Weird Statistics Questions

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I. Descriptive Statistics

1. How to represent and summarize a variable? Why do we calculate mean and standard deviation for a variable? How about if the variable is skewed or discrete?

The most accurate way to represent a variable is to visualize it as a distribution histogram (distribution is close to the raw data). We usually use mean (μ) to quantify the central tendency (typical value) of the distribution and $standard\ deviation\ (\sigma)$ to characterize the degree of dispersion/spread. In addition, we use quantiles to measure locations of a distribution.

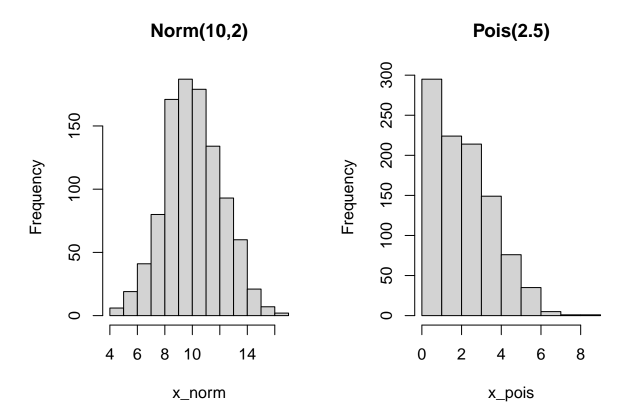
Notes:

- mean is just one of the measures of central tendency. Another two common measures are mode and median. Which measure is the most appropriate to use depends on the questions and also the shape of the distribution.
- standard deviation is defined according to the mean. If mean is not clearly defined, there is no standard deviation either. Given the association between mean and standard deviation, it is also meaningless to compare standard deviations across data sets. To compare dispersion across groups, we can use the coefficient of variation: $CV = \sigma/\mu$.
- if the distribution curve can be described mathematically using a distribution formula, the parameters of the formula will be the accurate measures.

For example, a Normal distribution (bell curve) could be described as $X \sim \mathsf{Norm}(\mu, \sigma)$, while a Poisson distribution (right skewed curve) could be described as $X \sim \mathsf{Pois}(\lambda)$. In a Normal distribution, mean and standard deviation are the two parameters that determine the distribution. Given the popularity of Normal distribution in the real world, the two parameters are commonly used measures to characterize data sets (represented as distribution histogram). In a Poisson distribution, λ is the only parameter (rate parameter), which is the total number of events (k) divided by the number of units (n) in the data $(\lambda = k/n)$.

In R, we use rnorm and rpois functions to generate Normal and Poisson distributions respectively, by setting the respective parameters.

```
x_norm = rnorm(1000, 10, 2) # mean = 10, sd = 2
x_pois = rpois(1000,2.5) # lambda = 2.5
par(mfrow=c(1,2))
hist(x_norm,main="Norm(10,2)")
hist(x_pois,main="Pois(2.5)")
```



As expected, the means of the generated data are close to the theoretical ones:

c(mean(x_norm),mean(x_pois))

[1] 10.06601 2.54900

It is interesting to calculate the standard deviations:

c(sd(x_norm),sd(x_pois))

[1] 2.058826 1.575462

The standard deviation of the Poisson distribution is 1.58. The variance is the standard deviation² = 2.48, which is close to the mean λ . In fact, E[X] = Var[X] for Poisson distribution. We will explain the formula soon below.

Question:

• As suggested in some text books, we should use *median* or *mode* to characterize the central tendency of skewed distributions. The median for x_pois is 2. Is this value informative, better than the *mean*?

mean and thus standard deviation are not well defined for categorical variables. The distribution could be described as $f(x = k) = p_k$. We use the sample function to generate two categorical variables by defining p_k s.

```
x_cat1 = sample(8:12, size = 1000, replace = TRUE, prob = c(0.25,0.1,0.3,0.25,0.1))
x_cat2 = sample(8:12, size = 1000, replace = TRUE, prob = c(0.1,0.25,0.3,0.1,0.25))
c(mean(x_cat1),mean(x_cat2)) #means
```

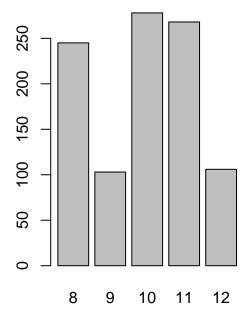
```
## [1] 9.887 10.153
```

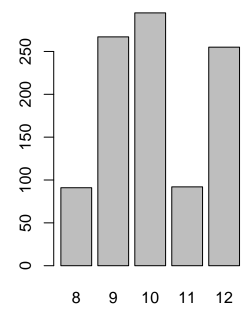
```
c(sd(x_cat1),sd(x_cat2)) #standard deviations
```

```
## [1] 1.328155 1.311988
```

Even though the *means* are hard to interpret (what does it mean if I say the average of race in Hong Kong is 3.1?), the above calculation indicates that the two variables have similar *means* and *standard deviations*. They are actually two different distributions, as presented below:

```
tbl_cat1 = table(x_cat1)
tbl_cat2 = table(x_cat2)
par(mfrow=c(1,2))
barplot(tbl_cat1)
barplot(tbl_cat2)
```





The best way is to calculate the frequency tables:

```
tbl_cat1 = data.frame(tbl_cat1)
tbl_cat1$Prob_x1 = tbl_cat1$Freq/sum(tbl_cat1$Freq)
tbl_cat1$Category = paste0(tbl_cat1$x_cat1)
t1 = tbl_cat1[,c("Category","Prob_x1")]

tbl_cat2 = data.frame(tbl_cat2)
tbl_cat2$Prob = tbl_cat2$Freq/sum(tbl_cat2$Freq)
t1$Prob_x2 = tbl_cat2$Prob
```

```
##
     Category Prob x1 Prob x2
## 1
                 0.245
            8
                          0.091
## 2
            9
                 0.103
                          0.267
                 0.278
                          0.295
## 3
           10
## 4
           11
                 0.268
                          0.092
## 5
           12
                 0.106
                          0.255
```

Question:

Are there any continuous distributions without mean or variance? See Pareto distribution.

If there are many distributions without "meaningful" mean or variance, why are they so popular measures in statistics? Let's introduce another concept – Expected Value.

- For discrete distributions: $E[X] = \sum_{k=1}^{n} x_k P(X = x_k)$.
- For continuous distributions: $E[X] = \int_{-\infty}^{\infty} x f(x) dx$.

```
x = c(8:12)
px_cat1 = tbl_cat1$Prob_x1 # estimated probs
px_cat2 = c(0.1,0.25,0.3,0.1,0.25) # true probs

EX_cat1 = sum(x*px_cat1) # expectation
EX_cat2 = sum(x*px_cat2) # expectation

EX_cat1 - mean(x_cat1) # the same
```

```
## [1] 0
```

```
EX_cat2 - mean(x_cat2) # slightly different
```

```
## [1] -0.003
```

Are there any cases that $E[X] \neq mean(X)$? Biased samples! The expectation, by definition, is true (of he population), while mean might not be true in biased samples.

For Normal distribution, $E[X] = \mu \approx mean$. For Poisson distribution, $E[X] = \lambda \approx mean$. If expectation is the only parameter for the distribution (e.g., λ in Poisson), the mean suffices to characterize the distribution. That means if we know the parameter λ , we can recover the distribution using the formula: $Pois(\lambda) = \frac{\lambda^k e^{-\lambda}}{k!}$. However, mean (μ) is not sufficient to characterize a Normal distribution because it varies with standard deviation (σ) too. For categorical distribution, we need k-1 probabilities (p_k) to characterize the distribution.

