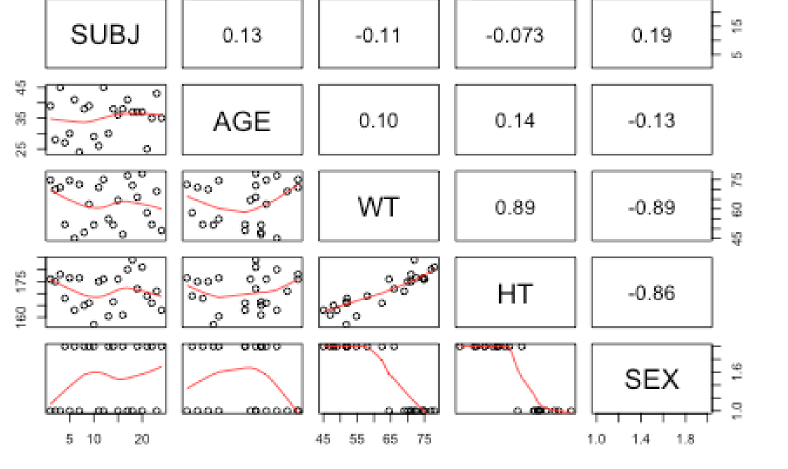
if(missing(cex.cor)) cex <- 1.5</pre>

text(0.5, 0.5, txt, cex = 1.5)

pairs(d.demog, lower.panel=panel.smooth, upper.panel=panel.cor)

}





2.3 하위수준 그림 함수

• points : 점추가

• lines : 선 추가

• abline : 기준선 추가

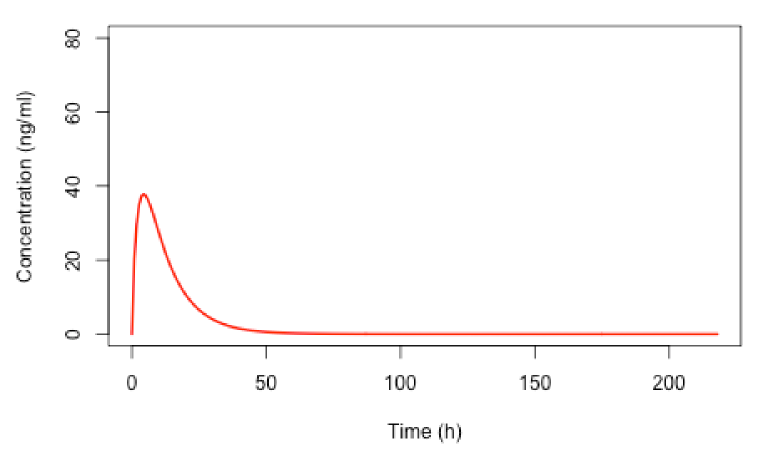
• mtext : 텍스트 추가

• legend : 설명 (legend) 추가

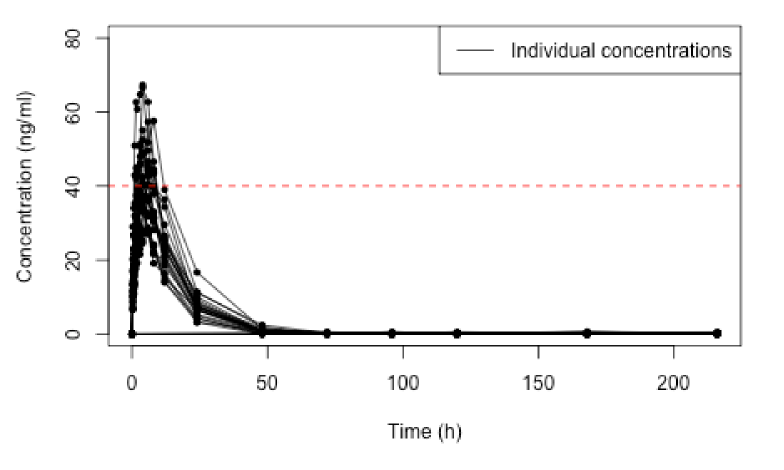
• polygon : polygon 추가

2.3.1 점, 선, 설명 추가 하기 {add}

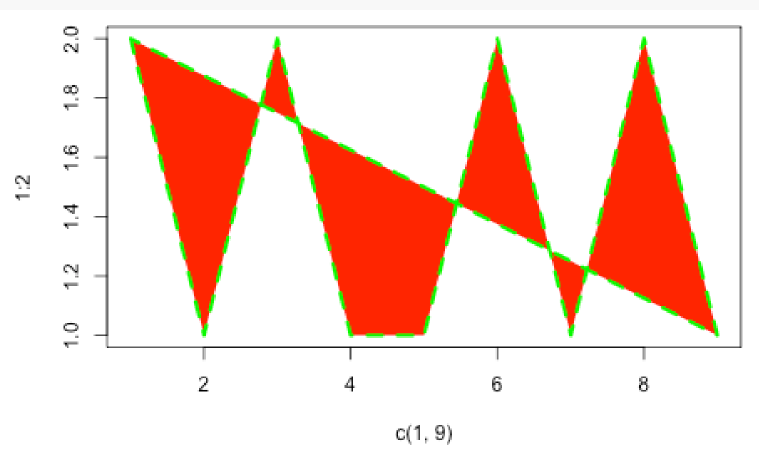
```
plot(pct.95$TIME, pct.95$PCT50, main="PK of Drug X"
    , type="l", xlab="Time (h)", ylab="Concentration (ng/ml)"
    , ylim=range(0,80), lty=1, col="red", lwd=2)
```



```
plot(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], main="PK of Drug X"
          , type="n", xlab="Time (h)", ylab="Concentration (ng/ml)"
          , ylim=range(0,80))
points(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], pch = 16, cex=0.8)
lines(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], col="black", lwd=1)
abline(40, 0, col="red", lty=2)  # abline(a,b): y=a+b*x
legend("topright", legend=c("Individual concentrations")
          , lty=1, col="black")
```



2.3.2 polygon 함수



2.4 그림 출력하기

2.4.1 pdf graphics devices

```
dev.off()
## quartz_off_screen
## 2
```

2.4.2 PNG graphics devices

Data Import / Export

2017-03-29 배균섭 교수님 강의

이번 시간에는 자료를 불러오고 조작을 가한 뒤 저장하는 방법에 대해 알아보겠습니다.

3.1 Read.csv

setwd 명령어를 통해서 자료가 있는 작업 공간을 설정할 수 있습니다. 설정 후에서는 dir()을 통해 파일의 이름을 확인 할 수 있습니다. read.csv를 통해서 자료를 R에서 사용할 수 있게 됩니다.

```
setwd("D:/Rt")
dir()
mydata <- read.csv("MyData2017.csv", as.is=TRUE)</pre>
```

3.2 Theoph 데이타

R에 기본적으로 들어있는 Theoph 약동학 자료에 대해 살펴보겠습니다.

```
1 79.6 4.02 0.57 6.57
## 3
           1 79.6 4.02 1.12 10.50
## 4
## 5
           1 79.6 4.02 2.02
                              9.66
           1 79.6 4.02 3.82
## 6
                              8.58
           1 79.6 4.02 5.10 8.36
## 7
           1 79.6 4.02 7.03 7.47
## 8
           1 79.6 4.02
## 9
                        9.05 6.89
## 10
           1 79.6 4.02 12.12
                              5.94
## 11
           1 79.6 4.02 24.37
                              3.28
tail(Theoph, n = 11)
       Subject Wt Dose
##
                         Time conc
## 122
            12 60.5 5.3
                         0.00 0.00
## 123
           12 60.5 5.3 0.25 1.25
## 124
           12 60.5
                   5.3 0.50 3.96
## 125
           12 60.5
                    5.3
                         1.00 7.82
## 126
           12 60.5
                   5.3 2.00 9.72
## 127
           12 60.5
                   5.3 3.52 9.75
                   5.3 5.07 8.57
## 128
           12 60.5
## 129
           12 60.5
                    5.3 7.07 6.59
## 130
           12 60.5 5.3 9.03 6.11
## 131
           12 60.5 5.3 12.05 4.57
            12 60.5 5.3 24.15 1.17
## 132
```

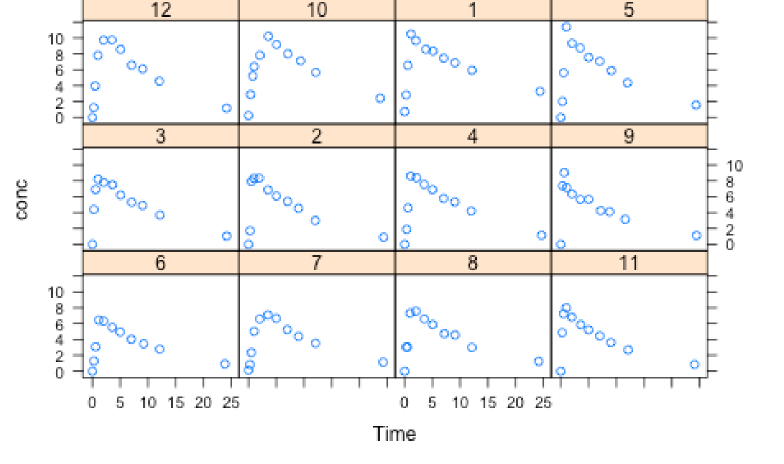
R console에서 ?Theoph를 타이핑 치면 좀 더 자세한 정보를 얻을 수 있습니다.

3.3 lattice

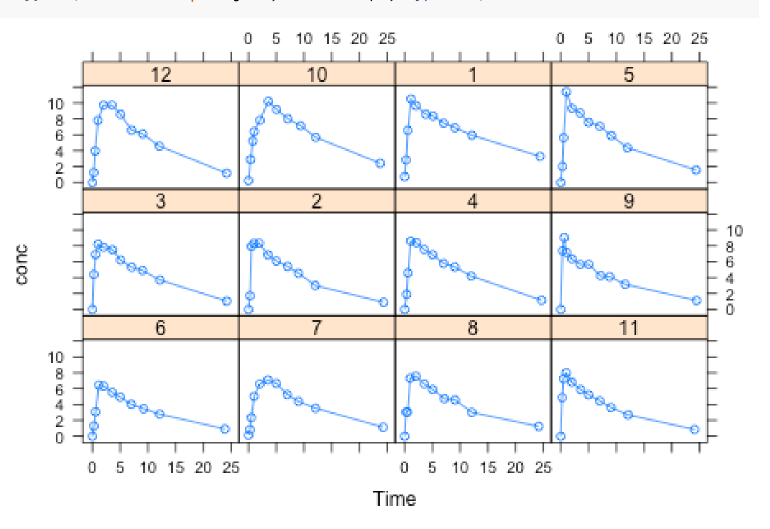
lattice 패키지를 불러온 뒤 그림을 그려보겠습니다. (Sarkar, 2017)

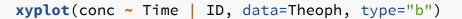
```
library(lattice) # trellis

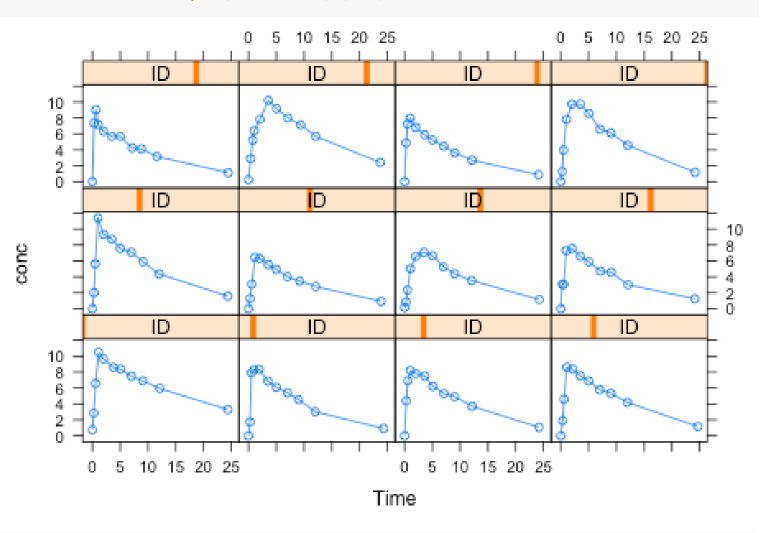
xyplot(conc ~ Time | Subject, data=Theoph)
```



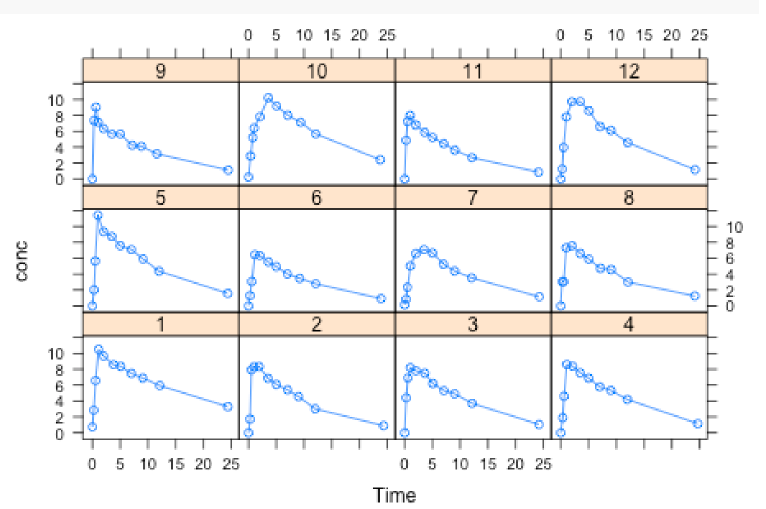








xyplot(conc ~ Time | as.factor(ID), data=Theoph, type="b")



```
write.csv(Theoph, "Theoph.csv", row.names=FALSE, quote=FALSE, na="")
```

3.4 Subseting and write.csv

자료를 편집하고, subset을 만들고 각각을 파일로 저장하는 방법에 대해 알아보겠습니다.

```
IDs = sort(unique(Theoph[,"ID"])) ; IDs
   [1] 1 2 3 4 5 6 7 8 9 10 11 12
nID = length(IDs) ; nID
## [1] 12
demog = unique(Theoph[,c("ID","Wt")])
colnames(demog) = c("ID", "BWT")
write.csv(demog, "1-demog.csv", row.names=FALSE, quote=FALSE, na="")
DV = Theoph[,c("ID","Time", "conc")]
colnames(DV) = c("ID", "TIME", "DV")
write.csv(DV, "3-DV.csv", row.names=FALSE, quote=FALSE, na="")
adm = cbind(IDs, rep(0, nID), rep(320, nID))
colnames(adm) = c("ID", "TIME", "AMT")
write.csv(adm, "2-adm.csv", row.names=FALSE, quote=FALSE, na="")
demog = read.csv("1-demog.csv", as.is=TRUE)
adm = read.csv("2-adm.csv", as.is=TRUE)
dv = read.csv("3-dv.csv", as.is=TRUE)
AdmDv = merge(adm, dv, by=intersect(colnames(adm), colnames(dv)), all=TRUE)
AdmDv
##
       ID
          TIME AMT
                       DV
## 1
       1 0.00 320
                    0.74
## 2
        1 0.25
                NA
                    2.84
## 3
        1 0.57
                 NA 6.57
        1 1.12
                NA 10.50
## 4
        1 2.02
## 5
                 NA 9.66
        1 3.82
                NA 8.58
## 6
                NA 8.36
## 7
        1 5.10
## 8
        1
         7.03
                 NA
                    7.47
## a
          9 05 NA 6 89
```

##	11	1	24.37	NA	3.28
##	12	2	0.00	320	0.00
##	13	2	0.27	NA	1.72
##	14	2	0.52	NA	7.91
##	15	2	1.00	NA	8.31
##	16	2	1.92	NA	8.33
##	17	2	3.50	NA	6.85
##	18	2	5.02	NA	6.08
##	19	2	7.03	NA	5.40
##	20	2	9.00	NA	4.55
##	21	2	12.00	NA	3.01
##	22	2	24.30	NA	0.90
##	23	3	0.00	320	0.00
##	24	3	0.27	NA	4.40
##	25	3	0.58	NA	6.90
##	26	3	1.02	NA	8.20
##	27	3	2.02	NA	7.80
##	28	3	3.62	NA	7.50
##	29	3	5.08	NA	6.20
##	30	3	7.07	NA	5.30
##	31	3	9.00	NA	4.90
##	32	3	12.15	NA	3.70
##	33	3	24.17	NA	1.05
##	34	4	0.00	320	0.00
##	35	4	0.35	NA	1.89
##	36	4	0.60	NA	4.60
##	37	4	1.07	NA	8.60
##	38	4	2.13	NA	8.38
##	39	4	3.50	NA	7.54
##	40	4	5.02	NA	6.88
##	41	4	7.02	NA	5.78
##	42	4	9.02	NA	5.33
##	43	4	11.98	NA	4.19
##	44	4	24.65	NA	1.15
##	45	5	0.00	320	0.00
##	46	5	0.30	NA	2.02
##	47	5	0.52	NA	5.63
##	48	5	1.00	NA	11.40
##	49	5	2.02	NA	9.33
##	50	5	3.50	NA	8.74
##	51	5	5.02	NA	7.56
##	52	5	7.02	NA	7.09
##	53	5	9.10	NA	5.90

	J		12.00	117.1	1.01
##	55	5	24.35	NA	1.57
##	56	6	0.00	320	0.00
##	57	6	0.27	NA	1.29
##	58	6	0.58	NA	3.08
##	59	6	1.15	NA	6.44
##	60	6	2.03	NA	6.32
##	61	6	3.57	NA	5.53
##	62	6	5.00	NA	4.94
##	63	6	7.00	NA	4.02
##	64	6	9.22	NA	3.46
##	65	6	12.10	NA	2.78
##	66	6	23.85	NA	0.92
##	67	7	0.00	320	0.15
##	68	7	0.25	NA	0.85
##	69	7	0.50	NA	2.35
##	70	7	1.02	NA	5.02
##	71	7	2.02	NA	6.58
##	72	7	3.48	NA	7.09
##	73	7	5.00	NA	6.66
##	74	7	6.98	NA	5.25
##	75	7	9.00	NA	4.39
##	76	7	12.05	NA	3.53
##	77	7	24.22	NA	1.15
##	78	8	0.00	320	0.00
##	79	8	0.25	NA	3.05
##	80	8	0.52	NA	3.05
##	81	8	0.98	NA	7.31
##	82	8	2.02	NA	7.56
##	83	8	3.53	NA	6.59
##	84	8	5.05	NA	5.88
##	85	8	7.15	NA	4.73
##	86	8	9.07	NA	4.57
##	87	8	12.10	NA	3.00
##	88	8	24.12	NA	1.25
##	89	9	0.00	320	0.00
##	90	9	0.30	NA	7.37
##	91	9	0.63	NA	9.03
##	92	9	1.05	NA	7.14
##	93	9	2.02	NA	6.33
##	94	9	3.53	NA	5.66
##	95	9	5.02	NA	5.67
##	96	9	7.17	NA	4.24
##	97	9	8.80	NA	4.11

```
## 99
        9 24.43
                   NA
                       1.12
## 100 10
            0.00 320
                       0.24
## 101 10
            0.37
                   NA
                       2.89
## 102 10
            0.77
                   NA
                       5.22
## 103 10
            1.02
                   NA
                       6.41
  104 10
##
            2.05
                       7.83
                   NA
##
  105 10
            3.55
                   NA 10.21
  106 10
##
            5.05
                   NA
                       9.18
##
  107 10
            7.08
                   NA
                       8.02
  108 10
##
            9.38
                   NA
                       7.14
  109 10 12.10
##
                   NA
                       5.68
## 110 10 23.70
                       2.42
                   NA
## 111 11
            0.00 320
                       0.00
## 112 11
            0.25
                   NA
                       4.86
## 113 11
            0.50
                       7.24
                   NA
## 114 11
            0.98
                       8.00
                   NA
## 115 11
            1.98
                       6.81
                   NA
## 116 11
            3.60
                   NA
                       5.87
## 117 11
            5.02
                   NA
                       5.22
## 118 11
            7.03
                   NA
                       4.45
## 119 11
            9.03
                       3.62
                   NA
## 120 11 12.12
                   NA
                       2.69
  121 11 24.08
                   NA
                       0.86
## 122 12
            0.00 320
                       0.00
## 123 12
            0.25
                   NA
                       1.25
## 124 12
            0.50
                   NA
                       3.96
## 125 12
            1.00
                   NA
                       7.82
## 126 12
            2.00
                       9.72
                   NA
## 127 12
            3.52
                   NA
                       9.75
## 128 12
            5.07
                   NA
                       8.57
## 129 12
            7.07
                       6.59
                   NA
## 130 12
            9.03
                       6.11
                   NA
## 131 12 12.05
                   NA
                       4.57
## 132 12 24.15
                   NA
                       1.17
자료를 병합(merge)해 보겠습니다.
```

```
DataAll
                 TIME AMT
                               DV
##
       ΙD
            BWT
## 1
        1 79.6
                 0.00 320
                             0.74
## 2
        1 79.6
                 0.25
                        NA
                             2.84
## 3
        1 79.6
                 0.57
                        NA
```

DataAll = merge(demog, AdmDv, by=c("ID"), all=TRUE)

n n		_	, , , ,	1,12	117.1	10.00
##	5	1	79.6	2.02	NA	9.66
##	6	1	79.6	3.82	NA	8.58
##	7	1	79.6	5.10	NA	8.36
##	8	1	79.6	7.03	NA	7.47
##	9	1	79.6	9.05	NA	6.89
##	10	1	79.6	12.12	NA	5.94
##	11	1	79.6	24.37	NA	3.28
##	12	2	72.4	0.00	320	0.00
##	13	2	72.4	0.27	NA	1.72
##	14	2	72.4	0.52	NA	7.91
##	15	2	72.4	1.00	NA	8.31
##	16	2	72.4	1.92	NA	8.33
##	17	2	72.4	3.50	NA	6.85
##	18	2	72.4	5.02	NA	6.08
##	19	2	72.4	7.03	NA	5.40
##	20	2	72.4	9.00	NA	4.55
##	21	2	72.4	12.00	NA	3.01
##	22	2	72.4	24.30	NA	0.90
##	23	3	70.5	0.00	320	0.00
##	24	3	70.5	0.27	NA	4.40
##	25	3	70.5	0.58	NA	6.90
##	26	3	70.5	1.02	NA	8.20
##	27	3	70.5	2.02	NA	7.80
##	28	3	70.5	3.62	NA	7.50
##	29	3	70.5	5.08	NA	6.20
##	30	3	70.5	7.07	NA	5.30
##	31	3	70.5	9.00	NA	4.90
##	32	3	70.5	12.15	NA	3.70
##	33	3	70.5	24.17	NA	1.05
##	34	4	72.7	0.00	320	0.00
##	35	4	72.7	0.35	NA	1.89
##	36	4	72.7	0.60	NA	4.60
##	37	4	72.7	1.07	NA	8.60
##	38	4	72.7	2.13	NA	8.38
##	39	4	72.7	3.50	NA	7.54
##	40	4	72.7	5.02	NA	6.88
##	41	4	72.7	7.02	NA	5.78
##	42	4	72.7	9.02	NA	5.33
##	43	4	72.7	11.98	NA	4.19
##	44	4	72.7	24.65	NA	1.15
##	45	5	54.6	0.00	320	0.00
##	46	5	54.6	0.30	NA	2.02
##	47	5	54.6	0.52	NA	5.63

11 11	-10		0 110	1.00	1171	11. 10
##	49	5	54.6	2.02	NA	9.33
##	50	5	54.6	3.50	NA	8.74
##	51	5	54.6	5.02	NA	7.56
##	52	5	54.6	7.02	NA	7.09
##	53	5	54.6	9.10	NA	5.90
##	54	5	54.6	12.00	NA	4.37
##	55	5	54.6	24.35	NA	1.57
##	56	6	80.0	0.00	320	0.00
##	57	6	80.0	0.27	NA	1.29
##	58	6	80.0	0.58	NA	3.08
##	59	6	80.0	1.15	NA	6.44
##	60	6	80.0	2.03	NA	6.32
##	61	6	80.0	3.57	NA	5.53
##	62	6	80.0	5.00	NA	4.94
##	63	6	80.0	7.00	NA	4.02
##	64	6	80.0	9.22	NA	3.46
##	65	6	80.0	12.10	NA	2.78
##	66	6	80.0	23.85	NA	0.92
##	67	7	64.6	0.00	320	0.15
##	68	7	64.6	0.25	NA	0.85
##	69	7	64.6	0.50	NA	2.35
##	70	7	64.6	1.02	NA	5.02
##	71	7	64.6	2.02	NA	6.58
##	72	7	64.6	3.48	NA	7.09
##	73	7	64.6	5.00	NA	6.66
##	74	7	64.6	6.98	NA	5.25
##	75	7	64.6	9.00	NA	4.39
##	76	7	64.6	12.05	NA	3.53
##	77	7	64.6	24.22	NA	1.15
##	78	8	70.5	0.00	320	0.00
##	79	8	70.5	0.25	NA	3.05
##	80	8	70.5	0.52	NA	3.05
##	81	8	70.5	0.98	NA	7.31
##	82	8	70.5	2.02	NA	7.56
##	83	8	70.5	3.53	NA	6.59
##	84	8	70.5	5.05	NA	5.88
##	85	8	70.5	7.15	NA	4.73
##	86	8	70.5	9.07	NA	4.57
##	87	8	70.5	12.10	NA	3.00
##	88	8	70.5	24.12	NA	1.25
##	89	9	86.4	0.00	320	0.00
##	90	9	86.4	0.30	NA	7.37
##	91	9	86.4	0.63	NA	9.03

##	93	9	86.4	2.02	NA	6.33
##	94	9	86.4	3.53	NA	5.66
##	95	9	86.4	5.02	NA	5.67
##	96	9	86.4	7.17	NA	4.24
##	97	9	86.4	8.80	NA	4.11
##	98	9	86.4	11.60	NA	3.16
##	99	9	86.4	24.43	NA	1.12
##	100	10	58.2	0.00	320	0.24
##	101	10	58.2	0.37	NA	2.89
##	102	10	58.2	0.77	NA	5.22
##	103	10	58.2	1.02	NA	6.41
##	104	10	58.2	2.05	NA	7.83
##	105	10	58.2	3.55	NA	10.21
##	106	10	58.2	5.05	NA	9.18
##	107	10	58.2	7.08	NA	8.02
##	108	10	58.2	9.38	NA	7.14
##	109	10	58.2	12.10	NA	5.68
##	110	10	58.2	23.70	NA	2.42
##	111	11	65.0	0.00	320	0.00
##	112	11	65.0	0.25	NA	4.86
##	113	11	65.0	0.50	NA	7.24
##	114	11	65.0	0.98	NA	8.00
##	115	11	65.0	1.98	NA	6.81
##	116	11	65.0	3.60	NA	5.87
##	117	11	65.0	5.02	NA	5.22
##	118	11	65.0	7.03	NA	4.45
##	119	11	65.0	9.03	NA	3.62
##	120	11	65.0	12.12	NA	2.69
##	121	11	65.0	24.08	NA	0.86
##	122	12	60.5	0.00	320	0.00
##				0.25		1.25
##				0.50		
##				1.00		
##				2.00		
##				3.52		
##	128	12	60.5	5.07	NA	8.57
##				7.07		
##				9.03		
##	131			12.05		
##	132	12	60.5	24.15	NA	1.17

Frequently Used Functions

2017-04-05 배균섭 교수님 강의

자주 쓰는 함수 및 명령어에 대해 알아보겠습니다.