Specifically, a binary variable with 0 and 1, we only need 1 probability (the proportion of 1) to describe the distribution $(E[X] = p, Var[X] = E[(X - E[X]^2)] = p(1 - p))$.

Therefore, to describe a (sampled) data precisely, we need to know the underlying distribution (that generated the observed data) first. And then, we could estimate the corresponding parameters based on the sample. But what should we do if we don't know the underlying distribution?

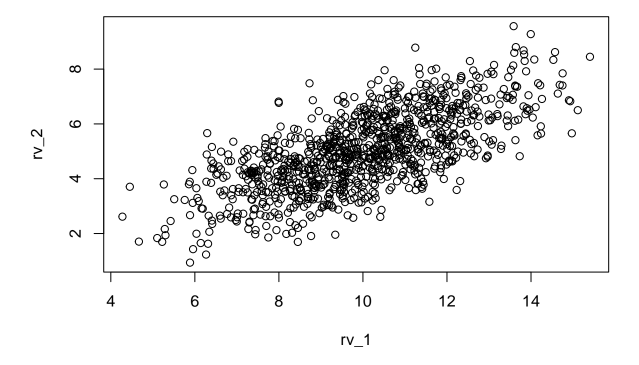
If the function is a probability distribution, then the first moment is the expected value (mean), the second central moment is the variance $(sd^2: Var[X] = E[(X - E[X]^2)])$, the third standardized moment is the skewness (skewness), and the fourth standardized moment is the kurtosis (kurtosis)... Even if the E(X) is not directly interpretable, it remains important when we consider the relationships between a sample and the population (see Q3).

Nevertheless, mean and sd could be interpreted in skewed distributions. Chebyshev's inequality says that at least $1-1/K^2$ of data from a sample must fall within K standard deviations from the mean, where K is any positive real number greater than one.

- For K=2 we have $1-1/K^2=1$ 1/4=3/4=75%. So Chebyshev's inequality says that at least 75% of the data values of any distribution must be within two *standard deviations* of the *mean*.
- For K = 3 we have $1 1/K^2 = 1$ 1/9 = 8/9 = 89%. So Chebyshev's inequality says that at least 89% of the data values of any distribution must be within three standard deviations of the mean.
- For K=4 we have $1-1/K^2=1$ 1/16=15/16=93.75%. So Chebyshev's inequality says that at least 93.75% of the data values of any distribution must be within four *standard deviations* of the *mean*.
- 2. We use correlation coefficients (why plural?) to quantify the (linear) relationships between two continuous variables, how can we show the relationships involving categorical/rank variables?

```
rv_1 = rnorm(1000,10,2)
rv_2 = 0.5*rv_1 + rnorm(1000)
cor.test(rv_1,rv_2)
```

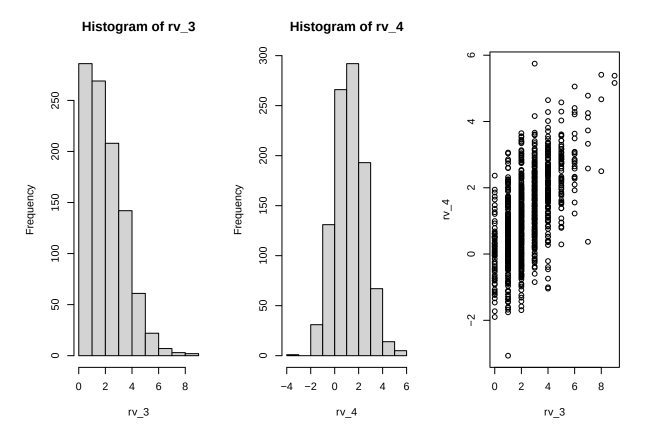
```
##
## Pearson's product-moment correlation
##
## data: rv_1 and rv_2
## t = 30.158, df = 998, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.6566285 0.7216161
## sample estimates:
## cor
## 0.6905133</pre>
```



The Pearson coefficient r=0.69 is larger than 0.5. Why? $r^2=0.48\approx 0.5$. Now, we generate a non-normal variable rv_3.

```
rv_3 = rpois(1000,2.5)
rv_4 = 0.5*rv_3 + rnorm(1000)

par(mfrow=c(1,3))
hist(rv_3)
hist(rv_4)
plot(rv_3,rv_4)
```



We have different ways to calculate the correlation between two continuous variables (rank, interval, or ratio): Pearson's r, Spearman's ρ , and Kendall's τ .Please check the method definitions here.

```
cor.test(rv_3,rv_4,method = "pearson")
##
   Pearson's product-moment correlation
##
##
## data: rv_3 and rv_4
## t = 23.687, df = 998, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
    0.5586884 0.6381638
## sample estimates:
##
        cor
## 0.599904
cor.test(rv_3,rv_4,method = "spearman")
## Warning in cor.test.default(rv_3, rv_4, method = "spearman"): Cannot compute
## exact p-value with ties
##
##
    Spearman's rank correlation rho
##
```

```
## data: rv_3 and rv_4
## S = 69355647, p-value < 2.2e-16
## alternative hypothesis: true rho is not equal to 0
  sample estimates:
##
         rho
## 0.5838657
cor.test(rv_3,rv_4,method = "kendall")
##
   Kendall's rank correlation tau
##
##
## data: rv_3 and rv_4
## z = 19.375, p-value < 2.2e-16
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
##
         tau
## 0.4449666
```

we can see that Pearson and Spearman correlations are roughly the same, but Kendall is very much different. That's because Kendall is a test of strength of **dependence** (i.e. one could be written as a linear function of the other), whereas Pearson and Spearman are nearly equivalent in the way they correlate normally distributed data. All of these correlations are correct in their result, it's just that Pearson and Spearman are looking at the data in one way, and Kendall in another.

In fact, the Pearson' r of the ranks of the original variables is equivalent to Spearman's ρ :

```
cor(rv_1,rv_2,method = "pearson") # pearson

## [1] 0.6905133

cor(rank(rv_1),rank(rv_2),method = "pearson") # ranks -> pearson

## [1] 0.6815647

cor(rv_1,rv_2,method = "spearman") #spearman
```

```
## [1] 0.6815647
```

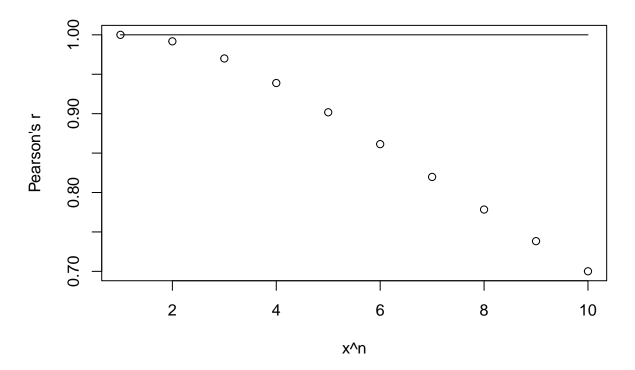
Pearson's correlation (r) is a measure of the **linear** relationship between two continuous random variables. It does not assume normality although it does assume finite variances and finite covariance (Cov(X,Y) = E[(X-E(X))(Y-E(Y))] = E[XY] - E[X]E[Y]). When the variables are Bivariate Normal (the sampling distribution of $r\sqrt{\frac{n-2}{1-r^2}}$ will follow a Student's t-distribution), Pearson's correlation provides a complete description of the association. In order words, violation of Bivariate Normal may influence the significance tests (when sample size is small).