4.1 Command

```
# 2017-04-05 R-intro.pdf Chapter 08
pois
# ?dbeta
dnorm(0)
pnorm(0)
1 - pnorm(1.96)
# ?pnorm
pnorm(1.96, lower.tail=FALSE)
qnorm(0.5)
qnorm(0.975)
format(qnorm(0.975), digits=22)
rnorm(5)
rnorm(5, 10, 1)
x = rnorm(100, 10, 1)
mean(x)
sd(x)
```

```
2*pt(-2.43, df = 1000)
qnorm(0.995)
qf(0.01, 2, 7, lower.tail = FALSE)
# ?fivenum
faithful
str(faithful)
eruptions
attach(faithful)
eruptions
waiting
stem(waiting)
sort(eruptions)
hist(eruptions)
hist(eruptions, seq(1.6, 5.2, 0.2), prob=TRUE)
lines(density(eruptions, bw=0.1))
rug(eruptions)
# ?hist
# ?density
lines(density(eruptions, bw="SJ"), lty=3)
plot(ecdf(eruptions), do.points=FALSE, verticals=TRUE)
# ?plot
ecdf(eruptions)
x = ecdf(eruptions)
Χ
str(x)
x()
plot(ecdf(eruptions), do.points=FALSE)
plot(ecdf(eruptions))
long <- eruptions[eruptions > 3]
x \leftarrow seq(3, 5.4, 0.01)
pnorm(x, mean=mean(long), sd=sqrt(var(long)))
# ?par
x \leftarrow rt(250, df = 5)
qqnorm(x); qqline(x)
curve(dnorm, -5, 5)
y = density(x)
```

```
cines (y, ccy-5)
# ?ppoints
ppoints(250)
ppoints(10)
qqplot(qt(ppoints(250), df = 5), x, xlab = "Q-Q plot for t dsn")
windows()
qqplot(qt(runif(250), df = 5), x, xlab = "Q-Q plot for t dsn")
# ?shapiro.test
# ?ks.test
# ?t.test
A = c(79.98, 80.04, 80.02, 80.04, 80.03, 80.03, 80.04, 79.97, 80.05, 80.03, 80.02, 80.
B = c(80.02, 79.94, 79.98, 79.97, 79.97, 80.03, 79.95, 79.97)
boxplot(A, B)
t.test(A, B)
var.test(A, B)
t.test(A, B, var.equal=TRUE)
wilcox.test(A, B)
plot(ecdf(A), do.points=FALSE, verticals=TRUE, xlim=range(A, B))
plot(ecdf(B), do.points=FALSE, verticals=TRUE, add=TRUE)
ks.test(A, B)
# Chapter 9 Grouping, loops and conditional execution
# { } does grouping
# Usefulness of loops: for >> while >> repeat
for (i in 1:10) {
 print(2*i)
}
for (i in 1:10) print(2*i)
while (
           ) {
# Statements
}
# # if ~ else ~
# if ( ) {
# # Statements 1
# } else {
# # Statements 2
```

```
# if ( ) # Statement1
# else # Statement2
# if ( ) {
# # Statements 1
# } else if ( ) {
# # Statements 2
# } else if ( ) {
# # Statements 3
# } else {
# # Statements 4
# }
#
# Chapter 10 Writing your own functions
Square = function(x=0)
{
 return(x*x)
twosam = function(y1, y2)
{
 n1 = length(y1)
 n2 = length(y2)
 yb1 = mean(y1)
 yb2 = mean(y2)
 s1 = var(y1)
 s2 = var(y2)
 s = ((n1 - 1)*s1 + (n2 - 1)*s2)/(n1 + n2 - 2)
 tst = (yb1 - yb2)/sqrt(s*(1/n1 + 1/n2))
 return (tst)
}
x = rnorm(10)
y = rt(10, 5)
twosam(x, y)
```

```
1. ccsc - runction(y1, y2)
{
  n1 = length(y1)
  n2 = length(y2)
  yb1 = mean(y1)
  yb2 = mean(y2)
  s1 = var(y1)
  s2 = var(y2)
  s = ((n1 - 1)*s1 + (n2 - 1)*s2)/(n1 + n2 - 2)
  tst = (yb1 - yb2)/sqrt(s*(1/n1 + 1/n2))
  DF = n1 + n2 - 2
  p.val = 2*(1 - pt(abs(tst), df=DF))
  Res = list(tst, DF, p.val, yb1, yb2)
  names(Res) = c("t", "df", "p-value", "mean of x", "mean of y")
  return (Res)
}
res = T.test(x, y)
t.test(x, y)
bslash = function(X, y)
{
 X = qr(X)
  return (qr.coef(X, y))
regcoeff = bslash(Xmat, yvar)
"%^%" = function(S, pow) with(eigen(S), vectors %*% (abs(values)^pow * t(vectors)))
M = matrix(c(2,1,1,2), nrow=2) ; M
M %^% 0.5
sqrtM = M%^{0}.5; sqrtM
sqrtM %*% sqrtM
```

```
fun1 = function(f, a, b, fa, fb, a0, eps, lim, fun)
  ## function 'fun1'is only visible inside 'area'
    d = (a + b)/2
    h = (b - a)/4
    fd = f(d)
    a1 = h * (fa + fd)
    a2 = h * (fd + fb)
    if (abs(a0 - a1 - a2) < eps | lim == 0)
      return (a1 + a2)
    else {
      return (fun(f, a, d, fa, fd, a1, eps, lim - 1, fun) + fun(f, d, b, fd, fb, a2, e
    }
  }
  fa = f(a)
  fb = f(b)
  a0 = ((fa + fb) * (b - a))/2
  fun1(f, a, b, fa, fb, a0, eps, lim, fun1)
}
area(dnorm, 0, 1)
integrate(dnorm, 0, 1)
pnorm(1) - pnorm(0)
f = function(x)
  y = 2 \times x
  print(x)
 print(y)
  print(z)
}
f(1)
z = 3
f(1)
cube = function(n) {
  sq = function() n*n
  n*sq()
```

```
cube(5)
open.account = function(total)
{
  list(
    deposit = function(amount)
      if(amount <= 0)</pre>
      stop("Deposits must be positive!\n")
      total <<- total + amount
      cat(amount, "deposited. Your balance is", total, "\n\n")
    },
    withdraw = function(amount)
      if(amount > total)
      stop("You don't have that much money!\n")
      total <<- total - amount
      cat(amount, "withdrawn. Your balance is", total, "\n\n")
    },
    balance = function()
      cat("Your balance is", total, "\n\n")
    }
  )
}
ross = open.account(100)
robert = open.account(200)
ross$balance()
robert$balance()
ross$deposit(50)
ross$balance()
ross$withdraw(500)
# More basic keywords and functions
1 \%in% c(1,2,3,4)
5 \% in\% c(1,2,3,4)
is.finite(Inf)
prod(1:3)
cummax(1:10)
```

Keyword	Bae Freq	Essential	Comment
?	Н	Y	
str	M	Y	strucutre

```
cummax(10:1)
# ?xor
x = 11:20
which (x==3)
which (x==13)
length(x)
y = "my string"
length(y)
nchar(y)
strsplit(y, " ")
strsplit(y, " ")[[1]]
substr(y, 4, 5)
sample(1:10)
sample(1:10, 20)
sample(1:10, 20, replace=TRUE)
sample(rep(1:10,2))
## Error: <text>:99:12: 예기치 않은 ')'입니다
## 98:
## 99: while (
                  )
##
```

Keyword	Bae Freq	Essential	Comment
%in%	M	Y	Value Matching
match	M	N	Value Matching
=	Н	Y	
<-	L	N	
«-	M	Y	
head	Н	N	
tail	M	N	
subset	L	N	Subsetting Vectors, Matrices and Data Frames
with	L	N	Evaluate an Expression in a Data Environment
assign	L	N	Assign a Value to a Name
get	L	N	Return the Value of a Named Object

TABLE 4.3: The basics - Comparison

Keyword	Bae Freq	Essential	Comment
all.equal	L	N	Test if Two Objects are (Nearly) Equal
identical	L	N	Test Objects for Exact Equality
!=, ==, >, >=, <, <=	Н	Y	Comparison Operator
is.na	Н	Y	
complete.cases	L	N	Find Complete Cases
is.finite	M	Y	

Keyword	Bae Freq	Essential	Comment
*,+,-,/,^	Н	Y	Math operator
%%	M	Y	Modulus
%/%	L	N	Integer division
abs	Н	Y	
sign	M	N	
acos	L	Y	
asin	L	Y	
atan	L	Y	
atan2	L	Y	
sin	L	Y	
cos	L	Y	
tan	L	Y	
ceiling	Н	Y	
floor	Н	N	
round	Н	Y	
trunc	Н	N	
signif	M	Y	rounds the values in its first argument to the specified num
exp	Н	Y	-
log	Н	Y	
log10	L	Y	
log2	L	Y	
sqrt	Н	N	
max	Н	Y	
min	Н	Y	
prod	L	N	
sum	Н	Y	
cummax	L	N	
cummin	L	N	
cumprod	L	N	
cumsum	L	N	
diff	L	N	
pmax	L	N	pairwise max
pmin	L	N	pairwise min
range	L	N	•
mean	H	Y	
median	Н	Y	
cor	Н	Y	
sd	Н	Y	

Keyword	Bae Freq	Essential	Comment
function	Н	Y	
missing	M	Y	Does a Formal Argument have a Value?
on.exit	L	Y	
return	Н	N	
invisible	L	N	Change the Print Mode to Invisible

TABLE 4.6: The basics - Logical - sets

Keyword	Bae Freq	Essential	Comment
&, ,!	Н	Y	
xor	L	Y	
all	L	Y	Are All Values True?
any	L	Y	Are Some Values True?
intersect	M	Y	
union	M	Y	
setdiff	L	Y	
setequal	L	Y	
which	L	N	Which indices are TRUE?

4.2 The basics

4.3 Common data structures

4.4 Statistics

4.5 Working with R

4.6 I/O

TABLE 4.7: The basics - Vectors and matrices

Keyword	Bae Freq	Essential	Comment
С	Н	Y	
matrix	Н	Y	
# automatic coercion rules character > numeric > logical	Н	Y	
length	Н	Y	
dim	Н	Y	
ncol	Н	N	
nrow	Н	N	
cbind	Н	Y	
rbind	Н	Y	
names	M	Y	
colnames	Н	Y	
rownames	M	Y	
t	Н	Y	
diag	Н	Y	
sweep	L	N	Sweep out A
as.matrix	Н	Y	
data.matrix	L	N	Convert a Da

TABLE 4.8: The basics - Making vectors

Keyword	Bae Freq	Essential	Comment
С	Н	Y	
rep	Н	Y	
rep_len	L	N	Replicate Elements of Vectors a
seq	M	Y	
seq_len	L	N	
seq_along	L	N	
rev	M	Y	
sample	Н	Y	
choose	Н	Y	
factorial	M	Y	
combn	L	N	Generate All Combinations of n
(is/as).(character/numeric/logical/)	Н	Y	

TABLE 4.9: The basics - Lists - data.frames

Keyword	Bae Freq	Essential	Comment
list	Н	Y	
unlist	L	Y	Flatten Lists
data.frame	Н	Y	
as.data.frame	Н	Y	
split	Н	Y	
expand.grid	L	N	Create a Data Frame from All Combinations of Factor V

TABLE 4.10: The basics - Control flow

Keyword	Bae Freq	Essential	Comment
if	Н	Y	
&&	L	Y	
(short circuiting)	L	Y	
for	Н	Y	
while	L	N	
next	M	Y	
break	M	Y	
switch	L	Y	
ifelse	L	N	Conditional Element Selection

TABLE 4.11: The basics - Apply - friends

Keyword	Bae Freq	Essential	Comment
lapply	L	N	Apply a Function over a List or Vector
sapply	L	N	user-friendly version and wrapper of lapply
vapply	L	N	similar to sapply, but has a pre-specified type of return value
apply	M	N	Apply Functions Over Array Margins
tapply	L	N	Apply a Function Over a Ragged Array
replicate	L	N	Apply a Function over a List or Vector

Keyword	Bae Freq	Essential	Comment
ISOdate	L	N	
ISOdatetime	L	N	
strftime	Н	Y	Date-time Conversion Functions to and from Chara
strptime	Н	Y	Date-time Conversion Functions to and from Chara
date	M	Y	
difftime	Н	Y	
julian	L	Y	Extract Parts of a POSIXt or Date Object
months	L	N	Extract Parts of a POSIXt or Date Object
quarters	L	N	Extract Parts of a POSIXt or Date Object
weekdays	L	N	Extract Parts of a POSIXt or Date Object
library(lubridate)	L	N	

TABLE 4.13: Common data structures - Character manipulation

Keyword	Bae Freq	Essential	Comment
grep	Н	Y	Pattern Matching and Replacement
agrep	L	N	Approximate String Matching (Fuzzy Matching)
gsub	M	Y	or use sub
strsplit	Н	Y	
chartr	L	N	Character Translation and Casefolding
nchar	M	Y	
tolower	M	Y	
toupper	Н	Y	
substr	Н	Y	
paste	Н	Y	
library(stringr)	L	N	

TABLE 4.14: Common data structures - Factors

Keyword	Bae Freq	Essential	Comment
factor	M	Y	
levels	M	Y	
nlevels	L	N	
reorder	L	N	Reorder Levels of a Factor
relevel	L	N	Reorder Levels of Factor
cut	L	Y	
findInterval	L	N	Find Interval Numbers or Indices
interaction	L	N	Compute Factor Interactions
ontions(stringsAsFactors = FALSF)	T.	N	

TABLE 4.15: Common data structures - Array manipulation

Keyword	Bae Freq	Essential	Comment
array	L	N	Multi-way Arrays
dim	Н	Y	
dimnames	M	Y	
aperm	L	N	Array Transposition
library(abind)	L	N	

TABLE 4.16: Statistics - Ordering and tabulating

Keyword	Bae Freq	Essential	Comment
duplicated	L	Y	Determine Duplicated Elements
unique	Н	Y	
merge	L	N	
order	Н	Y	
rank	L	Y	
quantile	L	Y	
sort	Н	Y	
table	M	Y	
ftable	L	Y	Flat Contingency Tables

TABLE 4.17: Statistics - Linear models

Keyword	Bae Freq	Essential	Comment
fitted	L	Y	Extract Model Fitted Values
predict	Н	Y	
resid	L	Y	Extract Model Residuals
rstandard	L	Y	Regression Deletion Diagnostics
lm	Н	Y	
glm	Н	Y	
hat	L	Y	
influence.measures	M	Y	Regression Deletion Diagnostics
logLik	L	Y	
df	M	Y	
deviance	M	Y	
formula	Н	Y	
~	Н	Y	
I	Н	Y	
anova	Н	Y	
coef	M	Y	
confint	M	Y	
VCOV	Н	Y	
contrasts	L	Y	Get and Set Contrast Matrices

TABLE 4.18: Statistics - Miscellaneous tests

Keyword	Bae Freq	Essential	Comment
apropos("\\.test\$")	L	Y	

TABLE 4.19: Statistics - Random variables

Keyword	Bae Freq	Essential	Commo
(q, p, d, r) * (beta, binom, cauchy, chisq, exp, f, gamma, geom,	Н	Y	

Keyword	Bae Freq	Essential	Comment
crossprod	L	Y	Matrix Crossproduct
tcrossprod	L	N	Matrix Crossproduct
eigen	Н	Y	
qr	L	Y	
svd	L	Y	
%*%	Н	Y	
%o%	L	Y	Outer Product of Arrays
outer	Н	Y	
rcond	L	N	Compute or Estimate the Condition Number of a Matrix
solve	Н	Y	Solve a System of Equations

TABLE 4.21: Working with R - Workspace

Keyword	Bae Freq	Essential	Comment
ls	Н	Y	List Objects
exists	M	Y	
rm	M	Y	
getwd	Н	Y	
setwd	Н	Y	
q	L	Y	
source	Н	Y	
install.packages	Н	Y	
library	Н	Y	
require	Н	Y	

TABLE 4.22: Working with R - Help

Keyword	Bae Freq	Essential	Comment
help	L	N	
?	Н	Y	
help.search	L	N	
apropos	L	Y	
RSiteSearch	L	N	Search for Key Words or Phrases in Documentation
citation	L	Y	
demo	L	Y	
example	L	Y	
vignette	L	Y	View, List or Get R Source of Package Vignettes

TABLE 4.23: Working with R - Debugging

Keyword	Bae Freq	Essential	Comment
traceback	L	Y	
browser	L	Y	Environment Browser
recover	L	Y	Browsing after an Error
options(error =)	L	Y	
stop	L	Y	Stop Function Execution
warning	Н	Y	
message	L	Y	
tryCatch	L	Y	
try	L	Y	

TABLE 4.24: I/O - Output

Keyword	Bae Freq	Essential	Comment
print	Н	Y	
cat	Н	Y	
message	L	Y	
warning	Н	Y	
dput	L	N	Write an Object to a File or Recreate it
format	Н	Y	
sink	L	Y	Send R Output to a File
capture.output	L	Y	Send Output to a Character String or File

 Keyword	Bae Freq	Essential	Comment
data	L	N	Loads specified data sets, or list the available data sets
count.fields	L	N	
read.csv	Н	Y	
write.csv	Н	Y	
read.delim	L	N	
write.delim	L	N	
read.fwf	L	N	
readLines	M	Y	
writeLines	M	Y	
readRDS	L	N	Serialization Interface for Single Objects
saveRDS	L	N	
load	L	Y	
save	L	Y	
library(foreign)	L	N	

TABLE 4.26: I/O - Files and directories

Keyword	Bae Freq	Essential	Comment		
dir	Н	Y			
basename	L	Y	removes all of the path up to and including the la		
dirname	L	Y			
tools::file_ext	L	Y			
file.path	L	Y			
path.expand	L	Y	Expand File Paths		
normalizePath	L	Y	Express File Paths in Canonical Form		
file.choose	L	Y			
file.copy	L	Y			
file.create	L	Y			
file.remove	L	Y			
file.rename	L	Y			
dir.create	L	Y			
file.exists	L	Y			
file.info	L	Y			
tempdir	L	Y			
tempfile	L	Y			
download.file	L	Y			
library(downloader)	L	N			

Rstudio and some useful packages 2 - ggplot2

9주차 강의 예정 자료입니다.

이번 시간에는 ggplot2의 사용법을 예제를 통해 알아보겠습니다. (Wickham and Chang, 2016) https://rpubs.com/kimwoohyung/ggplot2의 자료를 많이 참고하였습니다. https://rpubs.com/mccannecology/53464도 좋은 자료입니다.

5.1 Introduction

'data.frame':

먼저 다음과 같은 패키지가 필요합니다.

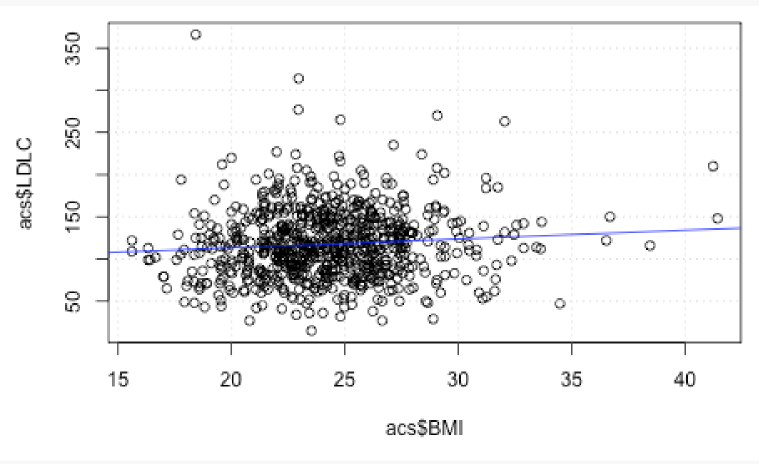
```
library(ggplot2)
library(gcookbook)
library(plyr)
library(reshape2)
library(moonBook)
```

```
"sex"
                                                                                       "Dx"
[1] "age"
                                              "cardiogenicShock" "entry"
[6] "EF"
                         "height"
                                              "weight"
                                                                  "BMI"
                                                                                       "obesi
[11] "TC"
                          "LDLC"
                                              "HDLC"
                                                                   "TG"
                                                                                        "DM"
                          "smoking"
[16] "HBP"
> str(acs)
```

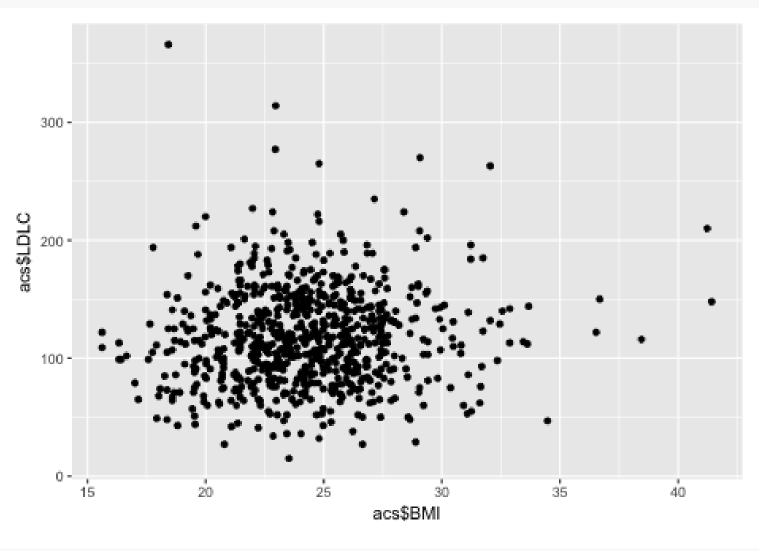
857 obs. of 17 variables:

```
$ Dx
                         "STEMI" "STEMI" "STEMI" ...
                  : chr
$ EF
                         18 18.4 20 21.8 21.8 22 24.7 26.6 28.5 31.1 ...
                   num
$ height
                         168 148 NA 165 162 153 167 160 152 168 ...
                  : num
                         72 48 NA 50 64 59 78 50 67 60 ...
$ weight
                  : num
$ BMI
                         25.5 21.9 NA 18.4 24.4 ...
                  : num
                         "Yes" "No" "No" "No" ...
$ obesity
                  : chr
$ TC
                  : num
                         215 NA NA 121 195 184 161 136 239 169 ...
$ LDLC
                         154 NA NA 73 151 112 91 88 161 88 ...
                  : int
$ HDLC
                  : int
                         35 NA NA 20 36 38 34 33 34 54 ...
$ TG
                  : int
                         155 166 NA 89 63 137 196 30 118 141 ...
                         "Yes" "No" "No" "No" ...
$ DM
                  : chr
                         "No" "Yes" "Yes" "No" ...
$ HBP
                  : chr
                         "Smoker" "Never" "Never" ...
$ smoking
                  : chr
```

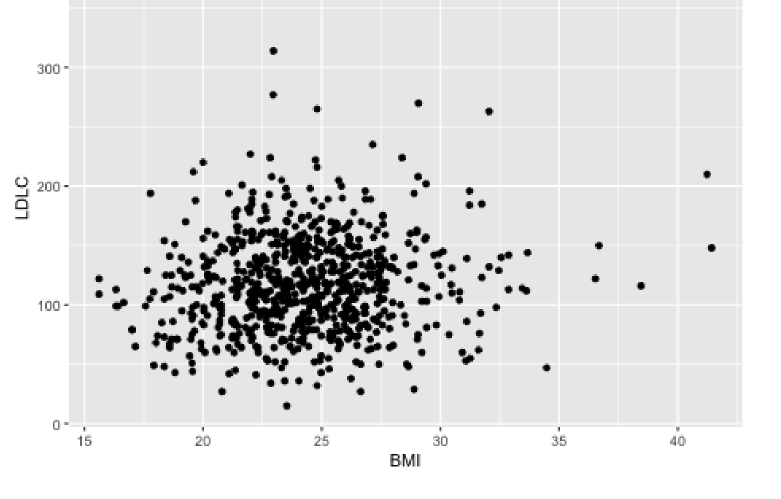
Scatter plot을 geom_point()를 사용해서 그려보도록 하겠습니다.



```
qplot(acs$BMI, acs$LDLC)
## Warning: Removed 106 rows containing missing values
## (geom point).
```