Spearman's correlation (ρ) applies to ranks and so provides a measure of a **monotonic** (could be non-linear) relationship between two continuous random variables. It is also useful with ordinal data and is robust to outliers (unlike Pearson's correlation). When correlating skewed variables, particularly highly skewed variables, a log or some other transformation often makes the underlying relationship between the two variables clearer. In such settings it may be that the raw metric is not the most meaningful metric

anyway. Spearman's ρ has a similar effect to transformation by converting both variables to ranks. From this perspective, Spearman's ρ can be seen as a quick-and-dirty approach (or more positively, it is less subjective) whereby you don't have to think about optimal transformations.

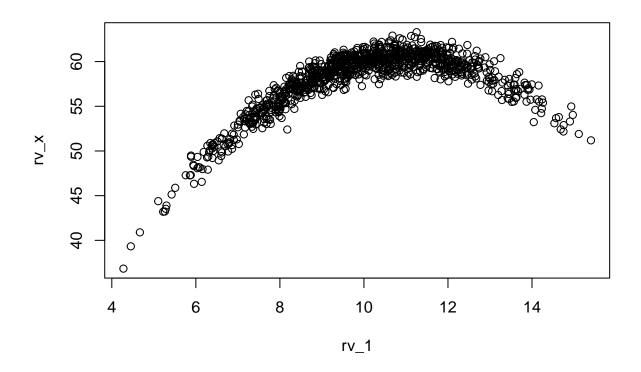
The two coefficients will be very different, if the relationship between the variable is non-linear. The following R code shows how the correlations between x and x^n differ over n.

```
rs = sapply(1:10,function(i)cor(rv_1,rv_1^i,method = "pearson"))
rhos = sapply(1:10,function(i)cor(rv_1,rv_1^i,method = "spearman"))
plot(rs,ylab = "Pearson's r",xlab="x^n")
lines(rhos)
```



Spearman's ρ vs. Kendall's τ (tau). These two are so much computationally different that you cannot directly compare their magnitudes. Spearman is usually higher by 1/4 to 1/3 and this makes one incorrectly conclude that Spearman is "better" for a particular dataset. The difference between ρ and τ is in their ideology, proportion-of-variance for ρ and probability for τ . ρ is a usual Pearson's r applied for ranked data. τ is more "general" than ρ , for ρ is warranted only when you believe the underlying (model, or functional in population) relationship between the variables is strictly **monotonic**. While τ allows for **non-monotonic** underlying curve and measures which monotonic "trend", positive or negative, prevails there overall. ρ is comparable with r in magnitude; τ is not.

```
rv_x = -0.5*rv_1^2+11*rv_1+rnorm(1000) # it is non-linear non-monotonic
plot(rv_1,rv_x)
```



```
cor.test(rv_1,rv_x, method = "pearson")
##
   Pearson's product-moment correlation
##
##
## data: rv_1 and rv_x
## t = 22.607, df = 998, p-value < 2.2e-16
\#\# alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.5394177 0.6215208
## sample estimates:
##
         cor
## 0.5819503
cor.test(rv_1,rv_x, method = "spearman")
##
##
    Spearman's rank correlation rho
##
## data: rv_1 and rv_x
## S = 76875234, p-value < 2.2e-16
## alternative hypothesis: true rho is not equal to 0
## sample estimates:
##
         rho
## 0.5387481
```

```
cor.test(rv_1,rv_x, method = "kendall")
```

```
##
## Kendall's rank correlation tau
##
## data: rv_1 and rv_x
## z = 19.576, p-value < 2.2e-16
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
## tau
## 0.4134214</pre>
```

Cosine similarity (a kind of dependence!) between two vectors A and B is defined as

$$CosSim = \frac{\sum (A_i B_i)}{(\sqrt{\sum A_i^2} \sqrt{\sum B_i^2})}$$

Given any two vectors, the similarity score could be calculated in the following way:

```
a = sum(rv_1*rv_x)
b = sqrt(sum(rv_1*rv_1))*sqrt(sum(rv_x*rv_x))
a/b
```

```
## [1] 0.9860533
```

Finally, how to quantify the "correlation" (dependence is a more appropriate term here) between two categorical variables:

```
tbl = matrix(data=c(65, 45, 20, 40), nrow=2, ncol=2, byrow=T)
dimnames(tbl) = list(City=c('B', 'T'), Gender=c('M', 'F'))
tbl
```

```
## Gender
## City M F
## B 65 45
## T 20 40
```

```
chi2 = chisq.test(tbl, correct=T)
c(chi2$statistic, chi2$p.value)
```

```
## X-squared
## 9.298484848 0.002293435
```

 χ^2 is NOT normalized. In a $k \times l$ contingency table, the theoretical maximal value of χ^2 is N(min(k,l)-1), So, we define **Cramer's** V: $V = \sqrt{\frac{\chi^2}{N(min(k,l)-1)}}$. V ranges from 0 to 1. Therefore,

```
sqrt(chi2$statistic/sum(tbl))
```

```
## X-squared
## 0.2338738
```

Alternatively, you can use the cramersv function in lsr package.

```
library(lsr)
cramersV(tbl)
```

```
## [1] 0.2338738
```

The **uncertainty coefficient** (also called entropy coefficient or Thiel's U) is a measure of nominal association. It is based on the concept of information entropy. the uncertainty coefficient ranges in [0,1]. And please read other methods.

```
library(DescTools)
UncertCoef(tbl, direction = "column")

## [1] 0.04435282

UncertCoef(tbl, direction = "row")

## [1] 0.04735172

UncertCoef(tbl, direction = "symmetric")
```

[1] 0.04580323

A final question: is it possible that the association between categorical variables is negative?

II. Inferential Statistics (Tests)

3. Why can we use a sample to infer the population, under what conditions? What is a sampling distribution? Is it observable? Why does large sample size work?

A simple random sample from a population could be used to estimate the population characteristics.

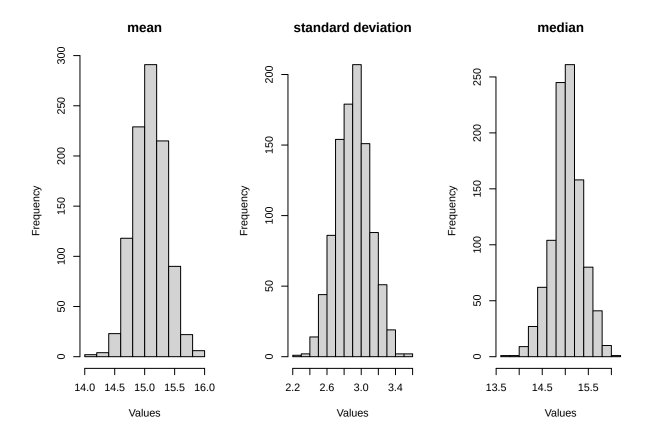
[1] 15.394123 2.776738 15.405100

The estimates are close to the true values, thought not always the same. Let's repeat sampling many times $(N \to +\infty)$ and plot the **sample means** (\overline{X}) -the means of all samples?

```
stats = matrix(NA,nrow = 1000,ncol = 3)

for (i in 1:1000){
    s = sample(rv_1,100) # sample size = 100
    stats[i,] = c(mean(s),sd(s),median(s))
}

par(mfrow=c(1,3))
hist(stats[,1],main="mean",xlab="Values")
hist(stats[,2],main="standard deviation",xlab="Values")
hist(stats[,3],main="median",xlab="Values")
```



We call these **sampling distributions (SMD)** of *means*, *sds*, and *medians*. All these distributions should be Normal. And then, it is straightforward to calculate the *mean* and *sd* of the distributions. For the first one, the sampling distribution of *means*:

```
mean(stats[,1])
```

[1] 15.08502

It is almost the same to the population mean. How about the means of of the second and third distributions (sampling distributions of standard deviations and medians)? The pattern remains!