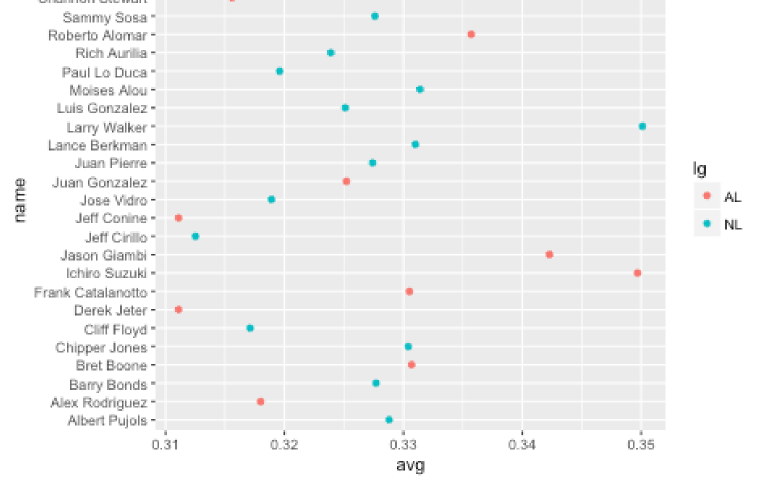


```
ggplot(acs, aes(x = BMI, y = LDLC)) + geom_point()
## Warning: Removed 106 rows containing missing values
## (geom_point).
```

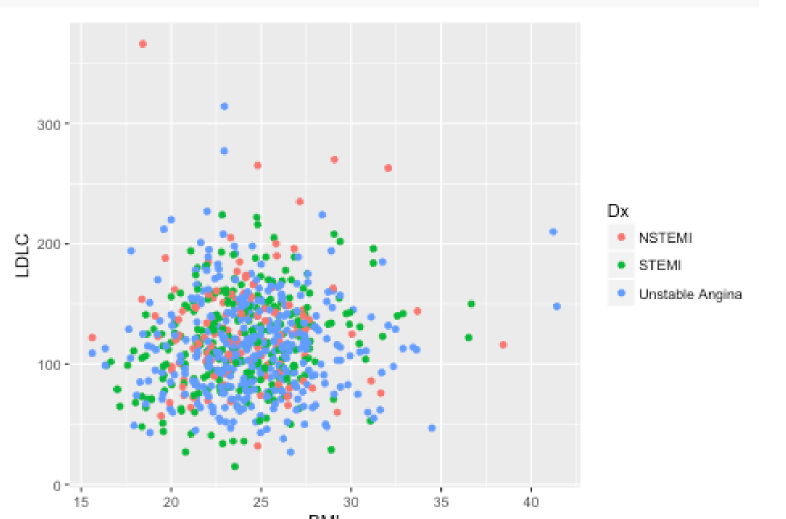


5.2 Part1

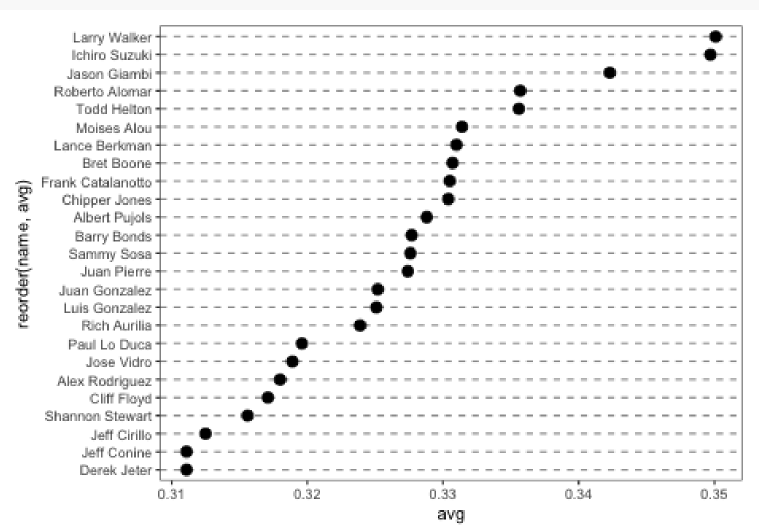
```
tophit <- tophitters2001[1:25,]
tophit <- tophit[,c("name", "lg", "avg")]
# y축이산형 그래프
ggplot(tophit, aes(x = avg, y = name, col=lg)) + geom_point()
```



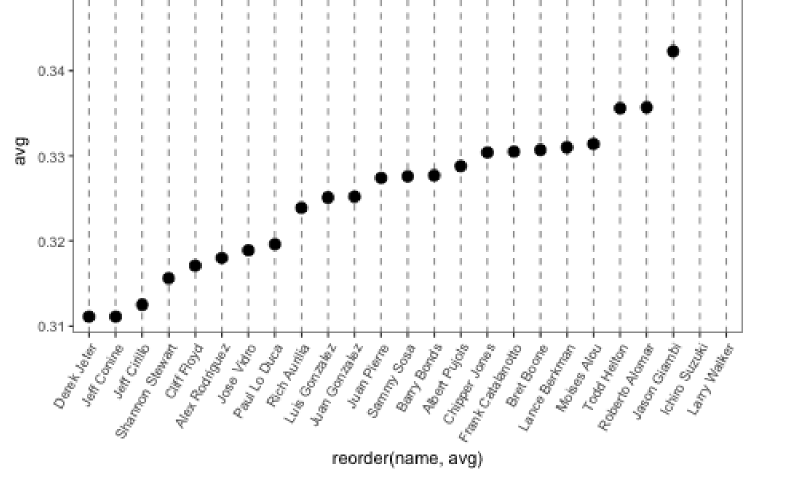
ggplot(acs, aes(x = BMI, y = LDLC, col=Dx)) + geom_point()
Warning: Removed 106 rows containing missing values
(geom_point).



그래프 정렬하기 & 그래프 격자 없애기 & 수평선 점선으로 바꾸기

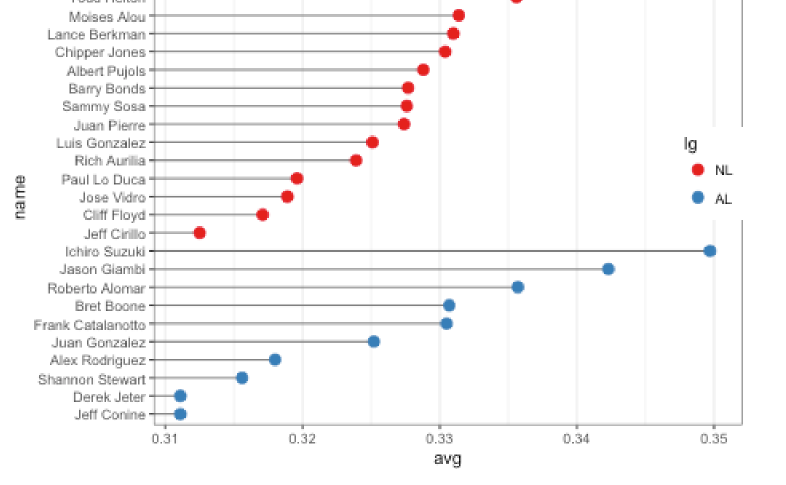


x, y축 바꿔서 그치기 & 그래프 격자 없애기 & 수직선 점선으로 바꾸기 & x축 값 정의 및 회전 ggplot(tophit, aes(x = reorder(name, avg), y = avg)) + geom_point(size = 3) + theme_bw theme(axis.text.x = element_text(angle = 60, hjust = 1), panel.grid.major.y = element_blank(), panel.grid.minor.y = element_blank(), panel.grid.major.x = element_line(color = "grey60", linetype = "dashed"))



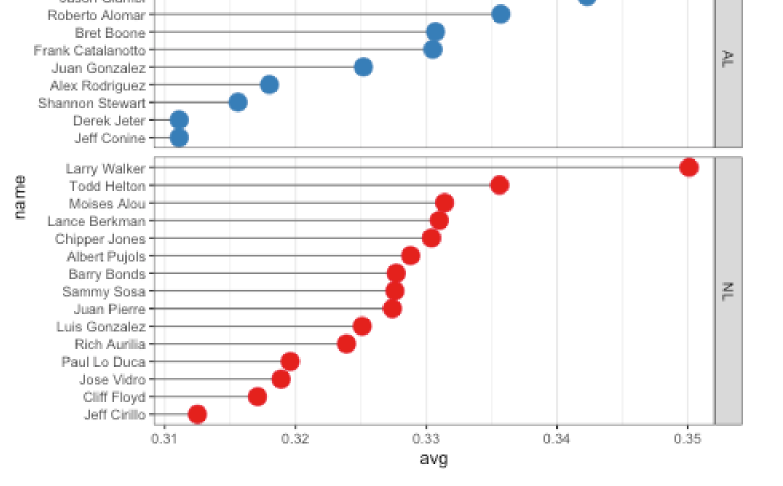
```
# 두개 변수로 정의하여 그리기
nameorder <- tophit$name[order(tophit$lg, tophit$avg)]
tophit$name <- factor(tophit$name, levels = nameorder)

# 격자 선이 그래프의 끝에서 끝까지 횡단하지 않고, 점까지만 가도록 표현 [ geom_segment ]
ggplot(tophit, aes(x = avg, y = name)) +
    geom_segment(aes(yend = name), xend = 0, color = "grey50") + geom_point(size = 3, ae
    scale_color_brewer(palette = "Set1", limits = c("NL", "AL")) + theme_bw() +
    theme(panel.grid.major.y = element_blank(), legend.position = c(1, 0.55), # 범례를 그
    legend.justification = c(1, 0.5))
```



그룹 별 그래프 분할

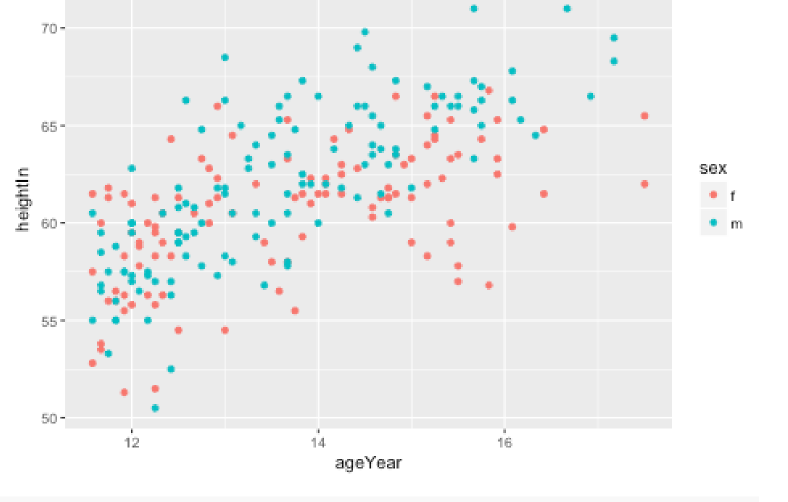
```
ggplot(tophit, aes(x = avg, y = name)) +
  geom_segment(aes(yend = name), xend = 0, color = "grey50") + geom_point(size = 5, ae
  scale_color_brewer(palette = "Set1", limits = c("NL", "AL"), guide = FALSE) + theme_
  theme(panel.grid.major.y = element_blank()) + facet_grid(lg ~ ., scales = "free_y",
```



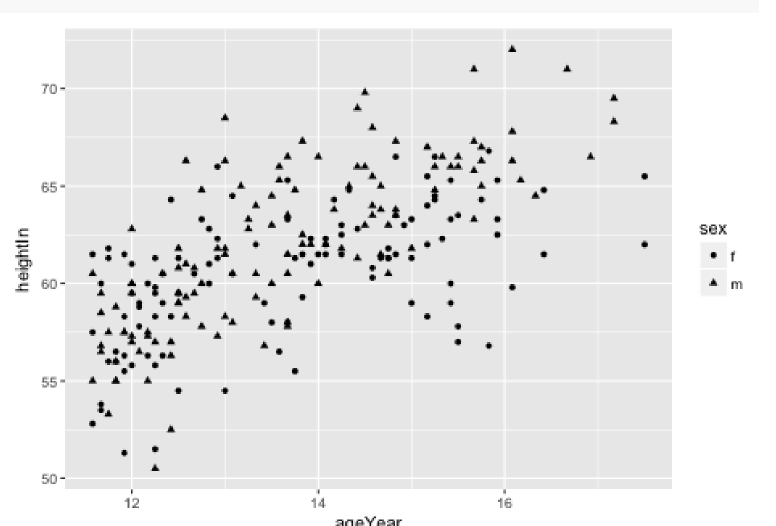
그룹 별 데이터 구별

색상기준

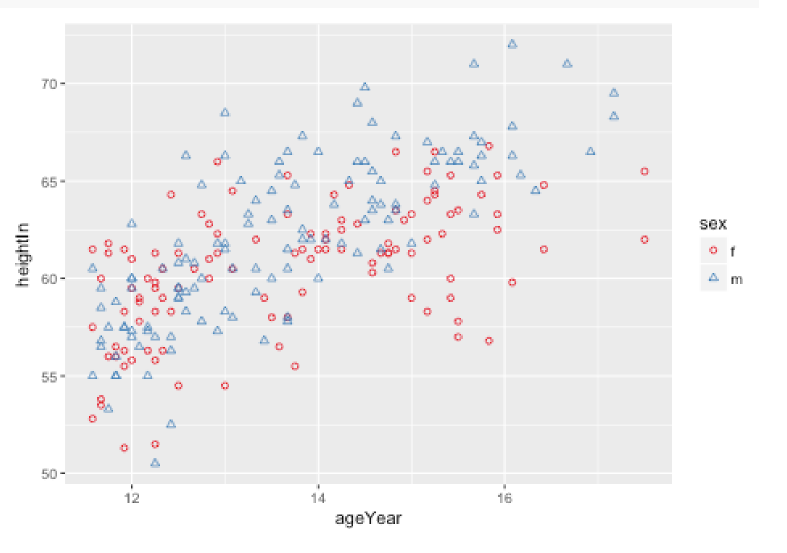
ggplot(heightweight, aes(x = ageYear, y = heightIn, color = sex)) + geom_point()



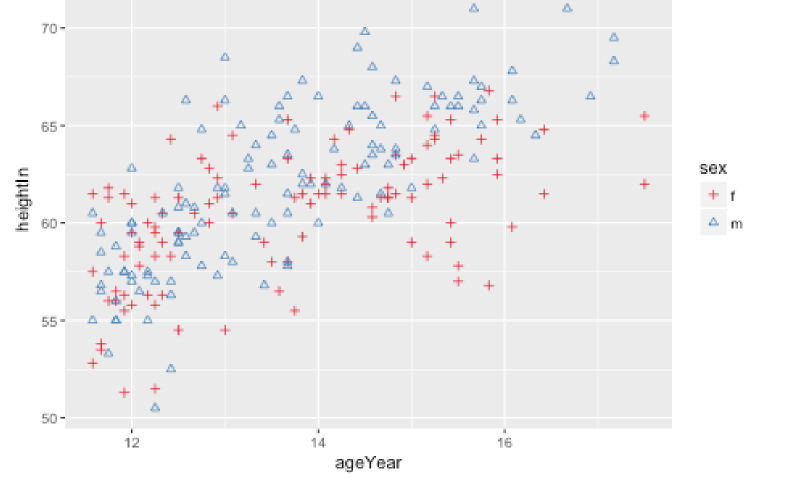
점 모양 기준
ggplot(heightweight, aes(x = ageYear, y = heightIn, shape = sex)) + geom_point()



```
ggplot(heightweight, aes(x = ageYear, y = heightIn, color = sex, shape = sex)) + geom_
scale_shape_manual(values = c(1,2)) + scale_color_brewer(palette = "Set1")
```



```
ggplot(heightweight, aes(x = ageYear, y = heightIn, color = sex, shape = sex)) + geom_
scale_shape_manual(values = c(3,2)) + scale_color_brewer(palette = "Set1")
```



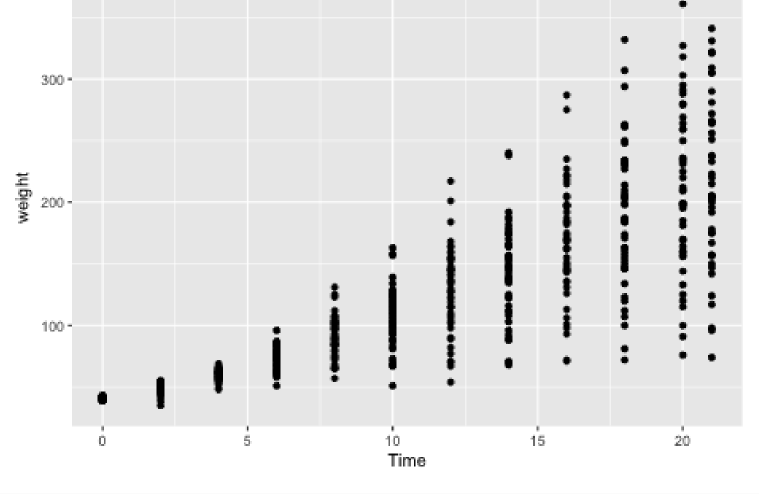
기준 정의에 따라 구별

```
hw <-heightweight
```

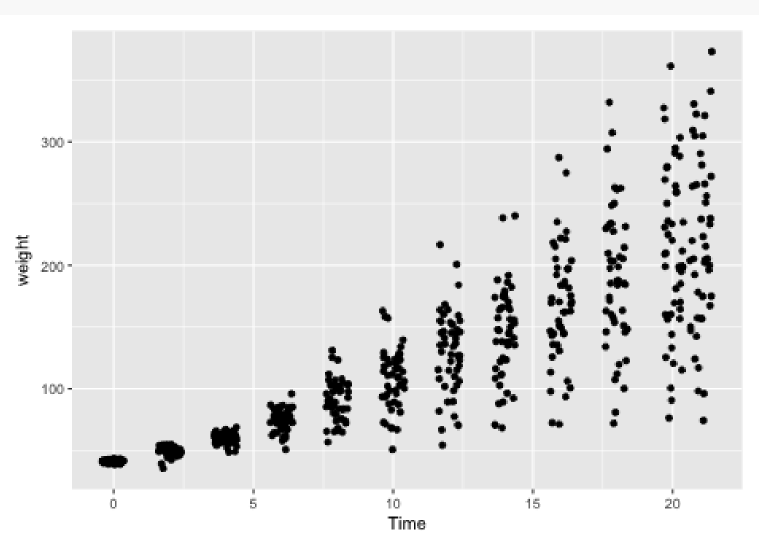
```
hw\$weightGroup <- cut(hw\$weightLb, breaks = c(-Inf, 100, Inf), labels = c("< 100", ">= c("< 100", ")= c(">= c(">= c("< 100", ")= c(">= c(
```

x축이 이산형일 때 점들을 랜덤하게 조금식 이동시켜 표현

```
ggplot(ChickWeight, aes(x = Time, y = weight)) + geom_point()
```

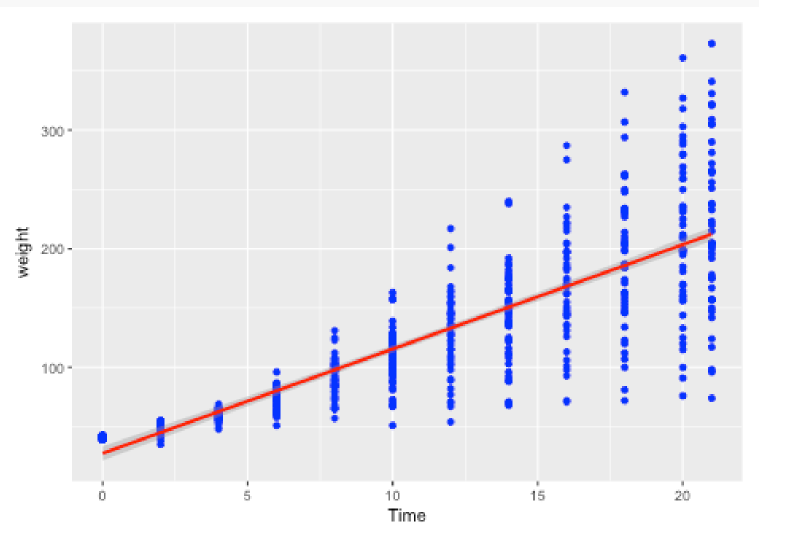




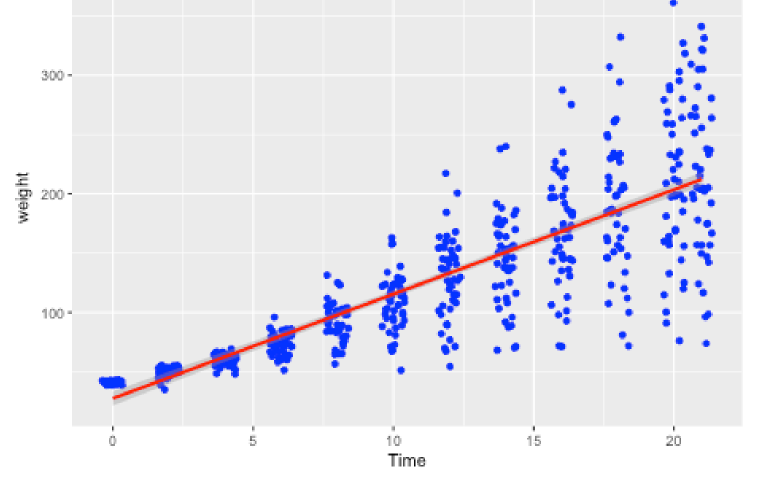


적합된 회귀선 추가하기

```
sp <- ggplot(ChickWeight, aes(x = Time, y = weight))
sp + geom_point(color = "blue") + stat_smooth(method = lm, se = TRUE, color = "red")</pre>
```

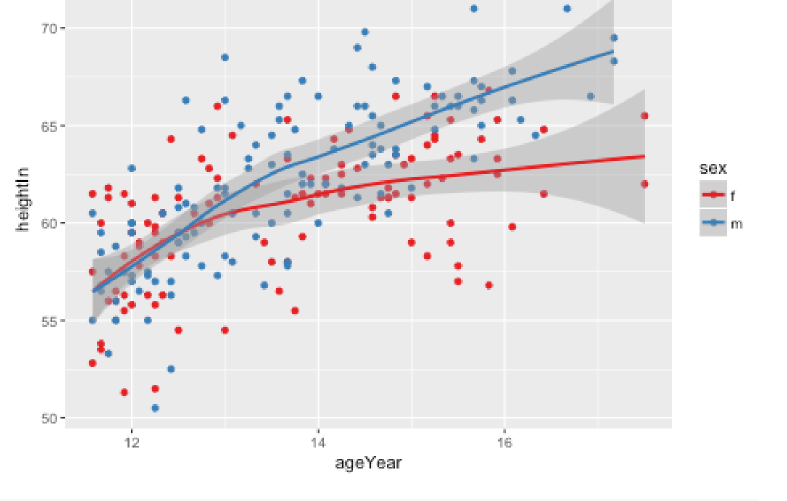


```
sp + geom_jitter(color = "blue") + stat_smooth(method = lm, se = TRUE, color = "red")
```



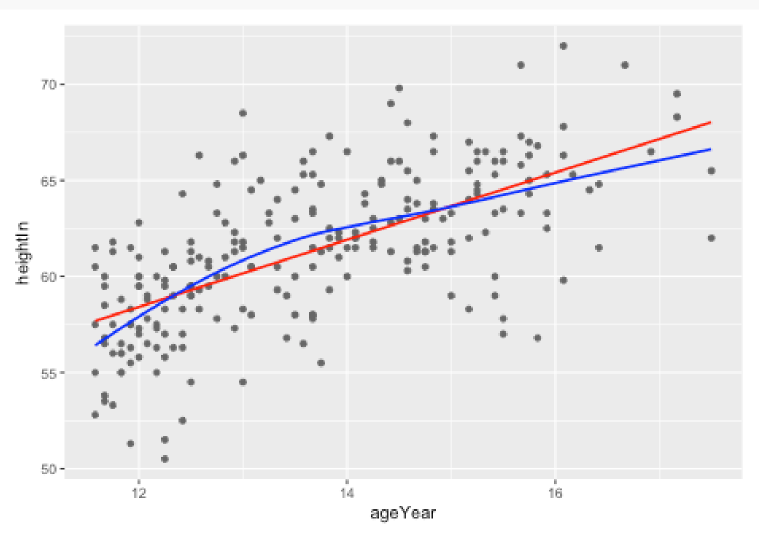
#그룹 별 회귀선 추가하기

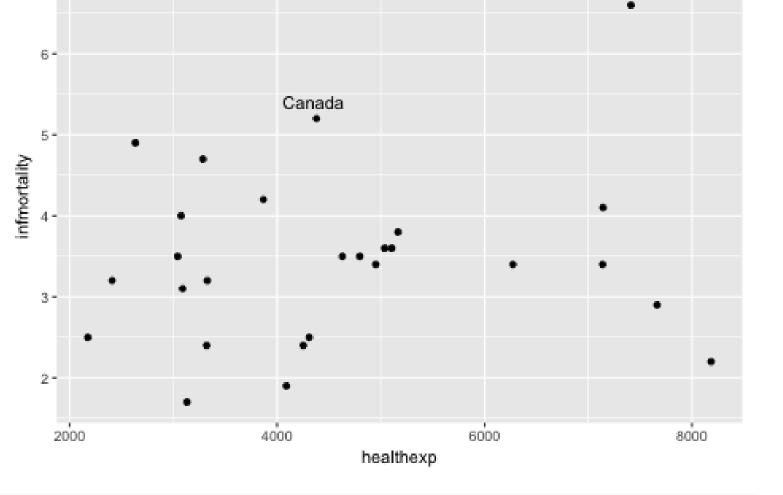
```
sps <- ggplot(heightweight, aes(x = ageYear, y = heightIn, color = sex)) +
    geom_point() + scale_color_brewer(palette = "Set1")
sps + geom_smooth()
## `geom_smooth()` using method = 'loess'</pre>
```



```
# 예측값 실제값 그래프로 표현하기 (함수)
predictvals <- function(model, xvar, yvar, xrange = NULL, sample = 100, ...){</pre>
  if(is.null(xrange)){
    if(any(class(model) %in% c("lm", "glm")))
      xrange <- range(model$model[[xvar]])</pre>
    else if(any(class(model) %in% "loess"))
      xrange <- range(model$x)</pre>
  }
  newdata <- data.frame(x = seq(xrange[1], xrange[2], length.out = sample))</pre>
  names(newdata) <- xvar</pre>
  newdata[[yvar]] <- predict(model, newdata = newdata, ...)</pre>
  newdata
}
modlinear <- lm(heightIn ~ ageYear, heightweight)</pre>
modloess <- loess(heightIn ~ ageYear, heightweight)</pre>
                <- predictvals(modlinear, "ageYear", "heightIn")</pre>
lm_predicted
loess_predicted <- predictvals(modloess, "ageYear", "heightIn")</pre>
sp <- ggplot(heightweight, aes(x = ageYear, y = heightIn)) +</pre>
```

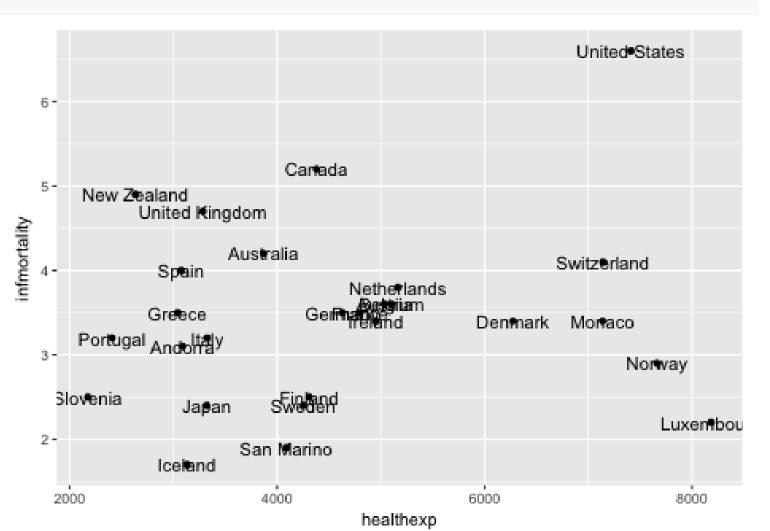
```
sp + geom_line(data = lm_predicted, color = "red", size = 0.8) +
geom_line(data = loess_predicted, color = "blue", size = 0.8)
```





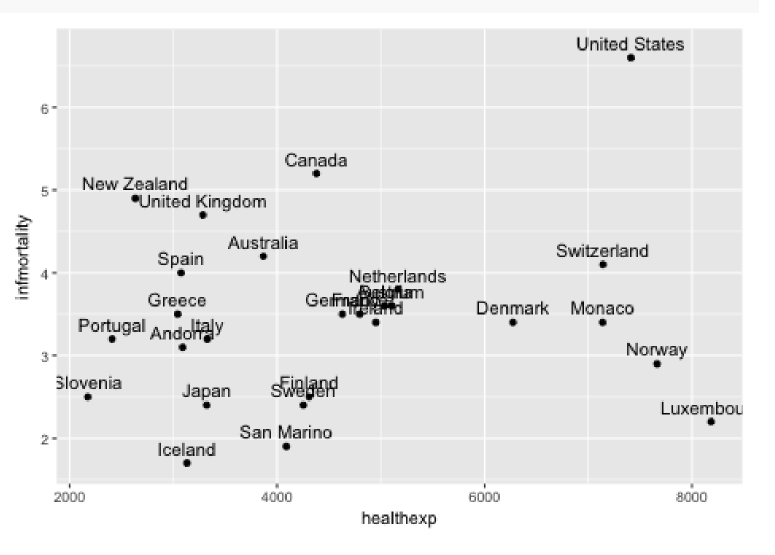
데이터 값을 라벨로 붙이기

sp + geom_text(aes(label = Name), size = 4)

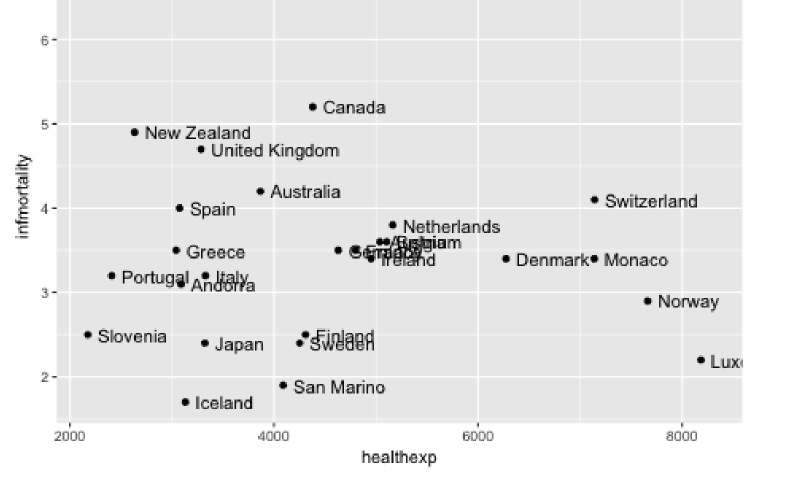


라벨의 위치를 데이터값보다 조금 크게 설정

```
sp + geom_text(aes(y = infmortality + 0.1, label = Name), size = 4, vjust = 0)
```



```
sp + geom_text(aes(x = healthexp + 100, label = Name), size = 4, hjust = 0)
```



특정 값만 라벨 붙이기

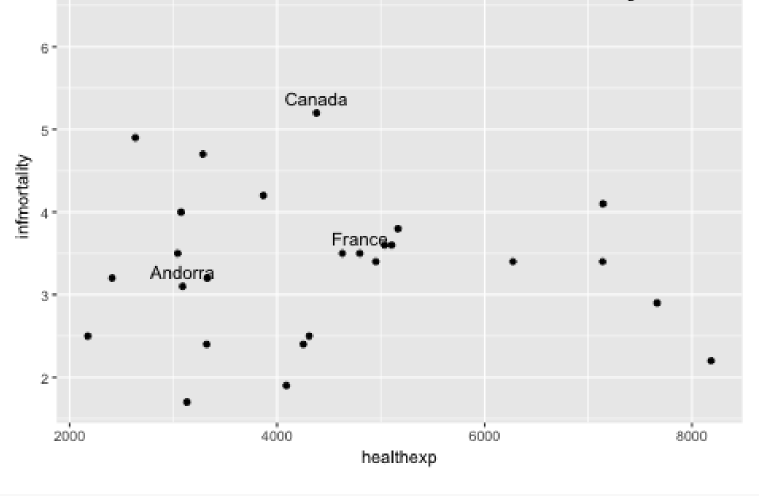
```
cdat <- subset(countries, Year == 2009 & healthexp > 2000)

cdat$Name1 <- cdat$Name
idx <- cdat$Name %in% c("Andorra", "France", "Canada")

cdat$Name1[!idx] <- NA

ggplot(cdat, aes(x = healthexp, y = infmortality)) + geom_point() + geom_text(aes(y = ## Warning: Removed 24 rows containing missing values

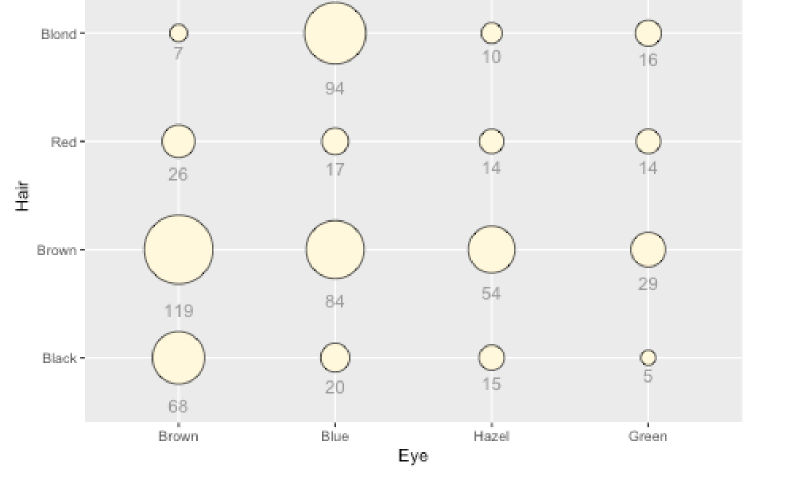
## (geom_text).</pre>
```



Warning: Removed 24 rows containing missing values (geom_text).

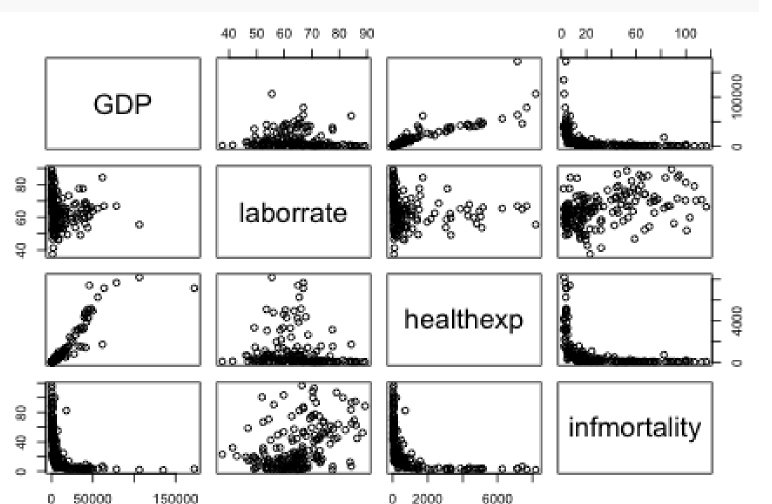
5.3 Part2

```
# 거품그래프(balloon plot)
hec <- HairEyeColor[,,"Male"] + HairEyeColor[,,"Female"]
hec <- melt(hec, value.name = "count")
ggplot(hec, aes(x = Eye, y = Hair)) + geom_point(aes(size = count), shape = 21, color
scale_size_area(max_size = 20, guide = FALSE) +
geom_text(aes(y = as.numeric(Hair)-sqrt(count)/22, label = count), vjust = 1, color
```

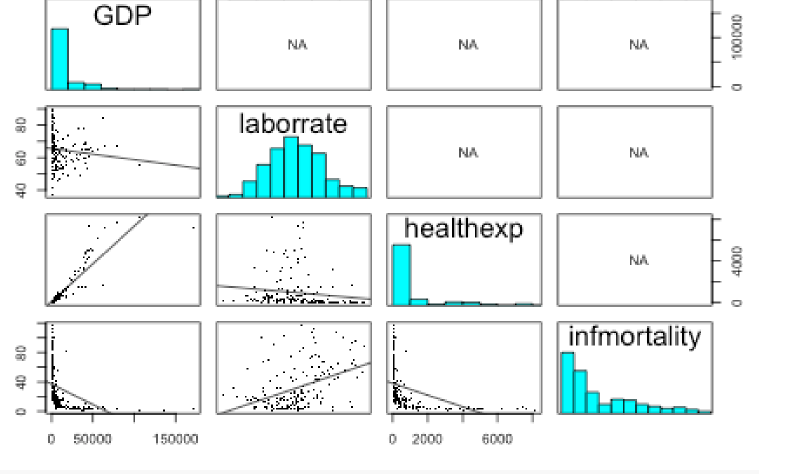


산점도 행렬 만들기

c2009 <- subset(countries, Year == 2009, select = c(Name, GDP, laborrate, healthexp, i
plot(c2009[,2:5])</pre>

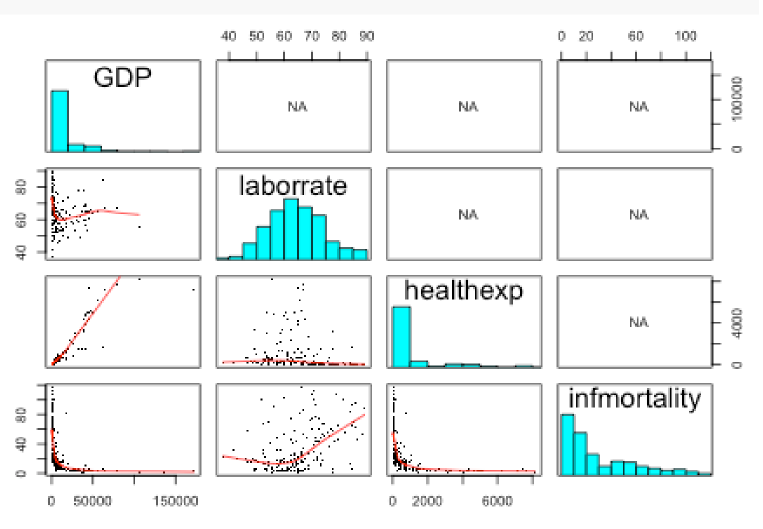


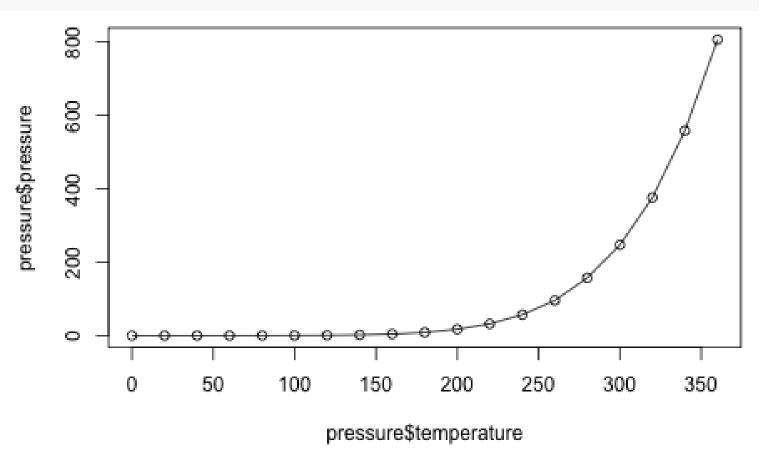
```
panel.hist <- function(x, ...)
{
  usr <- par("usr"); on.exit(par(usr))</pre>
  par(usr = c(usr[1:2], 0, 1.5) )
  h <- hist(x, plot = FALSE)
  breaks <- h$breaks; nB <- length(breaks)</pre>
  y <- h$counts; y <- y/max(y)</pre>
  rect(breaks[-nB], 0, breaks[-1], y, col = "cyan", ...)
panel.cor <- function(x, y, digits = 2, prefix = "", cex.cor, ...)</pre>
{
  usr <- par("usr"); on.exit(par(usr))</pre>
  par(usr = c(0, 1, 0, 1))
  r \leftarrow abs(cor(x, y))
  txt <- format(c(r, 0.123456789), digits = digits)[1]</pre>
  txt <- paste0(prefix, txt)</pre>
  if(missing(cex.cor)) cex.cor <- 0.8/strwidth(txt)</pre>
  text(0.5, 0.5, txt, cex = cex.cor * r)
}
panel.lm <- function(x, y, col = par("col"), bg = NA, pch = par("pch"), cex = 1, col.s
  points(x, y, pch = pch, col = col, bg = bg, cex = cex)
  abline(stats::lm(y ~ x), col = col.smooth, ...)
}
# 데이터에 대한 LOWESS선(평활선) 추가
pairs(c2009[,2:5], pch = ".", upper.panel = panel.cor, diag.panel = panel.hist, lower.
```



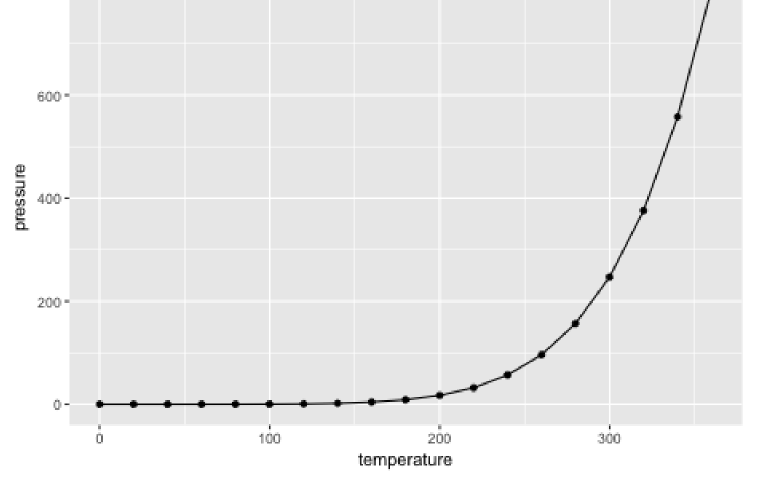
데이터에 대한 회귀적합선 추가

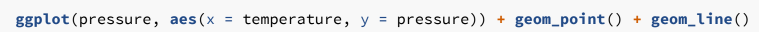
pairs(c2009[,2:5], pch = ".", upper.panel = panel.cor, diag.panel = panel.hist, lower.

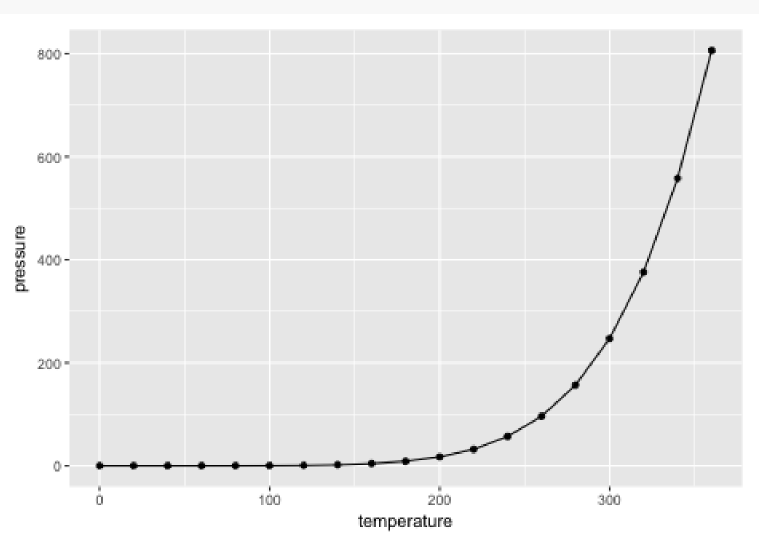




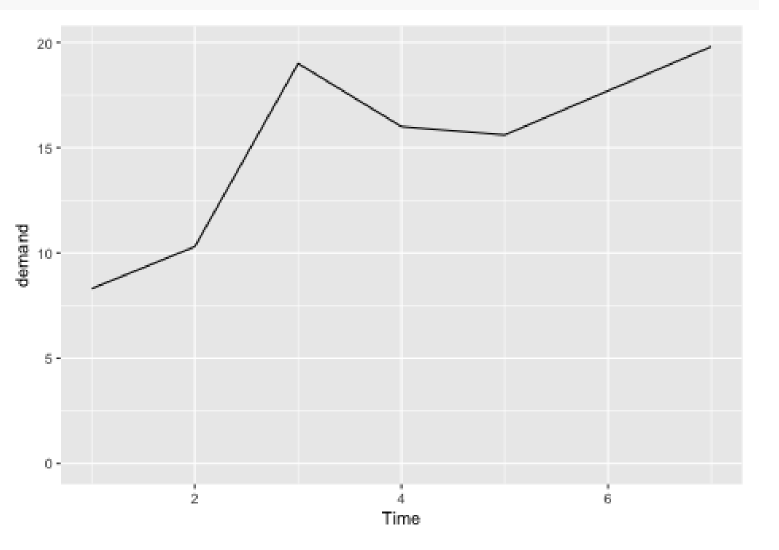
```
qplot(temperature, pressure, data = pressure, geom = c("line", "point"))
```



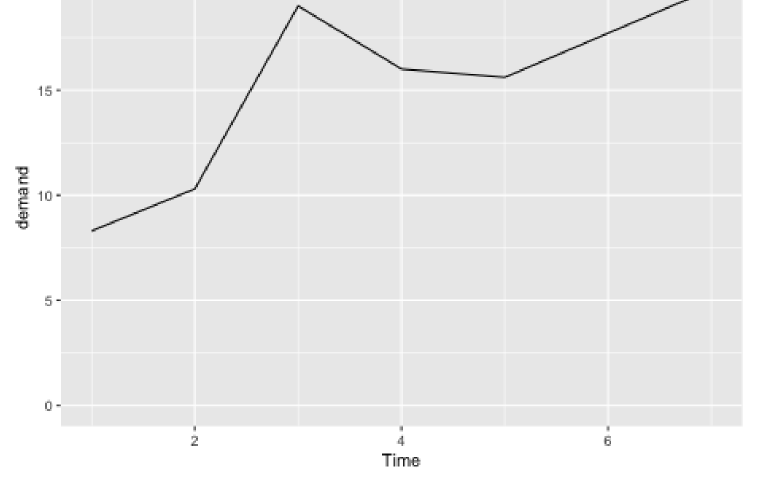




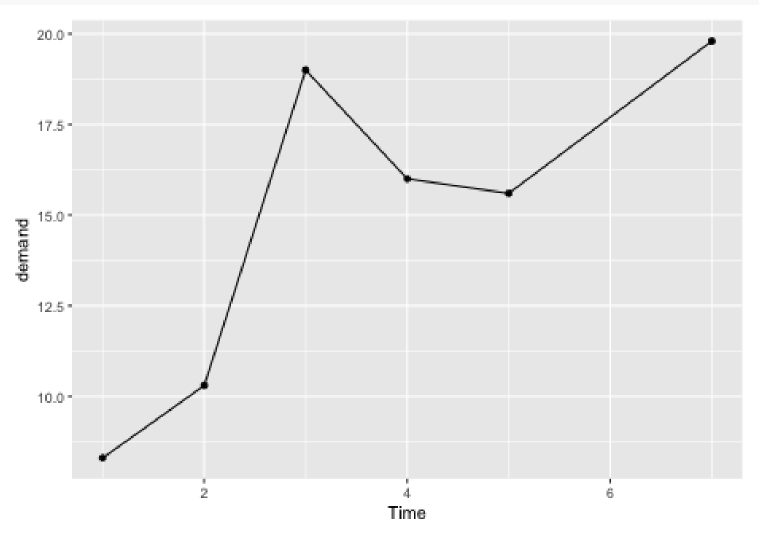
ggplot(BOD, aes(x = Time, y = demand)) + geom_line() + ylim(0, max(BOD\$demand))



```
ggplot(BOD, aes(x = Time, y = demand)) + geom_line() + expand_limits(y = 0)
```



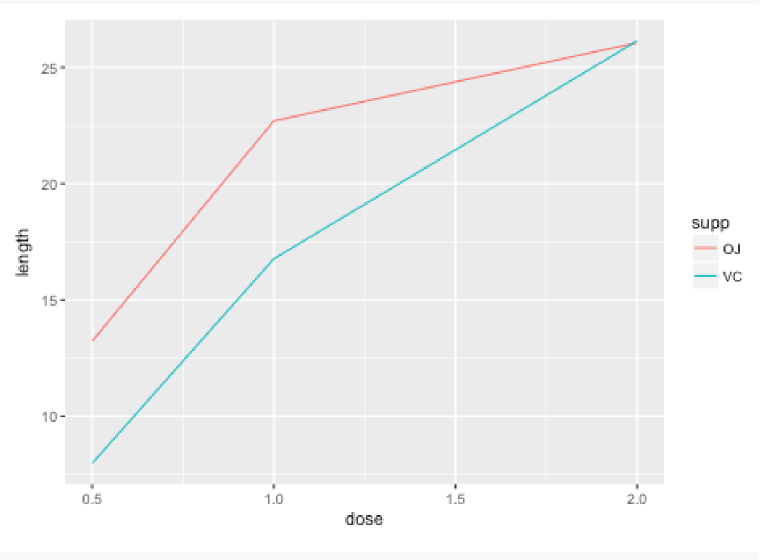




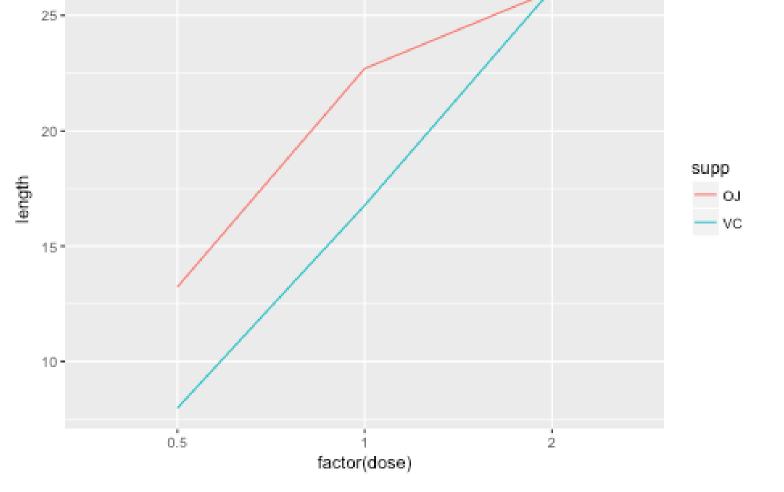
```
# 이산형 변수값에 따른 구분

tg <- ddply(ToothGrowth, c("supp", "dose"), summarize, length = mean(len))
# 색상으로 구분

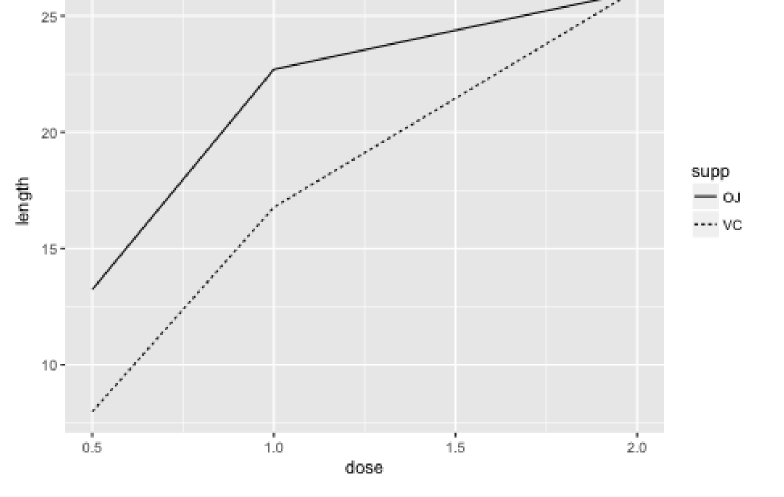
ggplot(tg, aes(x = dose, y = length, color = supp)) + geom_line()
```



```
ggplot(tg, aes(x = factor(dose), y = length, color = supp, group = supp)) + geom_line(
```