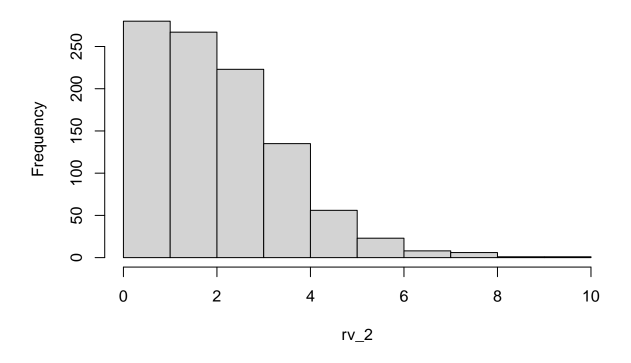
```
c(mean(stats[,2]),mean(stats[,3]))
```

[1] 2.906849 15.042164

Does the pattern apply to non-normal distributions? Let's simulate a Poisson variable:

```
rv_2 = rpois(1000,2.5)
hist(rv_2)
```

Histogram of rv_2

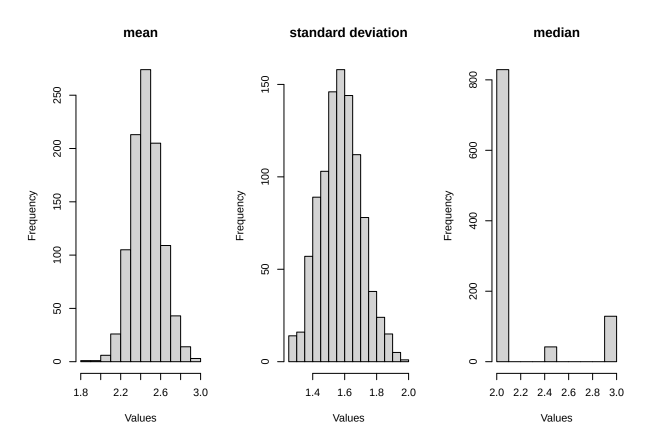


Repeat the previous process:

```
# samples
stats = matrix(NA,nrow = 1000,ncol = 3)

for (i in 1:1000){
    s = sample(rv_2,100) # sample size = 100
    stats[i,] = c(mean(s),sd(s),median(s))
}

par(mfrow=c(1,3))
hist(stats[,1],main="mean",xlab="Values")
hist(stats[,2],main="standard deviation",xlab="Values")
hist(stats[,3],main="median",xlab="Values")
```



```
c(mean(stats[,1]),mean(stats[,2]),mean(stats[,3]))
```

[1] 2.463200 1.574401 2.150000

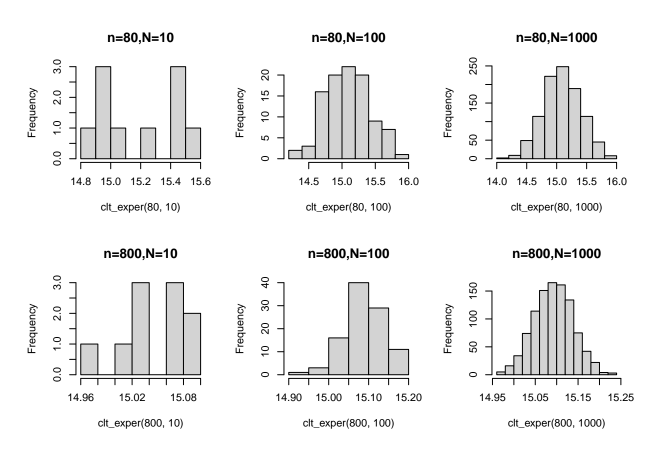
Sample size n vs. sampling times $N: N \to +\infty$, $SMD \to \mathsf{Norm}(\mu, \sigma/\sqrt{n})$; n approaches the population size, the *standard deviation* of the distribution approaches to 0. Why? **Central Limit Theorem (CLT)**: $\mu_s = \mu$, $\sigma_s = \frac{\sigma}{\sqrt{n}}$, where μ_s is the mean of the sampling distribution, σ_s is the standard deviation of the sampling distribution.

```
# define a function and change n and N:
clt_exper = function(n,N){
    means = c()
for (i in 1:N){
    s = sample(rv_1,n)
    means = c(means,mean(s))
}

return(means)
}

# plot the histograms
par(mfrow=c(2,3))
hist(clt_exper(80,10),main = "n=80,N=10")
hist(clt_exper(80,100),main = "n=80,N=100")
hist(clt_exper(80,100),main = "n=80,N=100")
hist(clt_exper(80,100)),main = "n=80,N=1000")
hist(clt_exper(800,100),main = "n=80,N=1000")
hist(clt_exper(800,100),main = "n=800,N=100")
```

```
hist(clt_exper(800,100), main = "n=800, N=100")
hist(clt_exper(800,1000), main = "n=800, N=1000")
```



As you can see, large sampling times N leads to more "normal" distributions, while large sample size n leads to smaller variance of the distributions.

```
c(mean(clt_exper(80,1000)),mean(clt_exper(800,1000)))
```

[1] 15.09374 15.08929

```
c(sd(clt_exper(80,1000)),sd(clt_exper(800,1000)))
```

[1] 0.31434754 0.04342242

The increase of sample size didn't change the means, however, it decreased the $standard\ deviation$ from 0.32 to 0.05

4. What are the relationships between standard errors, sampling errors, confidence interval, and confidence level?

The difference between the sample statistic and the population parameter is considered the **sampling error**. In a sampling distribution of means, where $\mu_s \to \mu$, when $\sigma_s \to 0$ $(n \to N)$.

```
sdm = clt_exper(800,1000)
ms = mean(sdm)
quantile(sdm,probs=c(0.025,0.975))
```

```
## 2.5% 97.5%
## 14.99830 15.17379
```

That means 95% of the sample means are within [14.95,15.14]. Given the sample distribution is a Normal distribution, 95% of the sample means should be around μ : $[\mu - 1.96\sigma, \mu + 1.96\sigma] \approx [\mu_s - 1.96\sigma_s, \mu_s + 1.96\sigma_s]$. The standard deviation of the sampling distribution (σ_s) is called **standard error** (SE).

```
ms-1.96*sd(sdm)
```

[1] 15.00335

```
ms+1.96*sd(sdm)
```

[1] 15.17775

In practice, it is nearly impossible to observe the sampling distribution directly, therefore, we don't know the mean of the sampling distribution (μ_s) , the standard deviation of sampling distribution (σ_s) , population mean (μ) , or population standard deviation (σ) . What we know is the mean of a sample (\bar{x}) , and the standard deviation of a sample (s).

For example, in sample 1 (s_1), the mean is 15.21, standard deviation is 2.84. We know both are different from the population mean and standard deviation (15.09, 2.91). But if we say the mean of any sample 15.21 is the true mean, how likely we are wrong? We should based on the **CLT**, imaging the sampling distribution, and then we have $(SE = \sigma_s = \frac{\sigma}{\sqrt{n}})$:

$$CI_{1-\alpha} = \overline{x} \pm Z_{\alpha/2} \cdot SE = \overline{x} \pm Z_{\alpha/2} \frac{\sigma}{\sqrt{n}}$$

We usually call $\pm Z_{\alpha/2} \cdot SE$ sampling error. If $\alpha = 5\%, 1 - \alpha = 95\%, Z_{\alpha/2} \approx 1.96, \sigma \approx s$ (it is just convenient to use s to replace σ), we will have the **95% confidence interval**:

$$CI_{95\%} = \overline{x} \pm 1.96 \frac{s}{\sqrt{n}}$$

```
m = mean(s_1)
s = sd(s_1)
m-1.96*s/sqrt(100)
```

[1] 14.65423

```
m+1.96*s/sqrt(100)
```

[1] 15.76668

You will find that the true mean 15.09 is within the 95% confidence interval [14.65,15.77]. Is it possible that for some ((random) samples, the true mean is not in the calculated confidence interval? The answer is yes and we even know the probability IS 5%.

5. What are the differences among T-test, Z-test, F-test, χ^2 -test etc.?

A test statistic (TS) is a measure of the difference between the observed data and what we expected from the null hypothesis (by chance). The test statistic gets bigger (in absolute value) as the observed data looks unusual compared the null hypothesis. So a large test statistics cast doubt on the null hypothesis. There are many different kinds of tests.

$$TS = \frac{Observed value - Expected value}{SE}$$

If TS>1.96=>p<.05; TS>2.58=>p<.01 Why? For example, in Z-test, $Z=\frac{\overline{x}-u_0}{SE}\sim \mathsf{Norm}(0,1).$ Therefore, $Z=\frac{\overline{x}-u_0}{SE}>1.96.$ And then, $(\overline{x}-u_0)>1.96\cdot SE=>\overline{x}>u_0+1.96\cdot SE.$ In a Normal distribution, only 2.5% values could be larger than $1.96\cdot SE.$ So, it is very likely that $\overline{x}>u_0.$ In a Z-test, $SE=s/\sqrt{n}.$

What are the differences? The distributions of the test statistics. When the sample size n is relatively small, $T = \frac{\overline{x} - u_0}{SE} \sim \mathsf{StudentT}(\nu, 0, 1)$, where $\nu = n - 1$ is the degree of freedom: $n \to +\infty$, then $\mathsf{StudenT} \to \mathsf{Norm}$

```
# whether mu>14
t.test(s_1,mu=14,alternative="less")
##
##
   One Sample t-test
##
## data: s_1
## t = 4.2653, df = 99, p-value = 1
## alternative hypothesis: true mean is less than 14
## 95 percent confidence interval:
##
        -Inf 15.68166
## sample estimates:
## mean of x
   15.21046
# whether the two samples have equal means:
t.test(s_1,s_2)
##
##
   Welch Two Sample t-test
##
## data: s_1 and s_2
## t = 1.5499, df = 197.93, p-value = 0.1228
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.1710462 1.4271512
## sample estimates:
## mean of x mean of y
   15.21046 14.58240
# we don't assume equal length of the two vectors:
t.test(s_1,rv_1)
##
##
   Welch Two Sample t-test
```

##

```
## data: s_1 and rv_1
## t = 0.40345, df = 120.73, p-value = 0.6873
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.4702333 0.7109370
## sample estimates:
## mean of x mean of y
## 15.21046 15.09010
```

Before we perform a T/Z-test, we need to check assumptions:

- measurement scales: ratio or interval
- simple random sample
- normality (the distribution of sample means!)
- large sample size
- homogeneity of variance (equal variance)

Why does it require a Normal distribution? The t-test is invalid for small samples from non-normal distributions, but it is valid for large samples from non-normal distributions (see online answers).

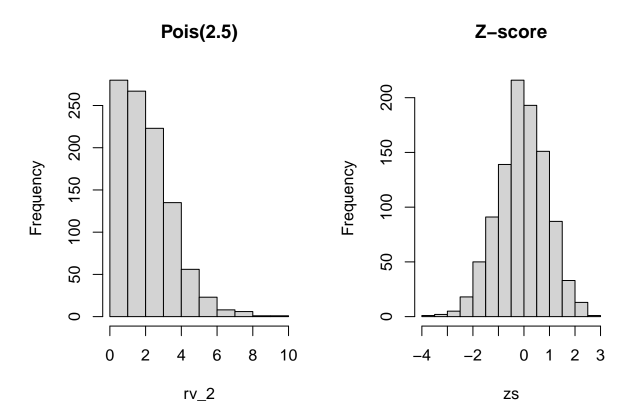
Let's try Poisson distribution:

- generate a variable with a Poisson distribution
- draw a random sample (sp)
- calculate the mean difference between the sample and the population: mean(sp)-mean(rv_2)
- standard error is the population sd divided by the square root of sample size
- get the Z score

```
for (i in 1:1000){
    sp = sample(rv_2,100) # rv_2 here is a poisson distribution we generated before
    m = mean(sp)-mean(rv_2)
    se = sd(sp)/sqrt(100)
    z = m/se

    zs= c(zs,z)
}

par(mfrow=c(1,2))
hist(rv_2, main="Pois(2.5)")
hist(zs,main = "Z-score")
```



The above example shows that the Z-score of a skewed variable is also normally distributed.

Why does it require equal variance (only for two-sample tests)? $T = \frac{\overline{x} - u_0}{SE} \sim \mathsf{StudentT}(\nu, 0, 1)$, where $SE = s/\sqrt(n)$. We extend this to two-sample-mean comparison: $T = \frac{\overline{x}_1 - \overline{x}_2}{SE} \sim \mathsf{StudentT}(\nu, 0, 1)$. It is easy to calculate the sample means (\overline{x}_1 and \overline{x}_2) and sample $standard\ deviations$ (s_1 and s_2). The problem is how to calculate SE. If the two samples are independent to each other, according to the **Variance Sum Law**:

$$\sigma_{x1\pm x2}^2 = \sigma_{x1}^2 + \sigma_{x2}^2$$

Therefore, $SE_{x1-x2} = \sqrt{s_1^2/n_1 + s_2^2/n_2}$. If equal variances are assumed, then $SE_{x1-x2} = \sqrt{\frac{(n_1-1)s_1^2 + (n_2-1)s_2^2}{n_1 + n_2 - 2}}$ (pooled variance estimate). The following example tests two samples from different distributions (different means and variances too).

```
zs_un = c() # z scores unequal variance
zs_eq = c() # z scores equal variance

for (i in 1:1000){
    sp_1 = sample(rv_1,100) # rv_1 normal
    sp_2 = sample(rv_2,100) # rv_2 poisson

# mean difference
    m = mean(sp_1)-mean(sp_2)

# sample sds
    se_1 = sd(sp_1)
```

```
se_2 = sd(sp_2)

# unequal var.
se_un = sqrt(se_1^2/100+se_2^2/100)
# equal var.
se_eq = sqrt(99*(se_1+se_2)/198)

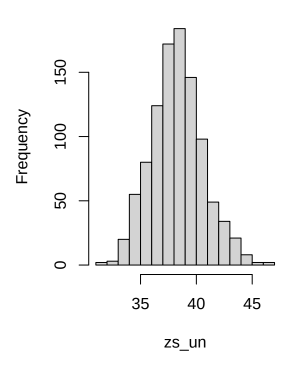
z_un = m/se_un
z_eq = m/(se_eq*sqrt(2/100))

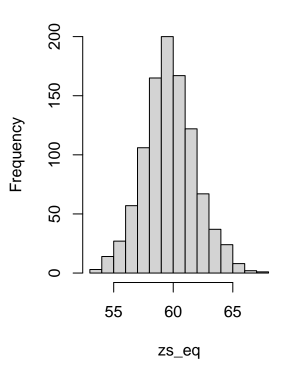
zs_un = c(zs_un,z_un)
zs_eq = c(zs_eq,z_eq)
}

par(mfrow=c(1,2))
hist(zs_un,main="Z-score (Unequal Variance)")
hist(zs_eq,main="Z-score (Equal Variance)")
```

Z-score (Unequal Variance)

Z-score (Equal Variance)





```
c(mean(zs_un),mean(zs_eq))
```

[1] 38.30293 59.70167

Both scores followed the Normal distribution, while it appears the equal variance method tends to overestimate the significance than the unequal variance method, because mean(zs_eq) > mean(zs_un).