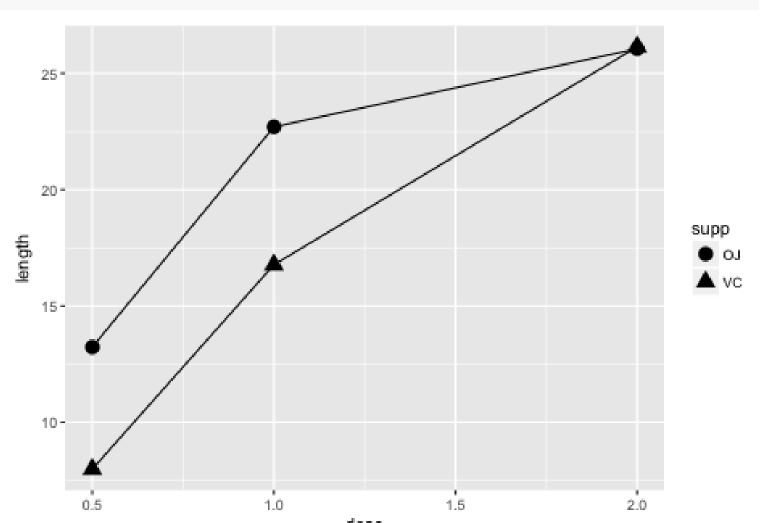


```
# group = supp 주의!:이 명령문이 없으면 데이터를 어떻게 묶어서 그릴지 모름
# Line type으로 구분
ggplot(tg, aes(x = dose, y = length, linetype = supp)) + geom_line()
```



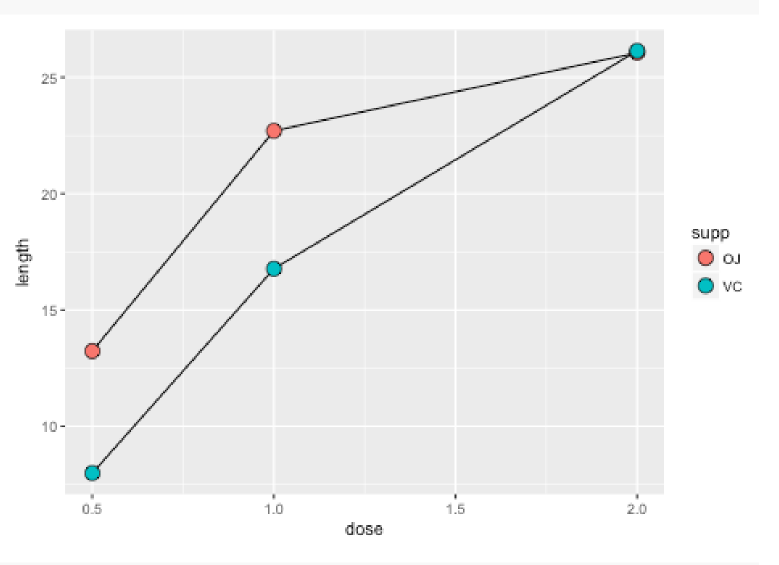
점 형태로 구분

ggplot(tg, aes(x = dose, y = length, shape = supp)) + geom_line() + geom_point(size =



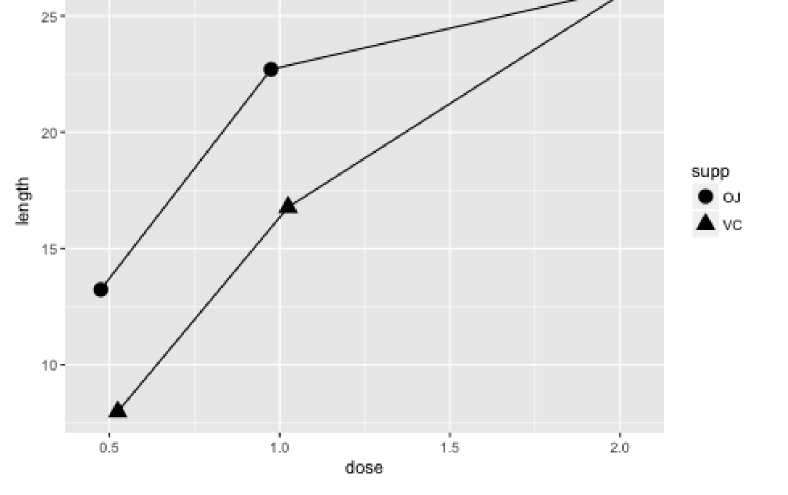
점 색상으로 구분

```
ggplot(tg, aes(x = dose, y = length, fill = supp)) + geom_line() + geom_point(size = 4
```



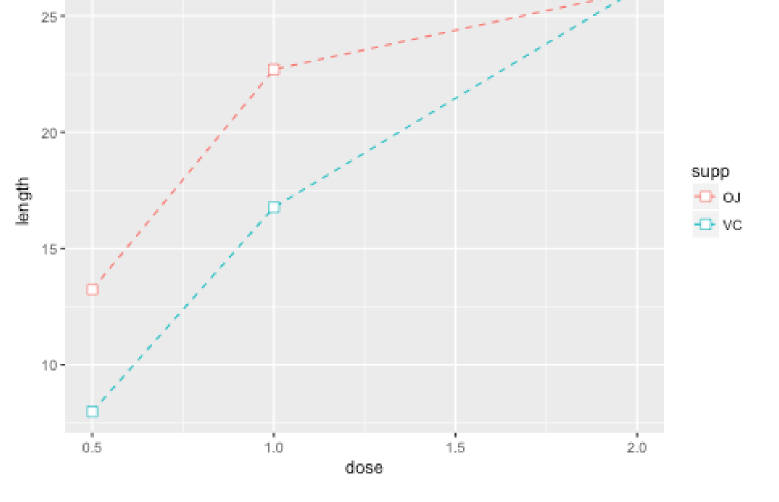
두 선이 겹칠때 하나의 선을 옆으로 이동시켜 표현

```
ggplot(tg, aes(x = dose, y = length, shape = supp)) + geom_line(position = position_do
geom_point(position = position_dodge(0.1), size = 4)
```

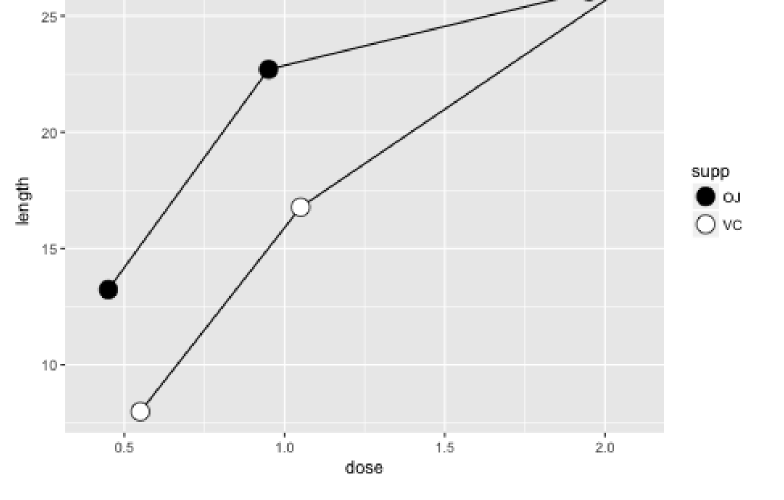


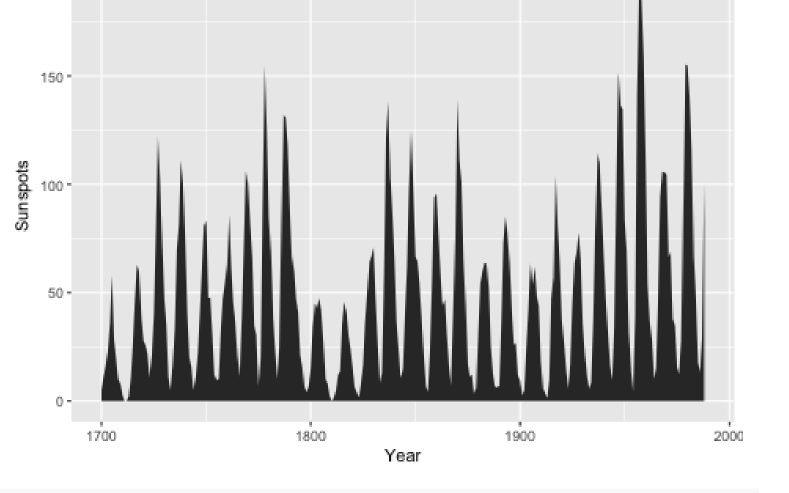
```
# 선형태바꾸기[linetype]

ggplot(tg, aes(x = dose, y = length, color = supp)) + geom_line(linetype = "dashed") +
geom_point(shape = 22, size = 3, fill = "white")
```

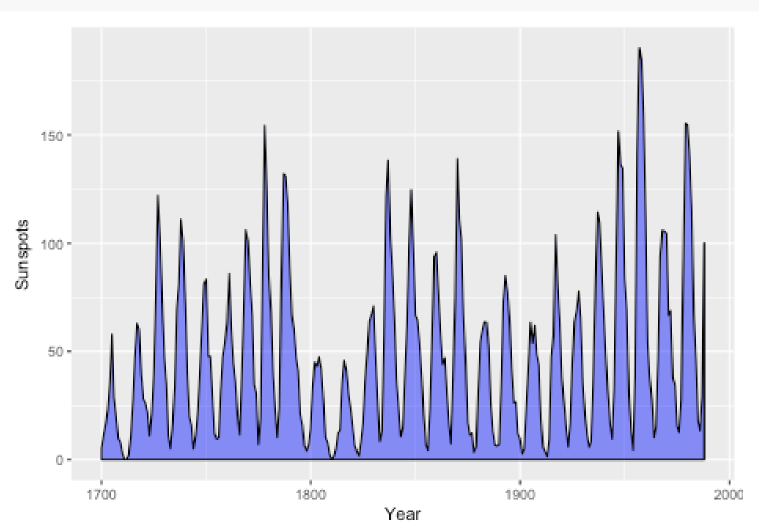


5.4 Part3



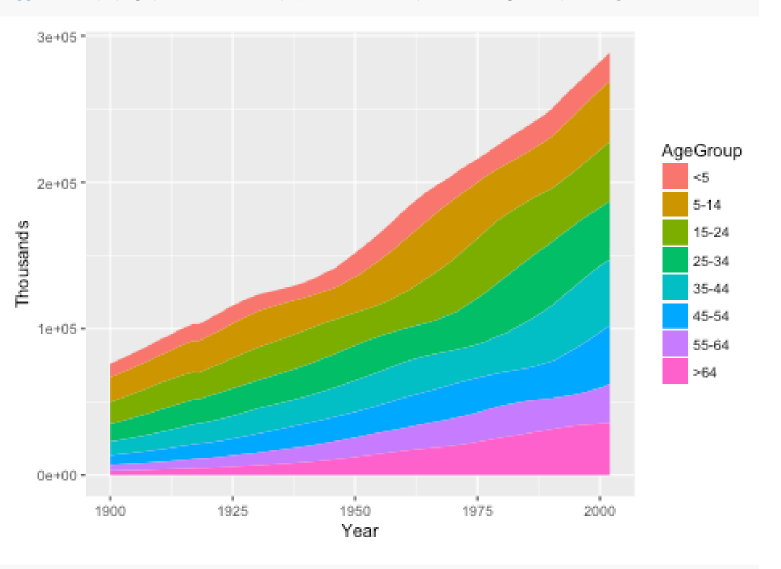






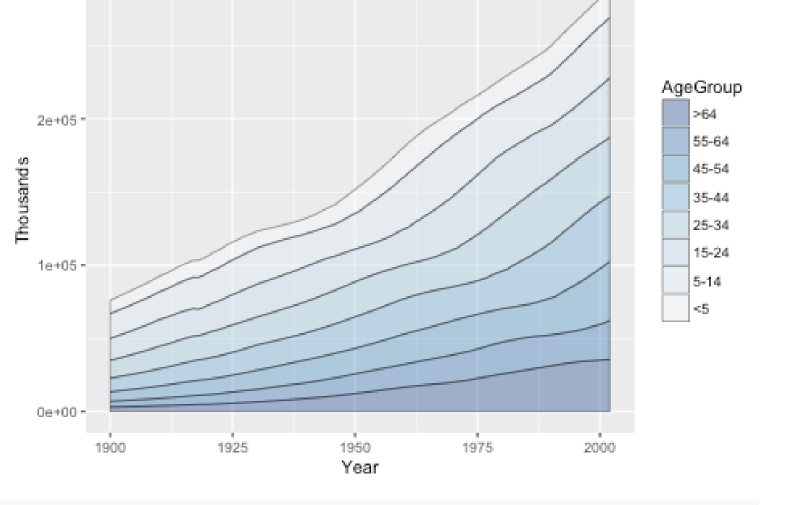
누적 영역 그래프 그리기

```
ggplot(uspopage, aes(x = Year, y = Thousands, fill = AgeGroup)) + geom_area()
```



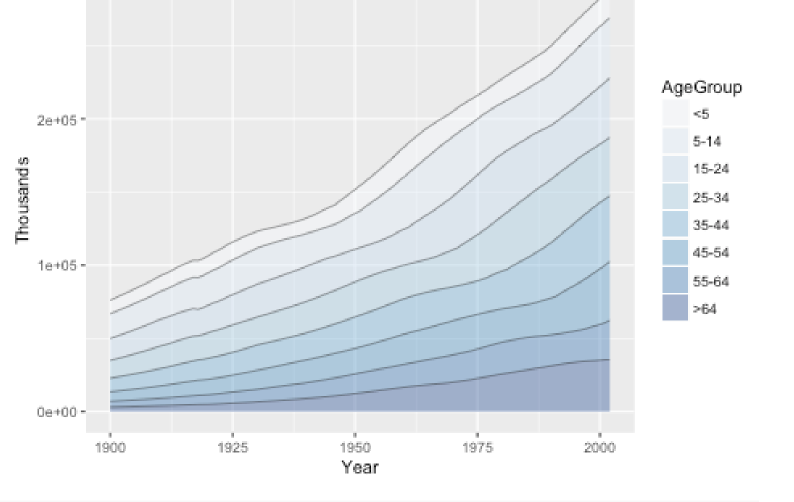
영역색상 그라데이션 넣기

```
ggplot(uspopage, aes(x = Year, y = Thousands, fill = AgeGroup)) + geom_area(color = "b
scale_fill_brewer(palette = "Blues", breaks = rev(levels(uspopage$AgeGroup)))
```



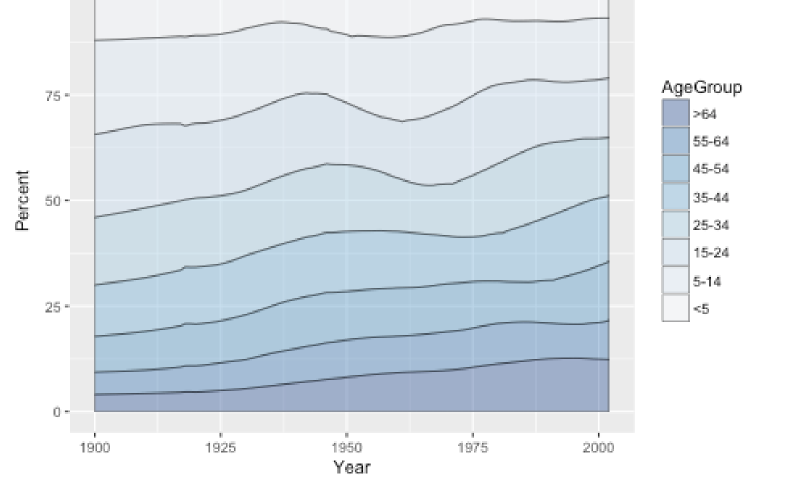
데이터 순서정렬하기 & 양쪽 테두리 지우기

```
ggplot(uspopage, aes(x = Year, y = Thousands, fill = AgeGroup, order = desc(AgeGroup))
geom_area(color = NA, alpha = 0.4) + scale_fill_brewer(palette = "Blues") + geom_lir
```

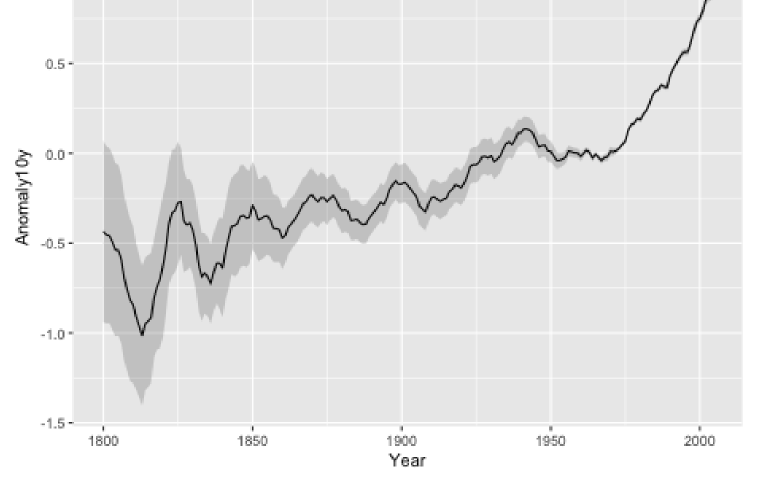


비율 누적 영역 그래프 그리기

```
uspopage_prop <- ddply(uspopage, "Year", transform, Percent = Thousands / sum(Thousand
ggplot(uspopage_prop, aes(x = Year, y = Percent, fill = AgeGroup)) + geom_area(color =
scale_fill_brewer(palette = "Blues", breaks = rev(levels(uspopage$AgeGroup)))</pre>
```

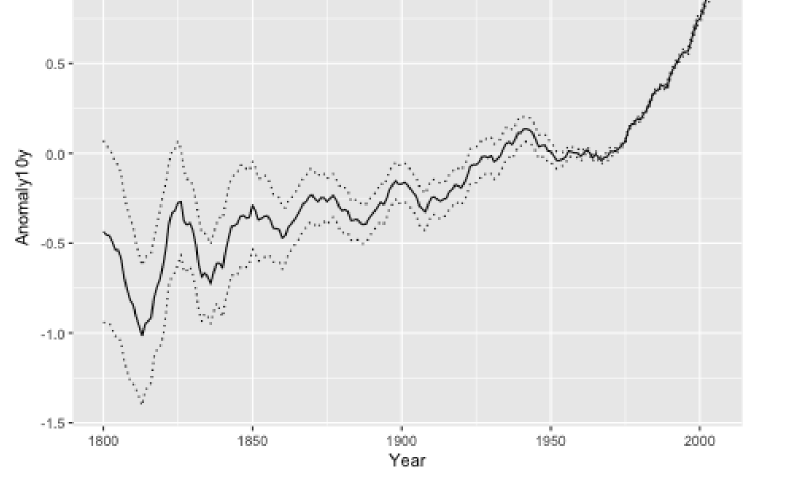


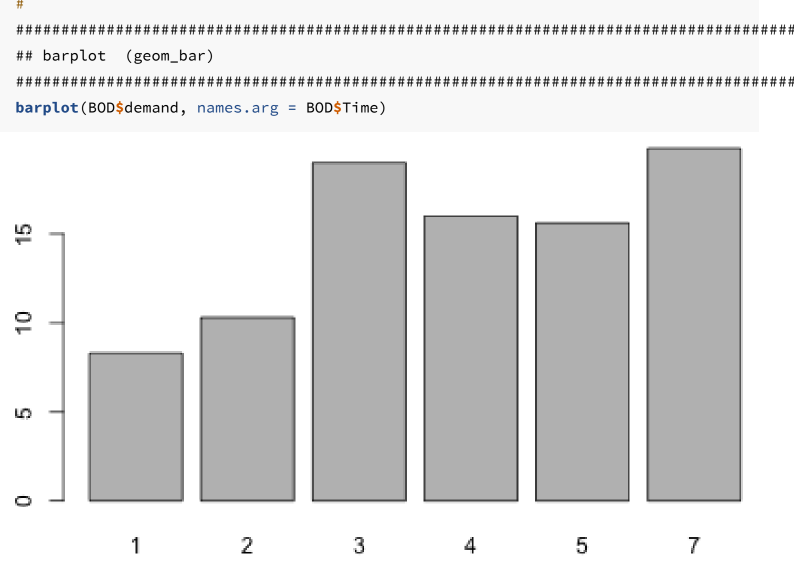
```
# 그래프에 신뢰 영역 추가하기
clim <- subset(climate, Source == "Berkeley", select = c("Year", "Anomaly10y", "Unc10y
# 신뢰영역 음영으로 표현
ggplot(clim, aes(x = Year, y = Anomaly10y)) +
geom_ribbon(aes(ymin = Anomaly10y - Unc10y, ymax = Anomaly10y + Unc10y), alpha = 0.2
geom_line()
```



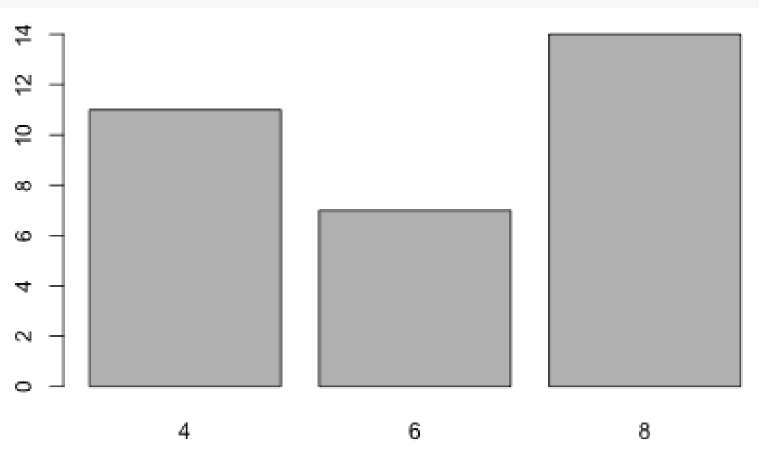
```
# 신뢰영역 점선으로 표현
```

```
ggplot(clim, aes(x = Year, y = Anomaly10y)) +
  geom_line(aes(y = Anomaly10y - Unc10y), linetype = "dotted") +
  geom_line(aes(y = Anomaly10y + Unc10y), linetype = "dotted") +
  geom_line()
```

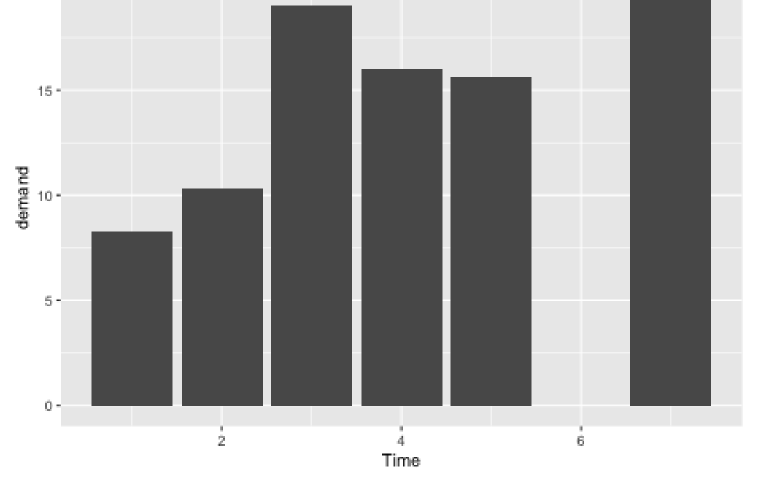




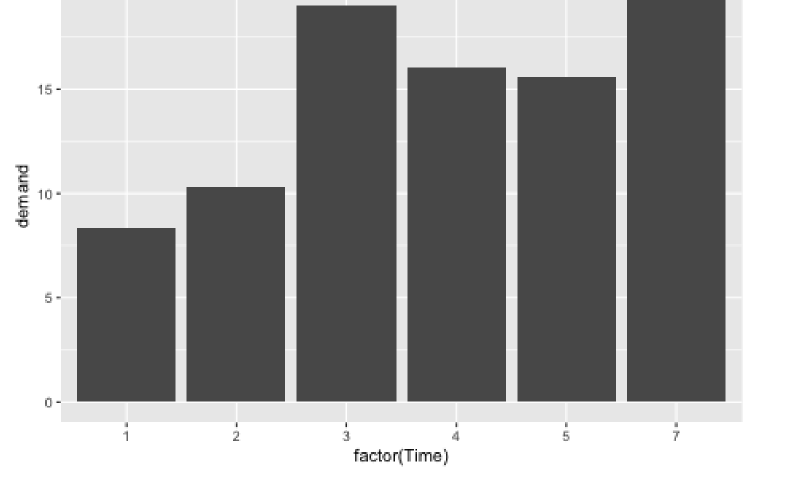
barplot(table(mtcars\$cyl))



```
# x값을 숫자로 인식
# qplot(BOD$Time, BOD$demand, geom = "bar", stat = "identity")
ggplot(BOD, aes(x = Time, y = demand)) + geom_bar(stat = "identity")
```

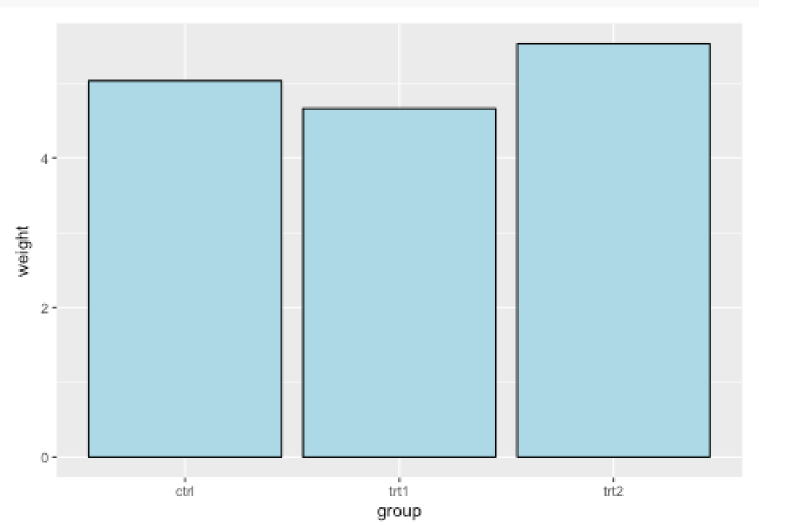


```
# x값을 요인으로 변환
# qplot(as.factor(BOD$Time), BOD$demand, geom = "bar", stat = "identity")
ggplot(BOD, aes(x = factor(Time), y = demand)) + geom_bar(stat = "identity")
```



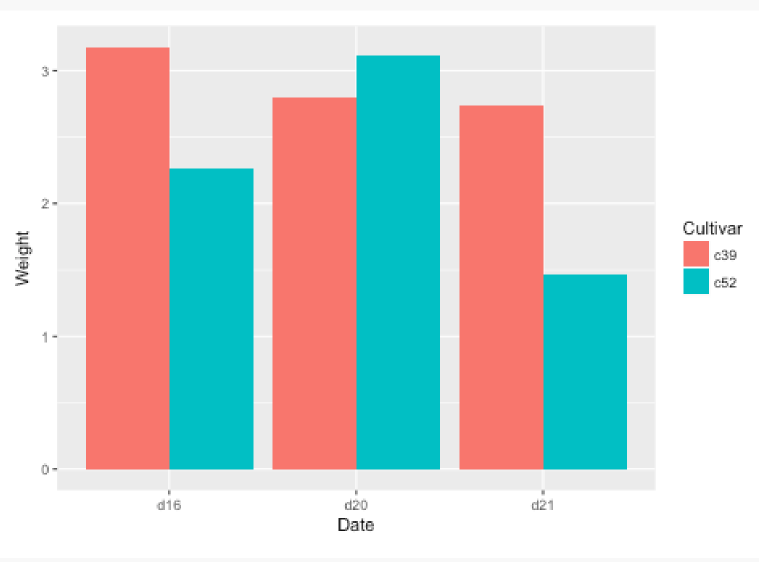
```
# 막대 색상 채우기/테두리 설정하기 (fill : 채우기/ colour(or color) : 테두리)

ggplot(pg_mean, aes(x = group, y = weight)) + geom_bar(stat = "identity", fill = "light")
```



막대 묶어서 표현하기(나누어 표현하고 싶은 변수를 색상으로 지정)

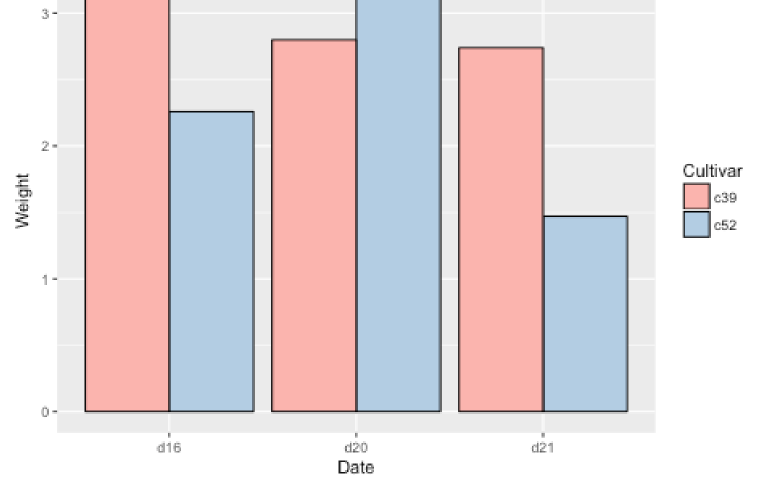
ggplot(cabbage_exp, aes(x = Date, y = Weight, fill = Cultivar)) + geom_bar(stat = "ide



```
# dodge : "피하다"라는 의미로 막대를 새로운 값을 기준으로 나누어 표현

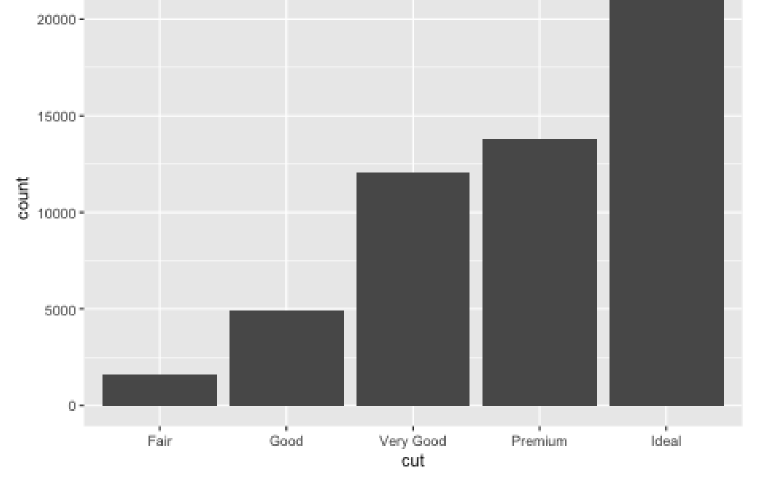
ggplot(cabbage_exp, aes(x = Date, y = Weight, fill = Cultivar)) + geom_bar(stat = "ide

scale_fill_brewer(palette = "Pastel1")
```



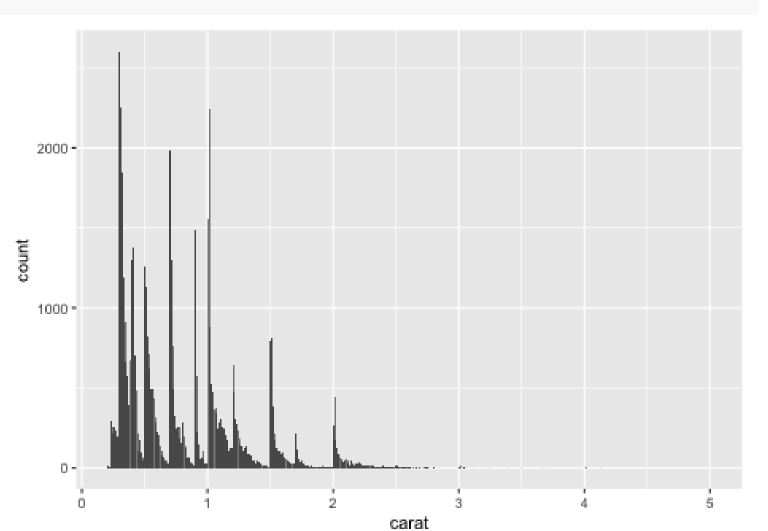
5.5 Part4

```
# 빈도수 막대 그래프 그리기
# x가 이산형
ggplot(diamonds, aes(x = cut)) + geom_bar()
```

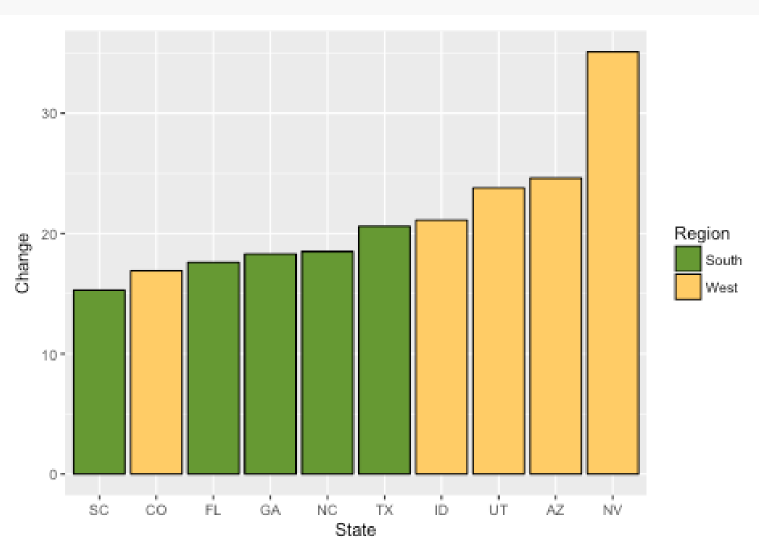


x가 연속형

ggplot(diamonds, aes(x = carat)) + geom_bar()



막대 색상 넣기/ 축 이름 정의하기(reorder) upc <- subset(uspopchange, rank(Change) > 40) ggplot(upc, aes(x = reorder(Abb, Change), y = Change, fill = Region)) + geom_bar(stat scale_fill_manual(values = c("#669933", "#FFCC66")) + xlab("State")

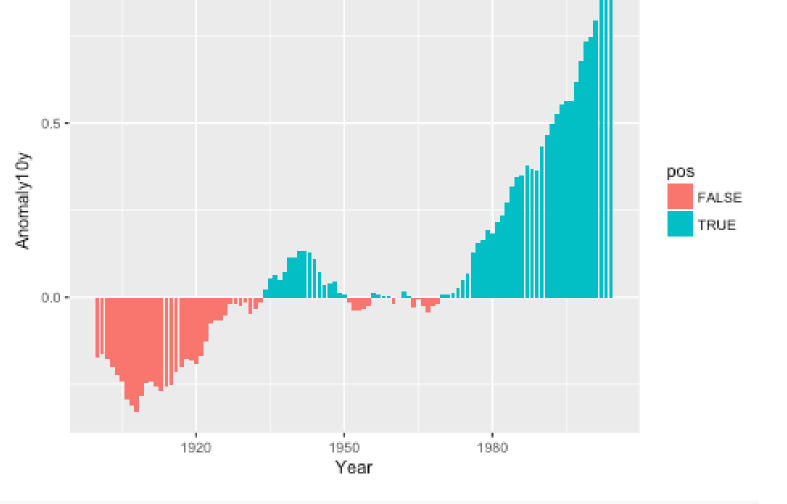


```
# 양수/음수 다른 색상으로 표현 (구분 inde를 만들어 색상 변수로 지정)

csub <- subset(climate, Source == "Berkeley" & Year >= 1900 )

csub$pos <- csub$Anomaly10y >= 0

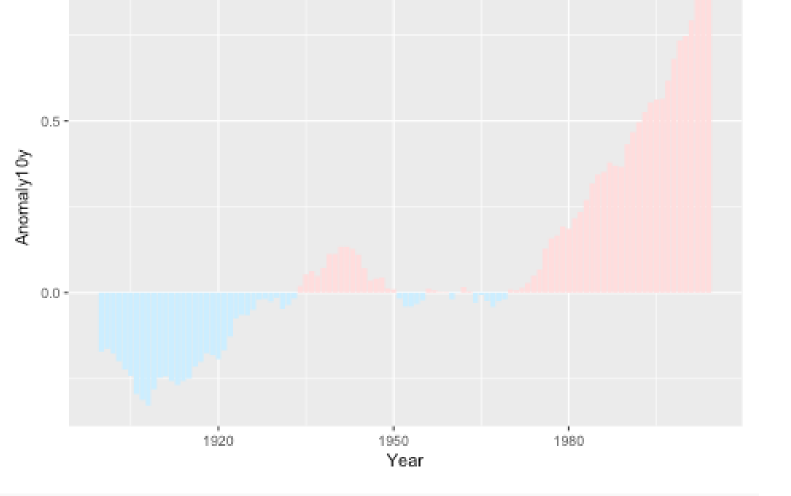
ggplot(csub, aes(x = Year, y = Anomaly10y, fill = pos)) + geom_bar(stat = "identity",
```



```
# 막대 그래프 테두리 두께 설정(size) / 범례 지우기(guide = FALSE)

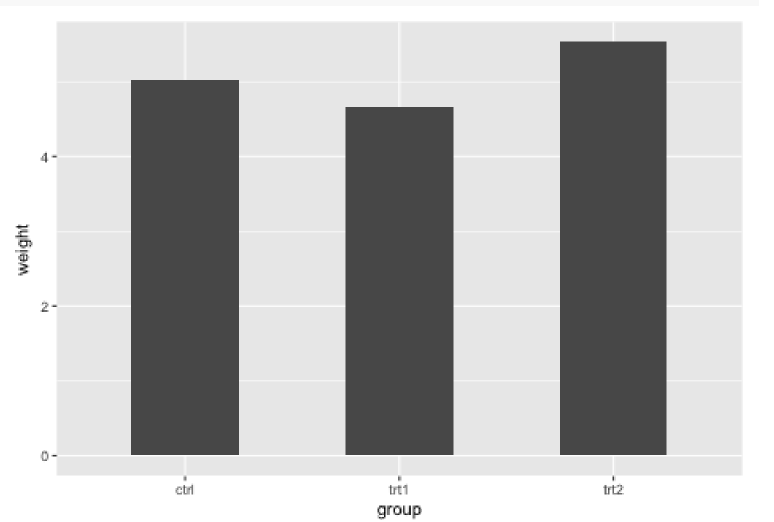
ggplot(csub, aes(x = Year, y = Anomaly10y, fill = pos)) + geom_bar(stat = "identity",

scale_fill_manual(values = c("#CCEEFF", "#FFDDDD"), guide = FALSE)
```



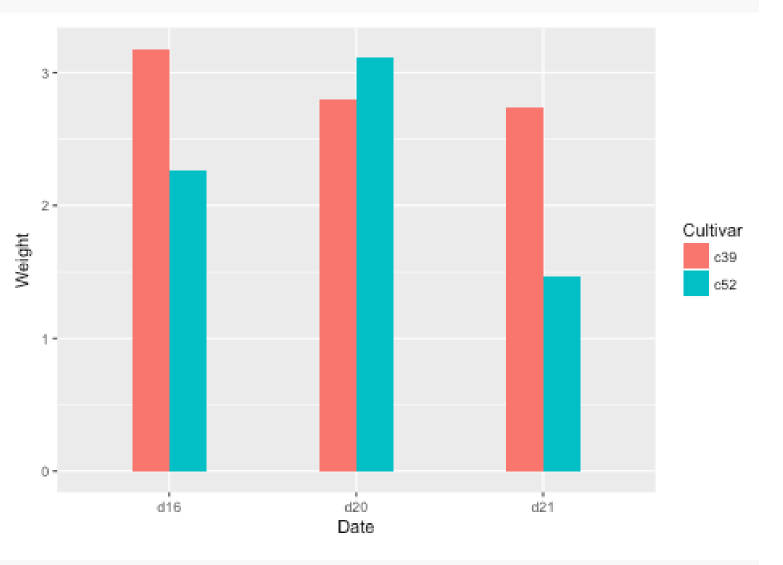
```
# 막대 너비/ 간격 조절하기(width : 최대 너비는 1)

ggplot(pg_mean, aes(x = group, y = weight)) + geom_bar(stat = "identity", width = 0.5)
```



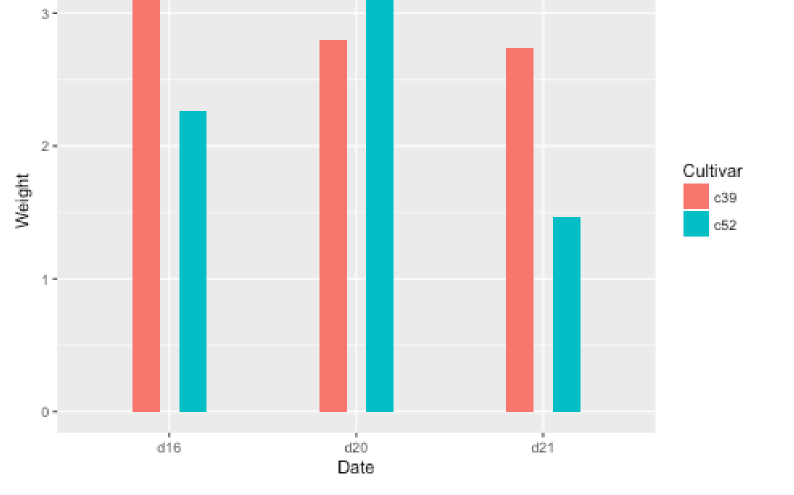
```
# 막대 그룹 간의 간격 조절하기 (default: 0.9)
```

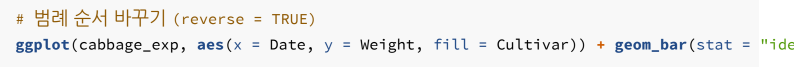
```
ggplot(cabbage_exp, aes(x = Date, y = Weight, fill = Cultivar)) + geom_bar(stat = "ide
```

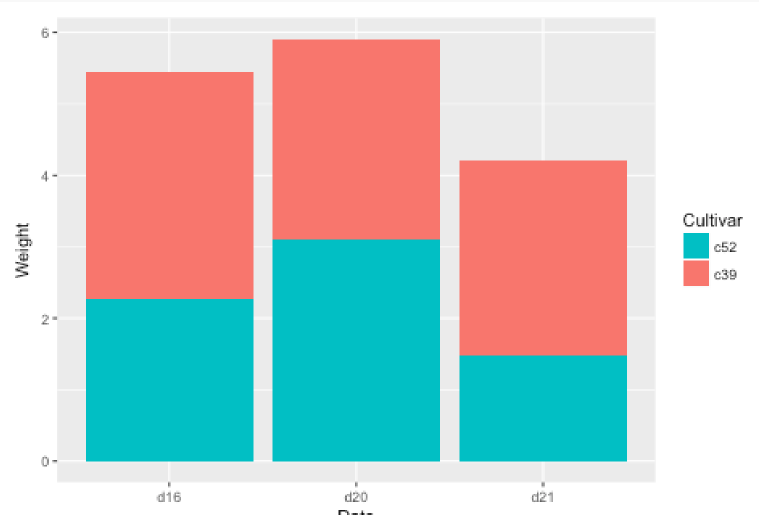


```
# 막대 그룹 내부간의 간격 조절 (default : 0.9)
```

ggplot(cabbage_exp, aes(x = Date, y = Weight, fill = Cultivar)) + geom_bar(stat = "ide

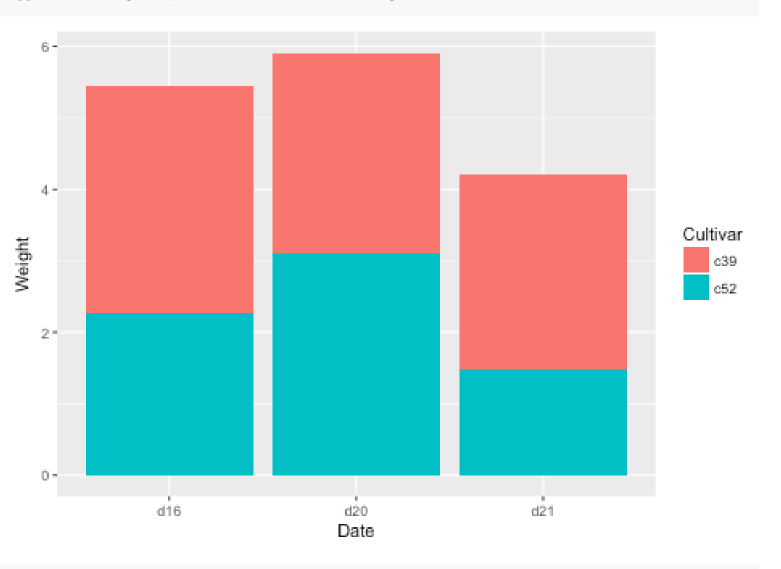






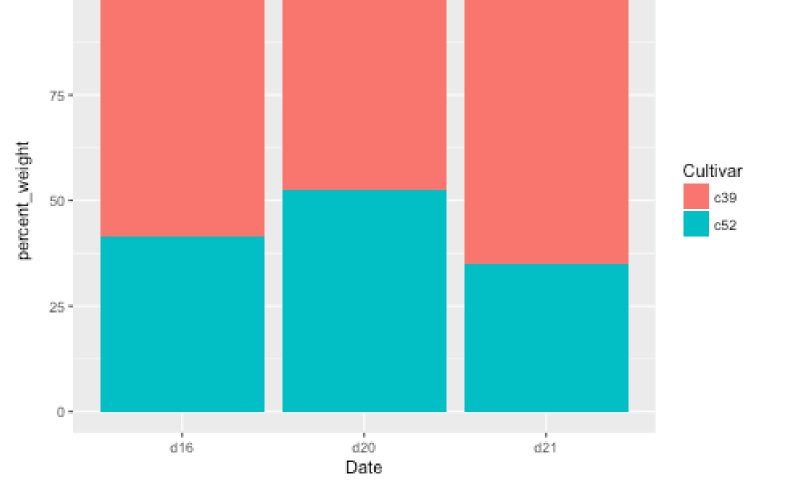
막대 쌓는 순서 바꾸기

ggplot(cabbage_exp, aes(x = Date, y = Weight, fill = Cultivar, order = desc(Cultivar))

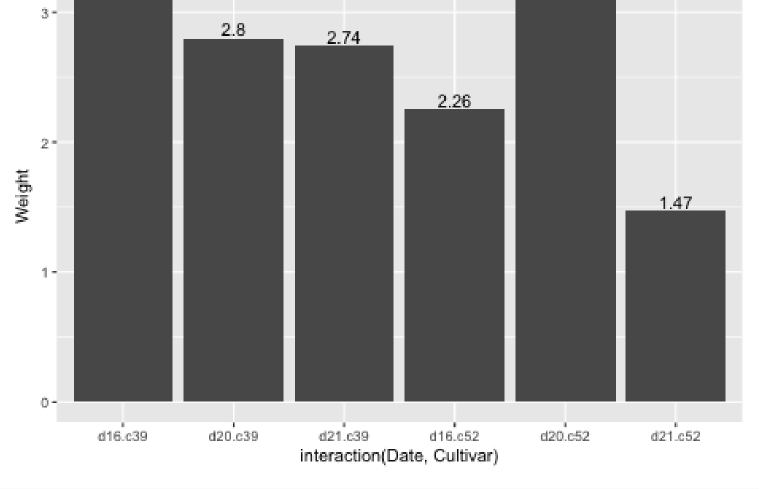


비율 누적 막대 그래프 그리기(막대 전체가 100%가 되도록)

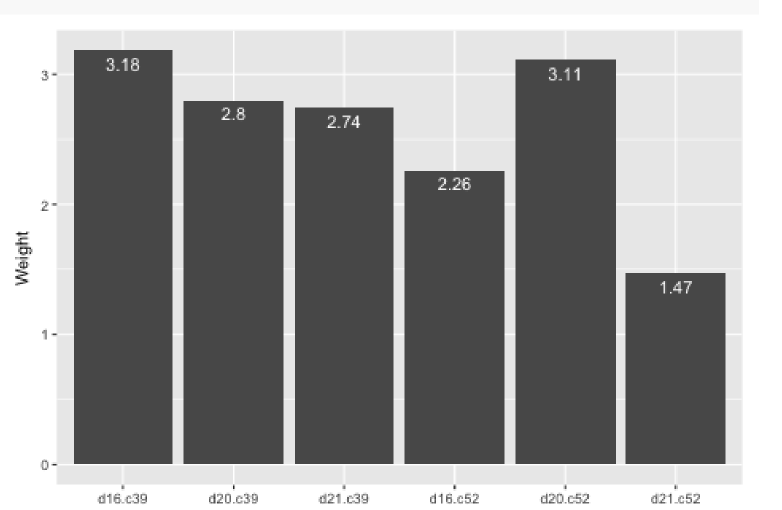
```
ce <- ddply(cabbage_exp, "Date", transform, percent_weight = Weight / sum(Weight) * 16
ggplot(ce, aes(x = Date, y = percent_weight, fill = Cultivar)) + geom_bar(stat = "ider</pre>
```



```
# 막대에 라벨 추가하기 [ geom_text ]
# 선 상단 [ vjust < 0 ]
ggplot(cabbage_exp, aes(x = interaction(Date, Cultivar), y = Weight)) + geom_bar(stat geom_text(aes(label = Weight), vjust = -0.2)
```