Based on "extensive" simulations from distributions either meeting or not meeting the assumptions imposed by a t-test, (normality and homogeneity of variance) that the Welch-tests performs equally well when the assumptions are met (i.e., basically same probability of committing alpha and beta errors) but outperforms the t-test if the assumptions are not met, especially in terms of power. Therefore, they recommend to always use the Welch-test if the sample size exceeds 30.

```
t.test(sp_1,sp_2,var.equal = T)
##
##
   Two Sample t-test
##
## data: sp_1 and sp_2
## t = 37.937, df = 198, p-value < 2.2e-16
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 11.53828 12.80362
## sample estimates:
## mean of x mean of y
   14.80095
               2.63000
t.test(sp_1,sp_2,var.equal = F)
##
##
   Welch Two Sample t-test
##
## data: sp_1 and sp_2
## t = 37.937, df = 174.67, p-value < 2.2e-16
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 11.53776 12.80414
## sample estimates:
## mean of x mean of y
  14.80095
               2.63000
```

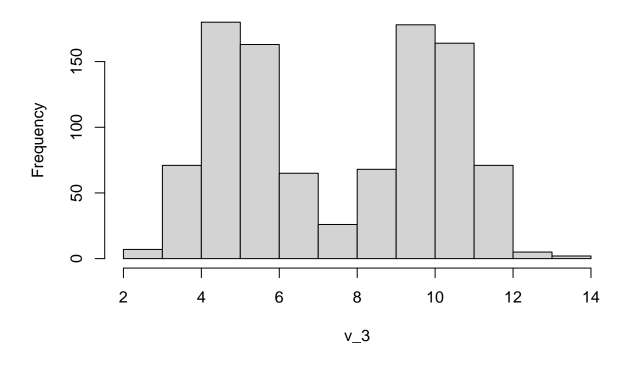
How about the two samples are dependent? Paired T-test. If more than two samples, use F-test in ANOVA (equal or unequal variances?). If both categorical variables, then $TS \to \chi^2(k)$, where k is the degree of freedom.

6. What are distribution-free tests (non-parametric tests)? We test the difference between means, can we test the difference between medians/variances? Can we test the difference between two categorical variables?

The problem that we cannot always use T-test is not because of its non-normality or unequal variances but *mean* is not well defined. Furthermore, for some test statistics (e.g., *mean difference*), we know they follow certain types of distributions. However, for others, we don't know. The distribution in distribution-free does not mean the distribution of the variables but the distribution of the test-statistics. The below distribution is a mixture of two Normal distributions. The *mean* is not the most important characteristic.

```
v_1 = rnorm(500,5,1)
v_2 = rnorm(500,10,1)
v_3 = c(v_1,v_2)
hist(v_3,main="Mixture of two normal distributions")
```

Mixture of two normal distributions

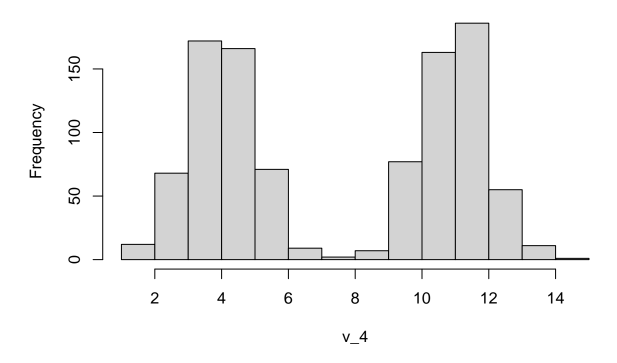


```
mean(v_3)
```

[1] 7.492717

```
# we define another variable in similar way:
v_4 = c(rnorm(500,4,1),rnorm(500,11,1))
hist(v_4)
```

Histogram of v_4



$mean(v_4)$

[1] 7.491419

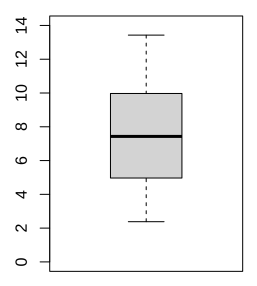
As expected, there is no difference in *means*.

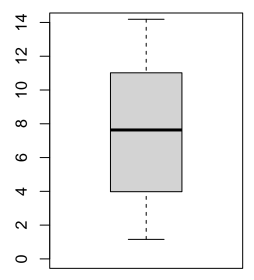
t.test(v_3,v_4) # no difference of mean

```
##
## Welch Two Sample t-test
##
## data: v_3 and v_4
## t = 0.0091126, df = 1841.1, p-value = 0.9927
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.2781582 0.2807551
## sample estimates:
## mean of x mean of y
## 7.492717 7.491419
```

How about quantiles? Are they different?

```
quantile(v_3)
##
          0%
                    25%
                              50%
                                         75%
                                                   100%
    2.379956
               4.969297
                         7.430170
                                    9.972587 13.428165
quantile(v_4)
                                                   100%
##
          0%
                    25%
                               50%
                                         75%
    1.155275
              3.979774 7.634006 11.017895 14.187821
par(mfrow=c(1,2))
boxplot(v_3, ylim=c(0, 14))
boxplot(v_4, ylim=c(0, 14))
```





Yes, quantiles are informative. And the boxplots also suggest that the variances are different. However, what are the standard errors (SEs) of quantiles (or other quantities if defined)? Even if we know the SE, it remains unknown about the distribution of the test statistics. An easy way is to to perform the randomization test, which is one of the most commonly used non-parametric tests.

The below code shows an example of testing variance difference using randomization test (use var.test for a parametric test):

- 1. given any two variables: x1, x2
- 2. calculate the difference of standard deviation: sd(x1)-sd(x2)

3. Permutation

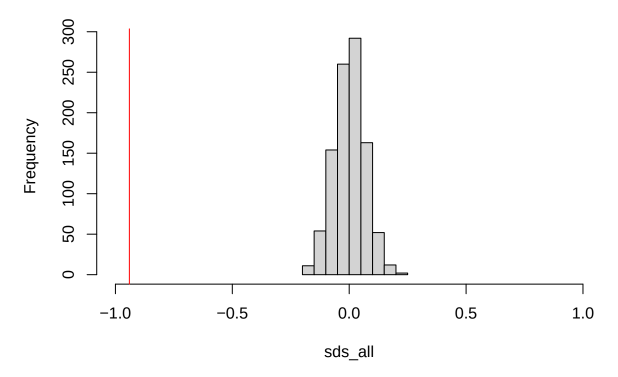
- combine two variables to a single one: values = c(x1,x2)
- randomly permute the values
- split the permuted values into two groups and calculate the difference again
- repeat many time
- 4. compare the real difference with the differences after permutation

```
sts = sd(v_3)-sd(v_4)
sds_all = c()

for (i in 1:1000){ #repeat many times
    values = c(v_3,v_4) # combine all values
    pv = sample(values) # randomly permute the values
    sds = sd(pv[1:500])-sd(pv[501:1000]) # split into two groups and calculate the difference of sds
    sds_all = c(sds_all,sds)
}

hist(sds_all,xlim=c(-1,1))
abline(v=sts,col="red")
```

Histogram of sds_all



```
table(sts<sds_all)/1000 # probability of true!
```

```
##
## TRUE
## 1
```

It means that the observed difference is smaller than 100% of the permuted differences. In other words, it is very unlikely the observed variance difference is caused by chance.