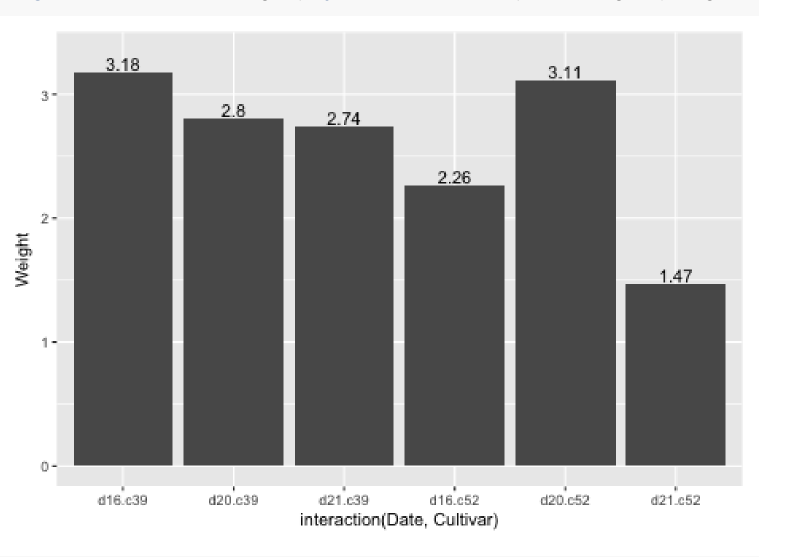


선하단[vjust > 0] ggplot(cabbage_exp, aes(x = interaction(Date, Cultivar), y = Weight)) + geom_bar(stat geom_text(aes(label = Weight), vjust = 1.5, color = "white")



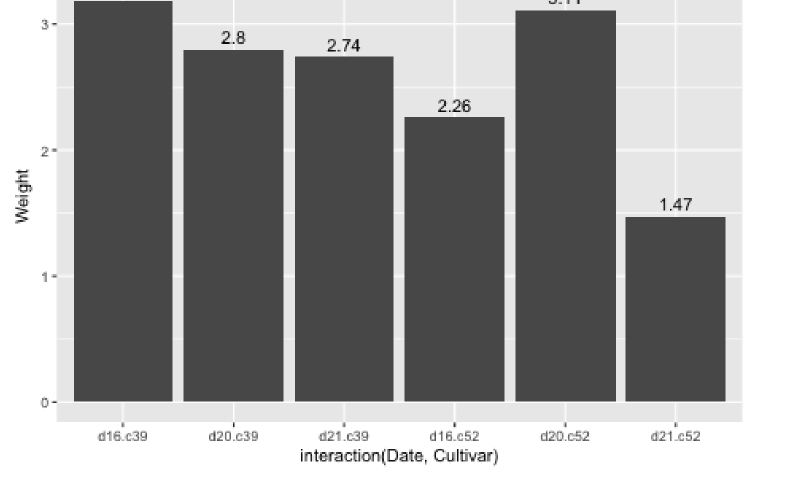
```
# 그래프 범위 설정
# max(데이터)로 지정
```

```
ggplot(cabbage_exp, aes(x = interaction(Date, Cultivar), y = Weight)) + geom_bar(stat
geom_text(aes(label = Weight), vjust = -0.2) + ylim(0, max(cabbage_exp$Weight) * 1.6
```



막대의 상단보다 조금 높은 위치로 지정(데이터에 따라 자동 조정)

```
ggplot(cabbage_exp, aes(x = interaction(Date, Cultivar), y = Weight)) + geom_bar(stat
geom_text(aes(y = Weight + 0.1, label = Weight))
```



```
# 누적 합계 그래프 그리기

ce <- arrange(cabbage_exp, Date, Cultivar)

ce <- ddply(ce, "Date", transform, label = cumsum(Weight))

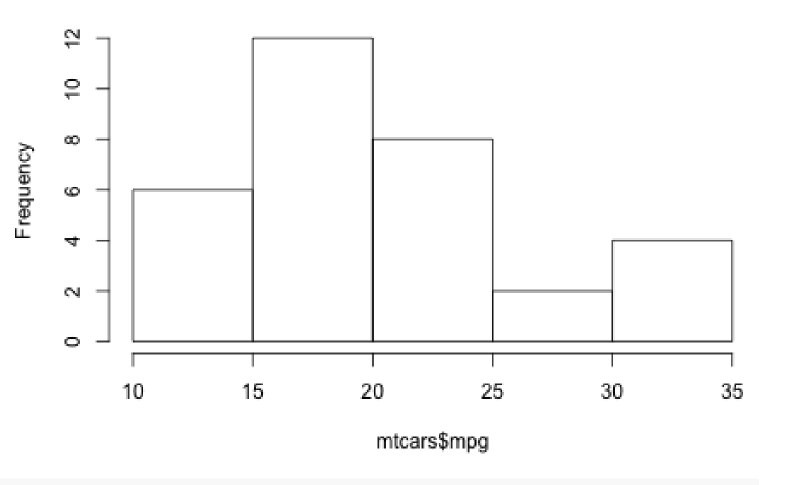
# ddply(data.frame, 그룹 기준 마지막 변수)

ggplot(ce, aes(x = Date, y = Weight, fill = Cultivar)) + geom_bar(stat = "identity", com_text(aes(y = label, label = paste(format(Weight, nsmall=2), "kg")), size = 6, worders(fill = guide_legend(reverse = TRUE)) + scale_fill_brewer(palette = "Pastel1")
```



histogram (geom_histogram)

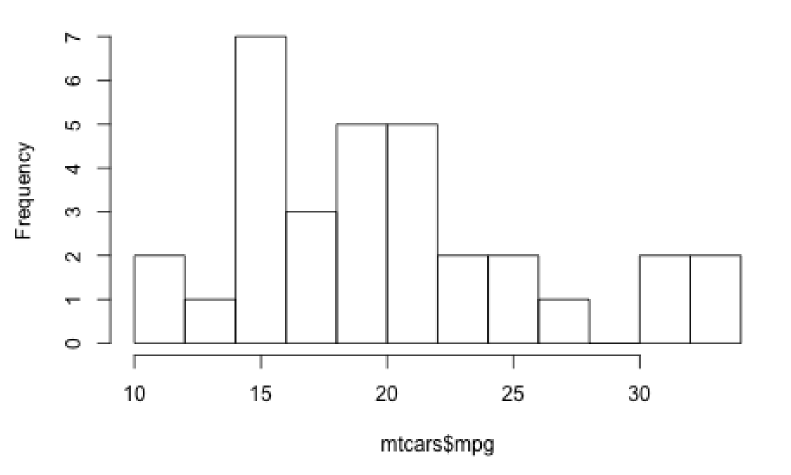
hist(mtcars\$mpg)



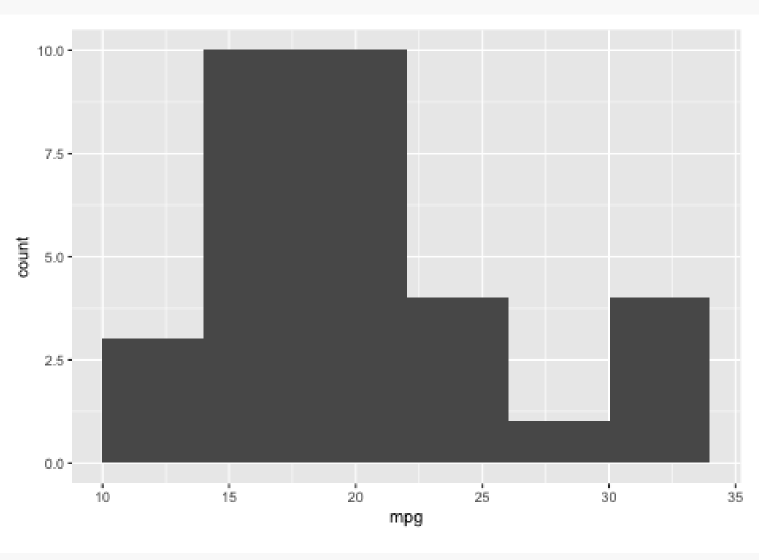
상자(bin) 개수 지정

hist(mtcars\$mpg, breaks = 10)

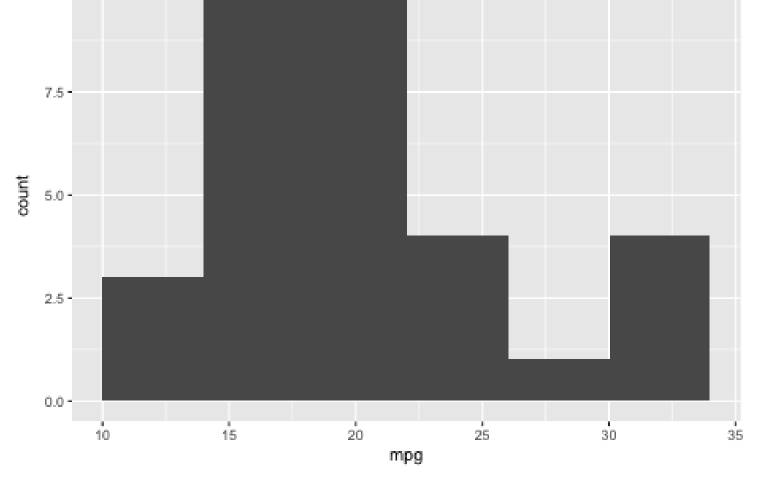
Histogram of mtcars\$mpg



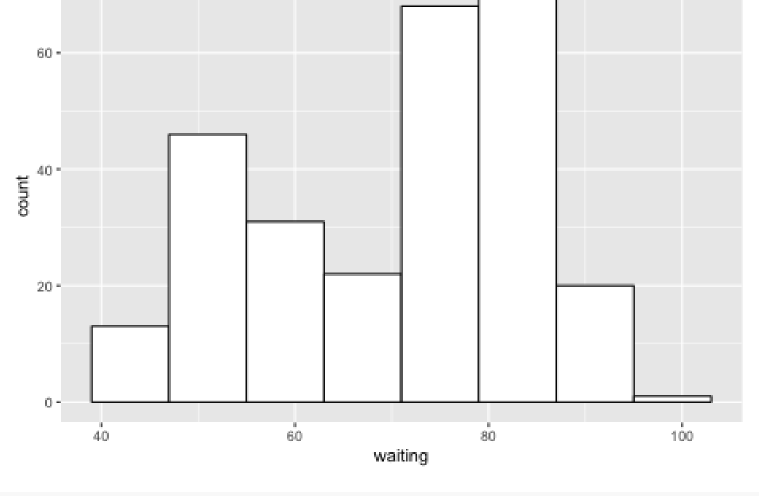
```
qplot(mpg, data = mtcars, binwidth = 4)
```



```
# ggplot(mtcars, aes(x = mpg)) + geom_histogram()
ggplot(mtcars, aes(x = mpg)) + geom_histogram(binwidth = 4)
```



```
h <- ggplot(faithful, aes(x = waiting))
h + geom_histogram(binwidth = 8, fill = "white", color = "black", origin = 31)
## Warning: `origin` is deprecated. Please use `boundary`
## instead.</pre>
```



```
# 그룹화된 데이터로 여러개의 히스토그램 만들기
library(MASS)

##

## Attaching package: 'MASS'

## The following object is masked from 'package:dplyr':

##

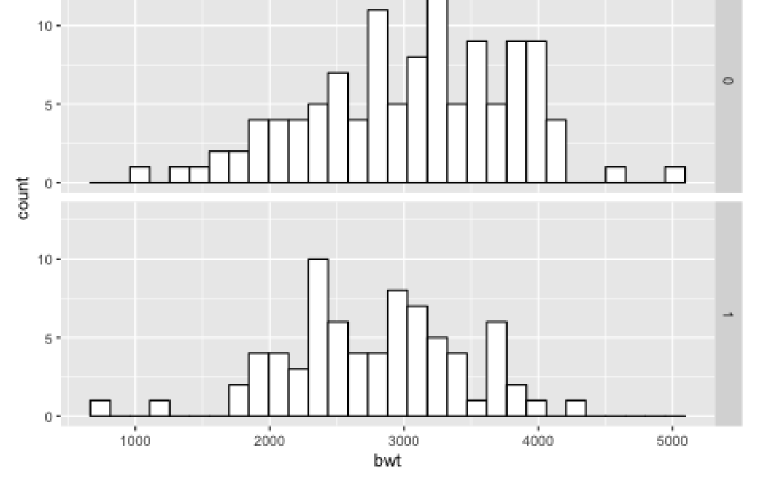
## select

ggplot(birthwt, aes(x = bwt)) + geom_histogram(fill = "white", color = "black") + face

## `stat_bin()` using `bins = 30`. Pick better value
```

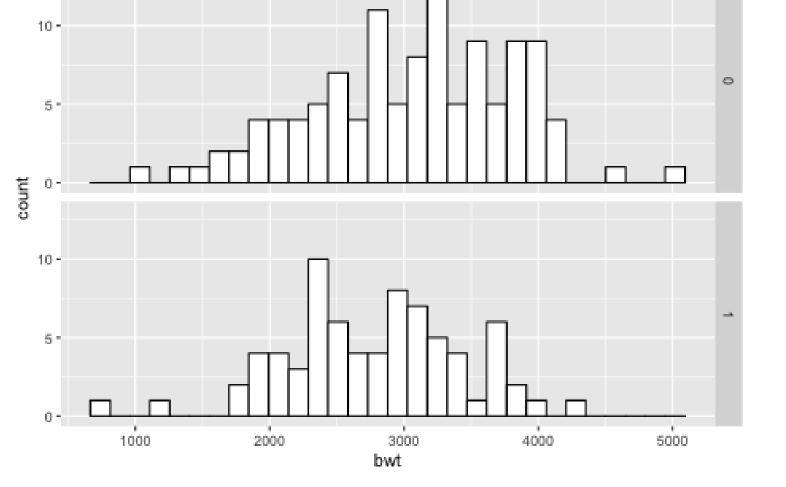
Warning: `origin` is deprecated. Please use `boundary` instead.

with `binwidth`.



```
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.

ggplot(birthwt, aes(x = bwt)) + geom_histogram(fill = "white", color = "black") + face
## `stat_bin()` using `bins = 30`. Pick better value
## with `binwidth`.
```



```
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.

# 그룹 라벨 정의
birthwt1 <- birthwt
birthwt1$smoke <- factor(birthwt1$smoke)

levels(birthwt1$smoke)

## [1] "0" "1"

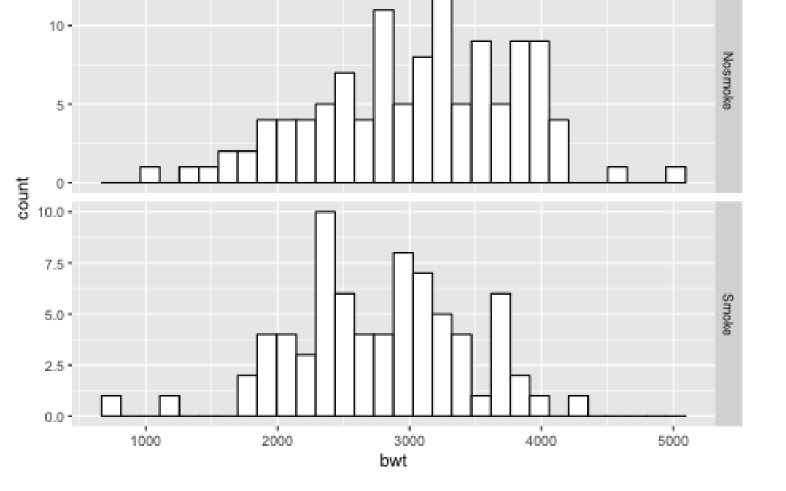
## [1] "0" "1"

birthwt$smoke <- revalue(birthwt1$smoke, c("0" = "Nosmoke", "1" = "Smoke"))

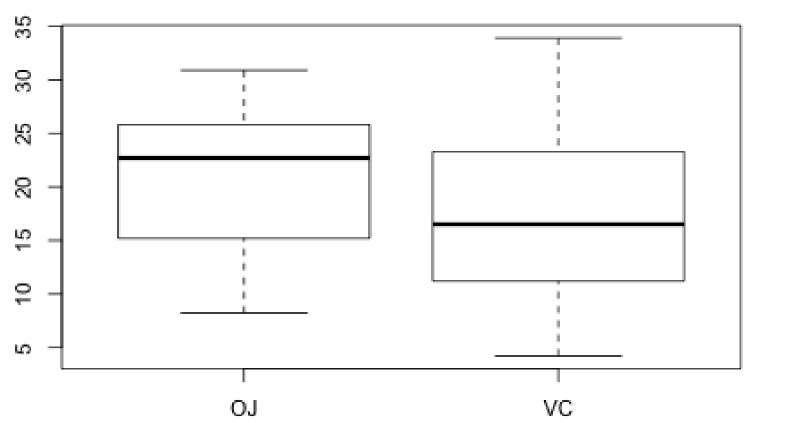
ggplot(birthwt, aes(x = bwt)) + geom_histogram(fill = "white", color = "black") + face

## `stat_bin()` using `bins = 30`. Pick better value
```

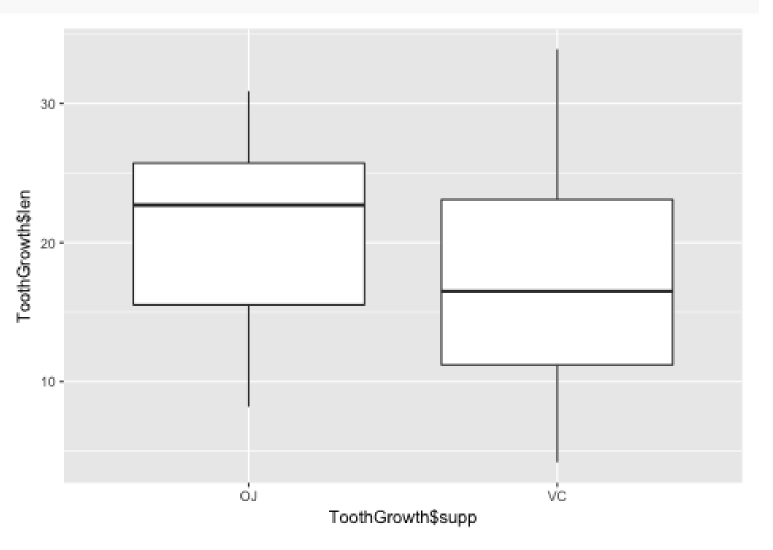
with `binwidth`.



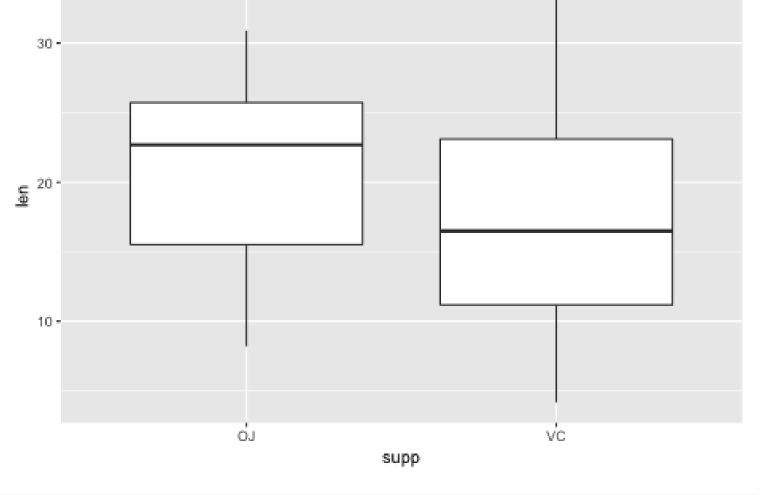
`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



```
qplot(ToothGrowth$supp, ToothGrowth$len, geom = "boxplot")
```



```
ggplot(ToothGrowth, aes(x = supp, y = len))+ geom_boxplot()
```



두 변수의 상호작용을 x축으로

OJ.0.5

VC.0.5

8

15

LO

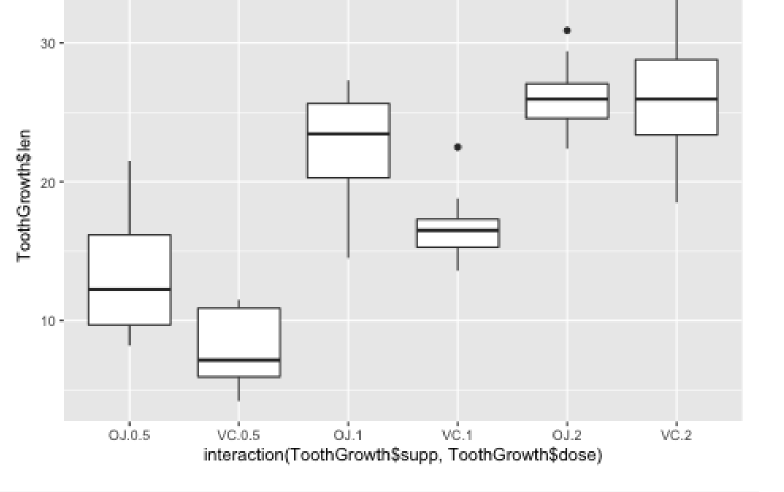
boxplot(len ~ supp + dose, data = ToothGrowth)

VC.1

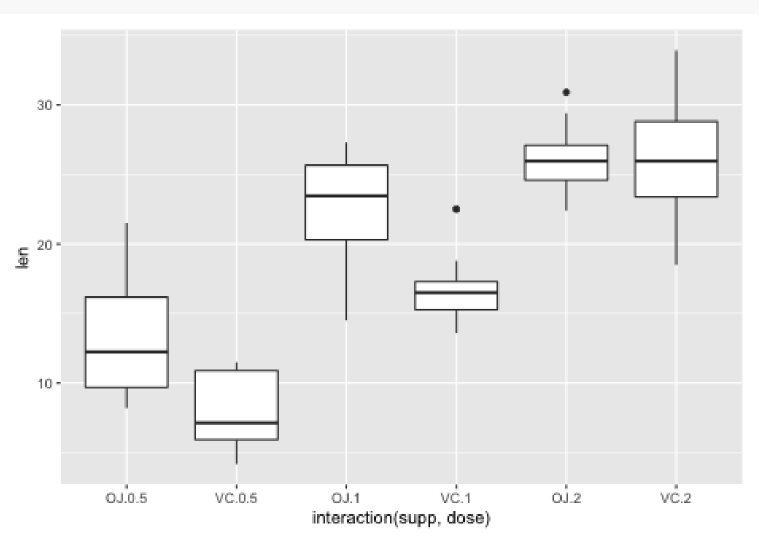
OJ.2

VC.2

OJ.1





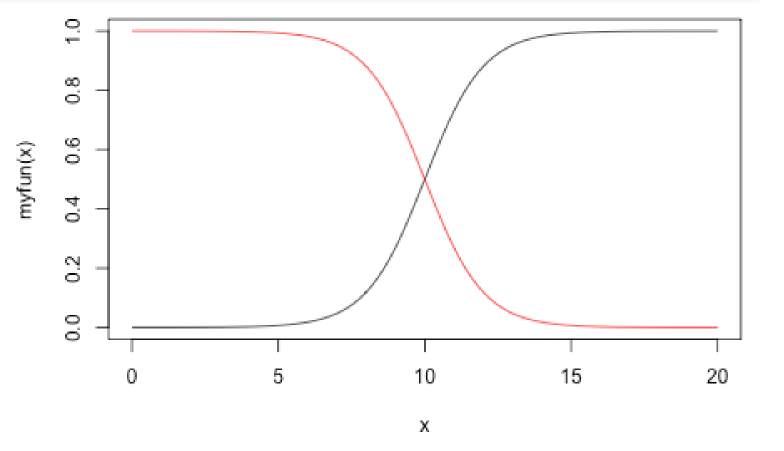


```
## 함수곡선 그리기 (stat_function)
```

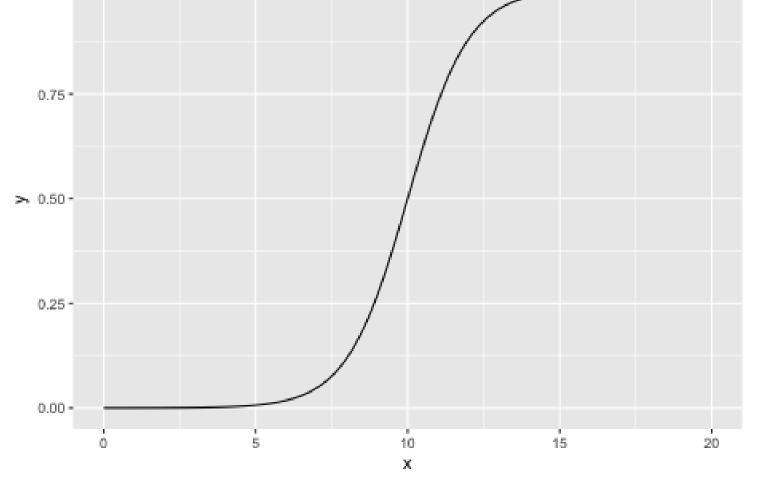
사용자함수 정의

```
myfun <- function(xvar){
   1/(1 + exp(-xvar + 10))
}

curve(myfun(x), from = 0, to = 20)
curve(1-myfun(x), add = TRUE, col = "red")</pre>
```



```
# qplot(c(0,20), fun = myfun, stat = "function", geom = "line")
ggplot(data.frame(x = c(0,20)), aes(x =x)) + stat_function(fun = myfun, geom = "line")
```



.1 Assignment 1

첨부한 concUnitConv-test.R과 유사한 R script를 실행하였을 때, concUnitConv-test.Rout 과 유사한 결과나 나오는 concUnitConv.R 파일을 작성하시오.

- 제출기한: 2017-05-10 18:00
- 제출방법: R scirpt와 output을 k@acr.kr¹, shan@acp.kr², sec@acp.kr³ 로 제출

.1.1 concUnitConv-test.R

```
concUnitConv() # Wrong input
concUnitConv("kg/L", "g/L") # Wrong input
concUnitConv("kg/L", "g/L") # Wrong input
concUnitConv("mg/kL", "g/L") # Wrong input

concUnitConv("mg/L", "ug/mL")
Theoph$conc * concUnitConv("mg/L", "ug/L")
Theoph$conc * concUnitConv("mg/L", "mg/mL")
Theoph$conc * concUnitConv("mg/L", "mmol/L") # Wrong input
Theoph$conc * concUnitConv("mg/L", "mmol/L", MW=-100) # Wrong input
Theoph$conc * concUnitConv("mg/L", "mmM", MW=180.164) # Wrong input
Theoph$conc * concUnitConv("mg/L", "mmol/L", MW=180.164)
Theoph$mM = Theoph$conc * concUnitConv("mg/L", "mmol/L", MW=180.164)
Theoph$mM * concUnitConv("mmol/L", "ug/L", MW=180.164)
Theoph$mM * concUnitConv("mmol/L", "ug/L", MW=180.164)
```

.1.2 concUnitConv-test.Rout

```
> source("D:/G/Desk/R/concUnitConv.R")
>
concUnitConv() # Wrong input
Error in concUnitConv(): Source concentration unit is not valid.
> concUnitConv("kg/L", "g/L") # Wrong input
```

¹mailto:k@acr.kr

²mailto:shan@acp.kr

```
The concentration ( Rg/L , g/L ) • Source amount is not supported.
> concUnitConv("g/kL", "g/L") # Wrong input
Error in concUnitConv("g/kL", "g/L") : Volume unit is not supported.
> concUnitConv("mg/L", "ug/mL")
1
> Theoph$conc * concUnitConv("mg/L", "ug/L")
       740
            2840 6570 10500
                              9660 8580
                                          8360 7470
                                                       6890
                                                             5940
                                                                   3280
                                                                               1720
                                                                                     7
  [1]
[28]
      7500
            6200 5300
                        4900
                              3700 1050
                                              0
                                                1890
                                                      4600
                                                             8600
                                                                   8380
                                                                         7540
                                                                               6880
[55]
      1570
               0 1290
                        3080
                              6440 6320
                                           5530
                                                 4940
                                                      4020
                                                             3460
                                                                   2780
                                                                         920
                                                                                150
[82]
            6590 5880
                        4730 4570 3000
                                                      7370
     7560
                                          1250
                                                             9030
                                                                   7140
                                                                         6330
                                                                               5660
[109]
      5680 2420
                        4860 7240 8000 6810
                                                 5870
                                                                                860
                      0
                                                      5220
                                                             4450
                                                                   3620
                                                                         2690
> Theoph$conc * concUnitConv("mg/L", "mg/mL")
  [1] 0.00074 0.00284 0.00657 0.01050 0.00966 0.00858 0.00836 0.00747 0.00689 0.00594
 [21] 0.00301 0.00090 0.00000 0.00440 0.00690 0.00820 0.00780 0.00750 0.00620 0.00530
[41] 0.00578 0.00533 0.00419 0.00115 0.00000 0.00202 0.00563 0.01140 0.00933 0.00874
[61] 0.00553 0.00494 0.00402 0.00346 0.00278 0.00092 0.00015 0.00085 0.00235 0.00502
[81] 0.00731 0.00756 0.00659 0.00588 0.00473 0.00457 0.00300 0.00125 0.00000 0.00737
[101] 0.00289 0.00522 0.00641 0.00783 0.01021 0.00918 0.00802 0.00714 0.00568 0.00242
[121] 0.00086 0.00000 0.00125 0.00396 0.00782 0.00972 0.00975 0.00857 0.00659 0.00611
> Theoph$conc * concUnitConv("mg/L", "mmol/L") # Wrong input
Error in concUnitConv("mg/L", "mmol/L") :
  Positive molecular weight should be given.
> Theoph$conc * concUnitConv("mg/L", "mmol/L", MW=-100) # Wrong input
Error in concUnitConv("mg/L", "mmol/L", MW = -100) :
  Positive molecular weight should be given.
> Theoph$conc * concUnitConv("mg/L", "mM", MW=180.164) # Wrong input
Error in concUnitConv("mg/L", "mM", MW = 180.164) :
 Target concentration unit is not valid.
Theoph$conc * concUnitConv("mg/L", "mmol/L", MW=180.164)
  [1] 0.0041073688 0.0157634156 0.0364667747 0.0582802336 0.0536178149 0.0476232766 0.
[13] 0.0095468573 0.0439044426 0.0461246420 0.0462356520 0.0380209143 0.0337470305 0.
 [25] 0.0382984392 0.0455140872 0.0432938878 0.0416287383 0.0344130903 0.0294176417 0.
[37] 0.0477342865 0.0465131769 0.0418507582 0.0381874292 0.0320818810 0.0295841567 0.
 [49] 0.0517861504 0.0485113563 0.0419617682 0.0393530339 0.0327479408 0.0242556782 0.
 [61] 0.0306942563 0.0274194623 0.0223130037 0.0192047246 0.0154303856 0.0051064586 0.
 [73] 0.0369663196 0.0291401168 0.0243666881 0.0195932595 0.0063830732 0.0000000000 0.
[85] 0.0262538576 0.0253657778 0.0166514953 0.0069381230 0.0000000000 0.0409071735 0.
 [97] 0.0228125486 0.0175395751 0.0062165582 0.0013321196 0.0160409405 0.0289736018 0.
```

[109] 0.0315268311 0.0134322062 0.0000000000 0.0269754224 0.0401856087 0.0444039875 0. [121] 0.0047734287 0.0000000000 0.0069381230 0.0219799738 0.0434048978 0.0539508448 0.

Theoph\$mM = Theoph\$conc * concUnitConv("mg/L", "mmol/L", MW=180.164)

> Theoph\$mM * concUnitConv("mmol/L", "ug/L", MW=180.164)

[-]	140	2040	0510	10300	3000	0300	0300	1410	0030	3340	3200	0	1120	'
[28]	7500	6200	5300	4900	3700	1050	0	1890	4600	8600	8380	7540	6880	5
[55]	1570	0	1290	3080	6440	6320	5530	4940	4020	3460	2780	920	150	
[82]	7560	6590	5880	4730	4570	3000	1250	0	7370	9030	7140	6330	5660	5
[109]	5680	2420	0	4860	7240	8000	6810	5870	5220	4450	3620	2690	860	
> Theo	ph \$ mM	* conc	UnitCo	onv("mm	ol/L",	"ug/m	L", MW	=180.1	64)					
[1]	0.74	2.84	6.57	10.50	9.66	8.58	8.36	7.47	6.89	5.94	3.28	0.00	1.72	7
[28]	7.50	6.20	5.30	4.90	3.70	1.05	0.00	1.89	4.60	8.60	8.38	7.54	6.88	5
[55]	4									0 40	2 70	0 00	0 1 5	
	1.57	0.00	1.29	3.08	6.44	6.32	5.53	4.94	4.02	3.46	2.78	0.92	0.15	0
				3.08 4.73										
[82]	7.56	6.59	5.88		4.57	3.00	1.25	0.00	7.37	9.03	7.14	6.33	5.66	5

>

As-is R Files

교수님께서 주신 원본 R 파일 입니다.

A.1 Lecture 3

```
Graphics
# 상위수준 그림 함수는 그림을 생성한다.
# 하위수준 그림 함수는 기존의 그림에 그림을 추가한다.
## 상위수준 그림 함수의 주요 인자 (arguments) ###
# main : 제목
# xlab/ylab : x축 및 y축 레이블
# xlim/ylim : x축 및 y축 범위
# col : 색깔
# lty : 선모양
# pch : 점 모양
# cex : 그림 성분의 크기
# lwd : 선 굵기
# type : 그림 타입
```

상위수준 그림 함수 ##########

```
WD <- "D:\\AMC\\Education\\UU\\2017\\R\\Graphics\\"
setwd(WD)
dta <- read.csv("PK.csv")</pre>
head(dta)
str(dta)
plot(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0])
plot(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], log="y")
plot(dta$TIME[dta$MDV==0], log(dta$DV[dta$MDV==0]))
plot(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0]
    , xlab="Time (hr)", ylab="Concentration (ng/mL)"
    , type="o", pch=2, col=1, main="PK time-course of Drug X"
    , x \lim = c(-2,218), y \lim = c(0,80)
plot(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], axes=F,
    , xlab="Time (hr)", ylab="Concentration (ng/mL)"
   , type="o", pch=2, col=1, main="PK time-course of Drug X"
    , x \lim = c(-2,218), y \lim = c(0,80)
axis(1, at=seq(0, 218, 24))
axis(2)
box()
d.demog <- read.csv("DEMOG.csv")</pre>
# histogram
hist(d.demog$HT)
hist(d.demog$HT, breaks=10)
hist(d.demog$HT, nclass=10)
# with density line
hist (d.demog$HT, probability=TRUE, breaks=10)
```

```
hist (d.demog$HT, probability=TRUE, breaks=9, xaxt="n"
      , main="Histogram for Height", xlab="Height (cm)", ylab="Probability (%)")
axis(1, at=seq(min(d.demog$HT), max(d.demog$HT), 3))
lines(density(d.demog$HT))
hist (d.demog$HT, probability=TRUE, breaks=9, xaxt="n"
      , main="Histogram for Height", xlab="Height (cm)", ylab="Probability (%)"
      , col = "lightblue", border = "pink")
axis(1, at=seq(min(d.demog$HT), max(d.demog$HT), 3))
lines(density(d.demog$HT))
########### Box-Whisker Plot ################
# Box-and-Whisker Plot
boxplot(d.demog$WT)
boxplot(d.demog$WT ~ d.demog$SEX)
boxplot(split(d.demog$WT, d.demog$SEX))
boxplot(WT ~ SEX, data=d.demog)
boxplot(d.demog$WT ~ d.demog$SEX
        , names=c("Male", "Female"), ylab="AGE, year", ylim=c(min(d.demog$WT)-2, max(d.
            , col="pink")
boxplot(d.demog$WT ~ d.demog$SEX
        , names=c("Male","Female"), ylab="AGE, year", ylim=c(min(d.demog$WT)-2, max(d.
            , col=c("lightblue", "salmon"), width=c(0.6, 1))
#varwidth: if varwidth is TRUE, the boxes are drawn with widths proportional
#to the square-roots of the number of observations in the groups.
boxplot(d.demog$WT ~ d.demog$SEX
        , names=c("Male", "Female"), ylab="AGE, year", ylim=c(min(d.demog$WT)-2, max(d.
            , col=c("lightblue", "salmon")
            , varwidth=TRUE)
```

```
barplot(d.demog$HT)
VADeaths
barplot(VADeaths, border = "dark blue")
barplot(VADeaths, col = rainbow(20))
barplot(VADeaths, col = heat.colors(8))
barplot(VADeaths, col = gray.colors(4))
barplot(VADeaths, col = gray.colors(4), log="x")
barplot(VADeaths, col = gray.colors(4), log="y")
barplot(VADeaths, col = gray.colors(4), log="xy")
################ pie chart ###################
drug.X.market <- c(0.12, 0.29, 0.32, 0.22, 0.11, 0.28)
names(drug.X.market) <- c("South Korea", "China", "USA", "Japan", "Austria", "EU")</pre>
pie(drug.X.market)
# matrix와 column 사이의 그림
pct.95 <- read.csv("pct95.csv")</pre>
matplot(pct.95[,1], pct.95[,2:ncol(pct.95)], pch=1)
matplot(pct.95[,1], pct.95[,2:ncol(pct.95)], pch=1, col=c(1,2,1), type="l", lty=1, lwc
###### Scatter plot matrices (pairs plots) ######
pairs(d.demog)
```

```
pairs(d.demog, panel = panel.smooth)
 panel.cor <- function(x, y, digits=2, prefix="", cex.cor)</pre>
  {
      usr <- par("usr"); on.exit(par(usr))</pre>
      par(usr = c(0, 1, 0, 1))
      r = (cor(x, y))
      txt <- format(c(r, 0.123456789), digits=digits)[1]</pre>
      txt <- paste(prefix, txt, sep="")</pre>
      if(missing(cex.cor)) cex <- 1.5</pre>
      text(0.5, 0.5, txt, cex = 1.5)
  }
pairs(d.demog, lower.panel=panel.smooth, upper.panel=panel.cor)
하위수준 그림 함수
# points : 점추가
# lines : 선 추가
# abline : 기준선 추가
# mtext : 텍스트 추가
# legend : 설명(legend) 추가
# polygon : polygon 추가
########## 점, 선, 설명 추가 하기 ##########
plot(pct.95$TIME, pct.95$PCT50, main="PK of Drug X"
    , type="l", xlab="Time (h)", ylab="Concentration (ng/ml)"
    , ylim=range(0,80), lty=1, col="red", lwd=2)
plot(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], main="PK of Drug X"
    , type="n", xlab="Time (h)", ylab="Concentration (ng/ml)"
    , ylim=range(0,80))
points(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], pch = 16, cex=0.8)
lines(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], col="black", lwd=1)
abline(40, 0, col="red", lty=2)
                                                         #abline(a,b): y=a+b*x
legend("topright", legend=c("Individual concentrations")
```

```
plot(c(1, 10), c(1, 6), type = "n")
polygon(c(2,8,8,2), c(5,4,3,2), col="lightgreen")
plot(c(1, 9), 1:2, type = "n")
polygon(1:9, c(2,1,2,1,1,2,1,2,1),
       col = c("red", "blue"),
       border = c("green", "yellow"),
       lwd = 3, lty = c("dashed", "solid"))
#--pdf graphics devices
pdf("PK_of_Drug_X.pdf")
plot(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], main="PK of Drug X"
    , type="n", xlab="Time (h)", ylab="Concentration (ng/ml)"
    , ylim=range(0,80))
points(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], pch = 16, cex=0.8)
lines(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], col="black", lwd=1)
abline(40, 0, col="red", lty=2)
                                                          #abline(a,b): y=a+b*x
legend("topright", legend=c("Individual concentrations")
      , lty=1, col="black")
dev.off()
#--PNG graphics devices
png("PK_of_Drug_X.png")
plot(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], main="PK of Drug X"
    , type="n", xlab="Time (h)", ylab="Concentration (ng/ml)"
    , ylim=range(0,80)
points(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], pch = 16, cex=0.8)
lines(dta$TIME[dta$MDV==0], dta$DV[dta$MDV==0], col="black", lwd=1)
abline(40, 0, col="red", lty=2)
                                                          #abline(a,b): y=a+b*x
legend("topright", legend=c("Individual concentrations")
      , lty=1, col="black")
```

A.2 Lecture 4

```
# 2017-03-29

setwd("D:/Rt")
dir()

mydata = read.csv("MyData2017.csv", as.is=TRUE)

Theoph
library(lattice) # trellis

xyplot(conc ~ Time | Subject, data=Theoph)

xyplot(conc ~ Time | Subject, data=Theoph, type="b")

Theoph[,"ID"] = as.numeric(as.character(Theoph[,"Subject"]))

xyplot(conc ~ Time | ID, data=Theoph, type="b")
```

```
write.csv(Theoph, "Theoph.csv", row.names=FALSE, quote=FALSE, na="")
IDs = sort(unique(Theoph[,"ID"])) ; IDs
nID = length(IDs) ; nID
demog = unique(Theoph[,c("ID","Wt")])
colnames(demog) = c("ID", "BWT")
write.csv(demog, "1-demog.csv", row.names=FALSE, quote=FALSE, na="")
DV = Theoph[,c("ID","Time", "conc")]
colnames(DV) = c("ID", "TIME", "DV")
write.csv(DV, "3-DV.csv", row.names=FALSE, quote=FALSE, na="")
adm = cbind(IDs, rep(0, nID), rep(320, nID))
colnames(adm) = c("ID", "TIME", "AMT")
write.csv(adm, "2-adm.csv", row.names=FALSE, quote=FALSE, na="")
demog = read.csv("1-demog.csv", as.is=TRUE)
adm = read.csv("2-adm.csv", as.is=TRUE)
dv = read.csv("3-dv.csv", as.is=TRUE)
AdmDv = merge(adm, dv, by=intersect(colnames(adm), colnames(dv)), all=TRUE)
DataAll = merge(demog, AdmDv, by=c("ID"), all=TRUE)
```

Ayptot (conc. Time | astractor (ib); data-incopii, type-b

A.3 Lecture 5

```
# 2017-04-05 R-intro.pdf Chapter 08

pois
# ?dbeta
dnorm(0)
pnorm(0)
1 - pnorm(1.96)
```

```
pnorm(1.96, lower.tail=FALSE)
qnorm(0.5)
qnorm(0.975)
format(qnorm(0.975), digits=22)
rnorm(5)
rnorm(5, 10, 1)
x = rnorm(100, 10, 1)
mean(x)
sd(x)
2*pt(-2.43, df = 13)
2*pt(-2.43, df = 1000)
qnorm(0.995)
qf(0.01, 2, 7, lower.tail = FALSE)
# ?fivenum
faithful
str(faithful)
eruptions
attach(faithful)
eruptions
waiting
stem(waiting)
sort(eruptions)
hist(eruptions)
hist(eruptions, seq(1.6, 5.2, 0.2), prob=TRUE)
lines(density(eruptions, bw=0.1))
rug(eruptions)
# ?hist
# ?density
lines(density(eruptions, bw="SJ"), lty=3)
plot(ecdf(eruptions), do.points=FALSE, verticals=TRUE)
# ?plot
ecdf(eruptions)
x = ecdf(eruptions)
str(x)
x()
plot(ecdf(eruptions), do.points=FALSE)
```

```
long <- eruptions[eruptions > 3]
x \leftarrow seq(3, 5.4, 0.01)
pnorm(x, mean=mean(long), sd=sqrt(var(long)))
# ?par
x \leftarrow rt(250, df = 5)
qqnorm(x); qqline(x)
curve(dnorm, -5, 5)
y = density(x)
lines(y, lty=3)
# ?ppoints
ppoints (250)
ppoints(10)
qqplot(qt(ppoints(250), df = 5), x, xlab = "Q-Q plot for t dsn")
windows()
qqplot(qt(runif(250), df = 5), x, xlab = "Q-Q plot for t dsn")
# ?shapiro.test
# ?ks.test
# ?t.test
A = c(79.98, 80.04, 80.02, 80.04, 80.03, 80.03, 80.04, 79.97, 80.05, 80.03, 80.02, 80.
B = c(80.02, 79.94, 79.98, 79.97, 79.97, 80.03, 79.95, 79.97)
boxplot(A, B)
t.test(A, B)
var.test(A, B)
t.test(A, B, var.equal=TRUE)
wilcox.test(A, B)
plot(ecdf(A), do.points=FALSE, verticals=TRUE, xlim=range(A, B))
plot(ecdf(B), do.points=FALSE, verticals=TRUE, add=TRUE)
ks.test(A, B)
# Chapter 9 Grouping, loops and conditional execution
# { } does grouping
# Usefulness of loops: for >> while >> repeat
for (i in 1:10) {
  print(2*i)
}
```

```
while ( ) {
# Statements
# # if ~ else ~
# if ( ) {
# # Statements 1
# } else {
# # Statements 2
# }
# if ( ) # Statement1
# else # Statement2
# if ( ) {
# # Statements 1
# } else if ( ) {
# # Statements 2
# } else if ( ) {
# # Statements 3
# } else {
# # Statements 4
# }
# Chapter 10 Writing your own functions
Square = function(x=0)
{
  return(x*x)
}
twosam = function(y1, y2)
{
  n1 = length(y1)
  n2 = length(y2)
  yb1 = mean(y1)
  yb2 = mean(y2)
  s1 = var(y1)
```

101 (1 III 1 1 1 0) princ(2 1)

```
s = ((n1 - 1)*s1 + (n2 - 1)*s2)/(n1 + n2 - 2)
  tst = (yb1 - yb2)/sqrt(s*(1/n1 + 1/n2))
  return (tst)
}
x = rnorm(10)
y = rt(10, 5)
twosam(x, y)
T.test = function(y1, y2)
{
  n1 = length(y1)
  n2 = length(y2)
  yb1 = mean(y1)
  yb2 = mean(y2)
  s1 = var(y1)
  s2 = var(y2)
  s = ((n1 - 1)*s1 + (n2 - 1)*s2)/(n1 + n2 - 2)
  tst = (yb1 - yb2)/sqrt(s*(1/n1 + 1/n2))
  DF = n1 + n2 - 2
  p.val = 2*(1 - pt(abs(tst), df=DF))
  Res = list(tst, DF, p.val, yb1, yb2)
  names(Res) = c("t", "df", "p-value", "mean of x", "mean of y")
  return (Res)
}
res = T.test(x, y)
t.test(x, y)
bslash = function(X, y)
{
  X = qr(X)
 return (qr.coef(X, y))
}
regcoeff = bslash(Xmat, yvar)
```

```
M = matrix(c(2,1,1,2), nrow=2) ; M
M %^% 0.5
sqrtM = M%^{\circ}.5; sqrtM
sqrtM %*% sqrtM
area = function(f, a, b, eps=1.0e-06, lim=10)
{
  fun1 = function(f, a, b, fa, fb, a0, eps, lim, fun)
  {
  ## function 'fun1'is only visible inside 'area'
    d = (a + b)/2
    h = (b - a)/4
    fd = f(d)
    a1 = h * (fa + fd)
    a2 = h * (fd + fb)
    if (abs(a0 - a1 - a2) < eps | lim == 0)
      return (a1 + a2)
    else {
      return (fun(f, a, d, fa, fd, a1, eps, lim - 1, fun) + fun(f, d, b, fd, fb, a2, e
    }
  }
  fa = f(a)
  fb = f(b)
  a0 = ((fa + fb) * (b - a))/2
  fun1(f, a, b, fa, fb, a0, eps, lim, fun1)
}
area(dnorm, 0, 1)
integrate(dnorm, 0, 1)
pnorm(1) - pnorm(0)
f = function(x)
  y = 2*x
  print(x)
  print(y)
  print(z)
```

- runction(3, pow) with (eigen(3), vectors % % (abs(vatues) pow

```
f(1)
z = 3
f(1)
cube = function(n) {
  sq = function() n*n
 n*sq()
cube(5)
open.account = function(total)
{
  list(
    deposit = function(amount)
    {
      if(amount <= 0)</pre>
      stop("Deposits must be positive!\n")
      total <<- total + amount
      cat(amount, "deposited. Your balance is", total, "\n\n")
    },
    withdraw = function(amount)
      if(amount > total)
      stop("You don't have that much money!\n")
      total <<- total - amount
      cat(amount, "withdrawn. Your balance is", total, "\n\n")
    },
    balance = function()
      cat("Your balance is", total, "\n\n")
    }
  )
}
ross = open.account(100)
robert = open.account(200)
ross$balance()
robert$balance()
ross$deposit(50)
```

```
# More basic keywords and functions
1 %in% c(1,2,3,4)
5 %in% c(1,2,3,4)
is.finite(Inf)
prod(1:3)
cummax(1:10)
cummax(10:1)
# ?xor
x = 11:20
which(x==3)
which(x==13)
length(x)
y = "my string"
length(y)
nchar(y)
strsplit(y, " ")
strsplit(y, " ")[[1]]
substr(y, 4, 5)
sample(1:10)
sample(1:10, 20)
sample(1:10, 20, replace=TRUE)
sample(rep(1:10,2))
```

ross\$withdraw(500)

R Tips

- Handbook of statistical distributions with applications¹ (Krishnamoorthy, 2006)
- Materials
 - https://cran.r-project.org/manuals.html
 - https://cran.r-project.org/doc/manuals/r-release/R-intro.pdf
- Changing defualt R console size and etc: 배균섭 교수님께서 알려주신 tip을 참고하여 video clip을 만들었습니다. 매일같이 마주하게 되는 R console이 너무 작게 느껴지 셨다면 다음의 동영상을 참고하셔서 초기 세팅 (Rconsole 파일)을 바꿔서 해결할 수 있습니다. https://youtu.be/uSunEN8W5Mo

B.1 Using Coursera

PAGK에 보낸 이메일을 그대로 옮겼습니다.

배균섭 교수님의 추천을 받아 다음과 같은 강의와 책을 공유하고자 합니다. Coursera.com에 유익한 R 강좌가 열렸습니다. "Mastering Software Development in R Specialization"이란 제목의 강좌인데 4개 Course를 무료로 들을 수 있게 되어있습니다. https://www.coursera.org/specializations/r 이것이 본래의 Link인데 여기서 각각의 Course를 찾거나 혹은 아래의 링크를 각각 클릭하여 하단에 나오는 "Audit"을 클릭하면 무료로 들을 수 있습니다.

Audit 버튼이 안보이신다고 하신 분들이 몇분 계셔서 첨언합니다. Coursera 회원가입하시고 로그인 한 뒤, Enroll Now를 누르시면 Audit 혹은 청강하기 라디오버튼을 보실 수있습니다. 앱에서도 마찬가지입니다. 이외에도 코세라에는 많은 유익한 강의가 있는 것 같습니다. 다만 코스(Course)의 묶음인 "Specialization" 에서는 유료등록(Enroll) 밖에

¹http://www.stat.rice.edu/~dobelman/textfiles/DistributionsHandbook.pdf

검색해서 "Audit(청강)" 하시면 무료로 강의를 들을 수 있습니다.

- https://www.coursera.org/learn/r-programming-environment
- https://www.coursera.org/learn/advanced-r
- https://www.coursera.org/learn/r-packages
- https://www.coursera.org/learn/r-data-visualization

https://bookdown.org/rdpeng/RProgDA/ 이 링크는 무료로 공개된 강의 책자입니다. 강의를 듣지 않고 책으로 보고 싶으신 분은 참고하시면 됩니다.

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- 1. Dr. Jekyll
- 2. Hyde

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