Another method is to use bootstrapping:

- 1. given two variables: x1, x2
- 2. calculate the difference: sd(x1)-sd(x2)
- 3. bootstrap:
 - resample from x1 and x2
 - calculate the difference in the subsamples
 - · repeat many times
- 4. compare the real difference with the differences after permutation

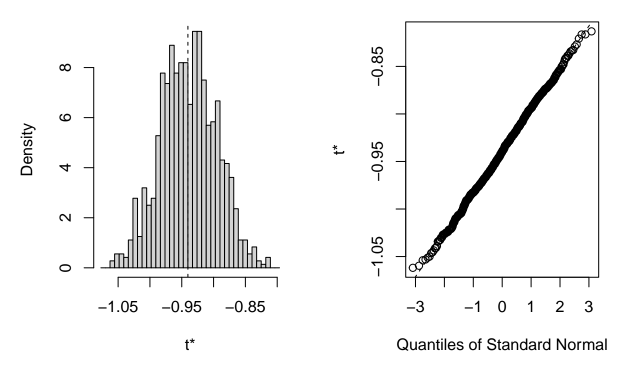
We don't need to do step 3 from the scratch because we can use the boot package in r.

```
library(boot)
data = data.frame(v_3, v_4)
sts = function(d,i){
 d2=d[i,]
 dis = sd(d2$v_3)-sd(d2$v_4)
 return(dis)
boot_sd = boot(data,sts,R=1000) # define the data, define the test statistic, repetition times
boot_sd
##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## boot(data = data, statistic = sts, R = 1000)
##
##
## Bootstrap Statistics :
##
         original
                                std. error
                        bias
## t1* -0.9403314 0.0009803804 0.04395312
boot.ci(boot.out = boot_sd, type = c("norm", "basic", "perc", "bca"))
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 1000 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = boot_sd, type = c("norm", "basic", "perc",
       "bca"))
##
```

```
##
## Intervals :
              Normal
                                   Basic
         (-1.0275, -0.8552)
                                (-1.0256, -0.8555)
##
  95%
##
## Level
                                    BCa
             Percentile
## 95%
         (-1.0252, -0.8551)
                                (-1.0260, -0.8556)
## Calculations and Intervals on Original Scale
```

plot(boot_sd)

Histogram of t



Randomization tests take the set of scores, randomize their ordering, and compute statistics from the results. Permutation tests do the same thing, but I reserve that label for tests in which we take all possible permutations of the data, rather than a subset or rearrangements. Bootstrapping resamples with replacement from a set of data and computes a statistic (such as the mean or median) on each resampled set. Bootstrapping is used primarily for parameter estimation. Bootstrapping is primarily focused on estimating population parameters, and it attempts to draw inferences about the population(s) from which the data came. Randomization approaches, on the other hand, are not particularly concerned about populations and/or their parameters. Instead, randomization procedures focus on the underlying mechanism that led to the data being distributed between groups in the way that they are (click here for a more detailed introduction).

Randomization Tests (Contingency Tables): (Fisher's Exact Test). Fisher's exact test is a statistical test used to determine if there are nonrandom associations between two categorical variables.

```
tbl = matrix(data=c(65, 45, 20, 40), nrow=2, ncol=2, byrow=T)
dimnames(tbl) = list(City=c('B', 'T'), Gender=c('M', 'F'))
tbl
##
        Gender
## City M F
##
      B 65 45
      T 20 40
##
chisq.test(tbl)
##
##
    Pearson's Chi-squared test with Yates' continuity correction
##
## X-squared = 9.2985, df = 1, p-value = 0.002293
fisher.test(tbl)
    Fisher's Exact Test for Count Data
##
##
## data: tbl
## p-value = 0.002162
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 1.427537 5.912307
## sample estimates:
## odds ratio
##
     2.870314
Odds ration: OR = \frac{65/20}{45/40} = \frac{65\times40}{45\times20}. The odds ratio shows how many times more positive cases (Male in city
B + Female NOT in B) occur than negative cases (Male NOT in B + Female in B).
Bootstrapping for correlation coefficients. Recall that we don't assume normality of variables to calculate
correlation coefficients (particularly Pearson's r). However, it might influence the significance test (or esti-
mation of confidence interval).
# generate two highly skewed variables
r1 = rweibull(1000, 0.5)
r2 = 0.5*r1 + rweibull(1000, 0.5)
data = data.frame(r1,r2)
```

```
# calculate the correlation coefficients
cor.test(r1,r2)

##
## Pearson's product-moment correlation
##
## data: r1 and r2
## t = 18.993, df = 998, p-value < 2.2e-16</pre>
```

```
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.4682172 0.5593796
## sample estimates:
         cor
## 0.5152544
cor.test(rank(r1),rank(r2))
##
  Pearson's product-moment correlation
##
##
## data: rank(r1) and rank(r2)
## t = 18.454, df = 998, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.4566869 0.5492191
## sample estimates:
##
         cor
## 0.5043997
```

Please note that the confidence interval of the rank correlation is more narrow than that of Pearson's correlation. Alternatively, we test the significance of the coefficients using bootstrapping (without assuming that how the sampling rs distributed).

```
# define the statistics:
rs_1 = function(d,i){
  d2=d[i,]
  r = cor(d2$r1,d2$r2,method="pearson")
  return(r)
}
rs_2 = function(d,i){
  d2=d[i,]
  r = cor(d2$r1,d2$r2,method="spearman")
  return(r)
# run bootstrapping
boot_r1 = boot(data,rs_1,R=1000) # define the data, define the test statistic, repetition times
boot_r2 = boot(data,rs_2,R=1000) # define the data, define the test statistic, repetition times
# obtain confidence interval
boot.ci(boot.out = boot_r1, type = c("norm", "basic", "perc", "bca"))
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 1000 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = boot_r1, type = c("norm", "basic", "perc",
       "bca"))
##
##
## Intervals :
```

```
Normal
                               Basic
## Level
## 95% ( 0.4086,  0.6275 ) ( 0.4113,  0.6223 )
## Level
          Percentile
                                BCa
       (0.4082, 0.6193) (0.4137, 0.6231)
## Calculations and Intervals on Original Scale
boot.ci(boot.out = boot_r2, type = c("norm", "basic", "perc", "bca"))
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 1000 bootstrap replicates
## CALL :
## boot.ci(boot.out = boot_r2, type = c("norm", "basic", "perc",
##
      "bca"))
##
## Intervals :
## Level Normal
                               Basic
## 95% ( 0.4542, 0.5554 ) ( 0.4559, 0.5584 )
## Level
          Percentile
                                BCa
## 95% ( 0.4504,  0.5529 ) ( 0.4513,  0.5535 )
## Calculations and Intervals on Original Scale
```

III. Regression (OLS)

7. Can regression models (why plural?) replace all the above-mentioned tests in section II?

The commonly used Ordinary Least Square (OLS) regression is just one of the many regression models, as we will explain the details in section IV. But as we have already known that Logistic Regression is different from OLS regression.

In section II, we introduced several statistics, basically comparing the difference of *means*: Z and T for two groups and F for more than two groups. In a regression framework, the dependent variable is a continuous variable, while the independent variable is a categorical variable.

Compare t.test with lm:

```
head(sleep) # load the sleep data
     extra group ID
##
## 1
       0.7
               1
                  1
     -1.6
## 2
               1
## 3 -0.2
                  3
               1
## 4 -1.2
               1 4
     -0.1
               1 5
## 5
## 6
       3.4
               1
t.test(extra~group,data=sleep,var.equal = T) # assume equal variance
##
##
   Two Sample t-test
## data: extra by group
## t = -1.8608, df = 18, p-value = 0.07919
## alternative hypothesis: true difference in means between group 1 and group 2 is not equal to 0
## 95 percent confidence interval:
## -3.363874 0.203874
## sample estimates:
## mean in group 1 mean in group 2
##
              0.75
                              2.33
summary(lm(extra~group,data=sleep))
##
## Call:
## lm(formula = extra ~ group, data = sleep)
##
## Residuals:
##
              1Q Median
                            3Q
                                  Max
## -2.430 -1.305 -0.580 1.455
                                3.170
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                 0.7500
                            0.6004
                                      1.249
                                              0.2276
                 1.5800
                            0.8491
                                      1.861
                                              0.0792 .
## group2
```

```
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## Residual standard error: 1.899 on 18 degrees of freedom
## Multiple R-squared: 0.1613, Adjusted R-squared: 0.1147
## F-statistic: 3.463 on 1 and 18 DF, p-value: 0.07919
Note that both t and p values are similar. In addition, the mean difference 2.33-0.75 = 1.58, which is the
regression coefficient.
How about the situation with more than 2 groups (using F-test in avo)?
head(npk)
##
     block N P K yield
## 1
         1 0 1 1 49.5
         1 1 1 0 62.8
## 2
## 3
         1 0 0 0 46.8
         1 1 0 1 57.0
## 4
## 5
         2 1 0 0 59.8
## 6
         2 1 1 1 58.5
aggregate(yield~block,npk,mean)
     block yield
## 1
         1 54.025
## 2
         2 57.450
## 3
         3 60.775
## 4
         4 50.125
         5 50.525
## 5
## 6
         6 56.350
summary(aov(yield~block,data=npk))
               Df Sum Sq Mean Sq F value Pr(>F)
##
## block
               5 343.3
                           68.66
                                  2.318 0.0861 .
## Residuals
               18 533.1
                           29.61
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
summary(lm(yield~block,data=npk))
##
```

Max

Call:

Residuals:

Min

Coefficients:

##

##

##

##

lm(formula = yield ~ block, data = npk)

1Q Median

-7.2250 -3.4937 -0.5375 2.1062 11.8750

3Q

Estimate Std. Error t value Pr(>|t|)

```
## (Intercept)
               54.025
                           2.721 19.855 1.09e-13 ***
## block2
                           3.848 0.890 0.3852
                3.425
## block3
                6.750
                           3.848
                                  1.754
                                          0.0964 .
## block4
               -3.900
                           3.848 -1.013
                                         0.3243
## block5
               -3.500
                           3.848 -0.910
                                          0.3751
                2.325
## block6
                           3.848
                                 0.604
                                         0.5532
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 5.442 on 18 degrees of freedom
## Multiple R-squared: 0.3917, Adjusted R-squared: 0.2228
## F-statistic: 2.318 on 5 and 18 DF, p-value: 0.08607
```

This is straightforward because 1m actually reported F-test statistics as well. They are exactly the same thing!

How about non-normal distribution?

3.574

```
# generate data
g1 = rpois(500,2.5)
g2 = rpois(500,3.5)
dv = c(g1,g2)
groups = c(rep(1,500),rep(0,500))
d = data.frame(dv,groups)

# compare the two outputs
t.test(dv~groups,d)

##
## Welch Two Sample t-test
##
## data: dv by groups
```

```
##
## data: dv by groups
## t = 9.7704, df = 967.5, p-value < 2.2e-16
## alternative hypothesis: true difference in means between group 0 and group 1 is not equal to 0
## 95 percent confidence interval:
## 0.8614805 1.2945195
## sample estimates:
## mean in group 0 mean in group 1</pre>
```

summary(lm(dv~groups,d))

```
##
## Call:
## lm(formula = dv ~ groups, data = d)
##
## Residuals:
## Min 1Q Median 3Q Max
## -3.574 -1.496 -0.496 1.426 6.504
##
## Coefficients:
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3.57400 0.07802 45.81 <2e-16 ***</pre>
```

2.496

```
## groups -1.07800 0.11033 -9.77 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.745 on 998 degrees of freedom
## Multiple R-squared: 0.0873, Adjusted R-squared: 0.08639
## F-statistic: 95.46 on 1 and 998 DF, p-value: < 2.2e-16</pre>
```

Testing for categorical variables (contingency tables) requires a regression model that we are less familiar with. We create a contingency table with two variables: city and gender. To test the dependence between the two, we use either χ^2 -test (parametric) or Fisher's exact test (non-parametric).

```
tbl = matrix(data=c(65, 45, 20, 40), nrow=2, ncol=2, byrow=T)
dimnames(tbl) = list(City=c('B', 'T'), Gender=c('M', 'F'))
tbl
##
       Gender
  City M F
##
      B 65 45
      T 20 40
chisq.test(tbl)
##
   Pearson's Chi-squared test with Yates' continuity correction
##
## data: tbl
## X-squared = 9.2985, df = 1, p-value = 0.002293
fisher.test(tbl)
##
##
   Fisher's Exact Test for Count Data
##
## data: tbl
## p-value = 0.002162
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 1.427537 5.912307
## sample estimates:
## odds ratio
     2.870314
##
```

Since this is just a 2 by 2 table, theoretically speaking, we can use logistic regression if we know the preaggregated data. We consider the frequencies in the contingency table the dependent variable and the city and gender as independent variables. Given the dependent variable is a count variable, as we will explain in section IV, we use a poisson regression to fit the data.

```
## log linear model
ctbl = data.frame(freq = c(65, 45, 20, 40), city = c("B","B","T","T"), gender = c("M","F","M","F"))
summary(logLM <- glm(freq ~city*gender, family=poisson(link="log"), data=ctbl))</pre>
```

```
##
## Call:
##
  glm(formula = freq ~ city * gender, family = poisson(link = "log"),
##
       data = ctbl)
##
## Deviance Residuals:
       0 0 0 0
## [1]
##
## Coefficients:
##
                 Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                   3.8067
                              0.1491 25.536
                                              < 2e-16 ***
                  -0.1178
                              0.2173
                                      -0.542
                                              0.58781
## cityT
## genderM
                   0.3677
                              0.1939
                                       1.896
                                              0.05793 .
## cityT:genderM -1.0609
                              0.3356
                                      -3.161
                                             0.00157 **
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
  (Dispersion parameter for poisson family taken to be 1)
##
##
      Null deviance: 2.5378e+01 on 3 degrees of freedom
## Residual deviance: 8.8818e-16 on 0 degrees of freedom
## AIC: 30.036
##
## Number of Fisher Scoring iterations: 3
```

The interaction coefficient (cityT:genderM = -1.0609) indicate the dependence between gender and city. In fact, log-linear is an extension of the χ^2 -test.

8. What is the degree of freedom? Is it possible to fit a regression model with $R^2=1$? What is over-fitting? Why should we avoid over-fitting?

Degrees of freedom (df) are the number of independent values that a statistical analysis can estimate. You can also think of it as the number of values that are free to vary as you estimate parameters. Typically, the degrees of freedom equals your sample size minus the number of parameters you need to calculate during an analysis (N-P). Degrees of freedom is a combination of how much data you have and how many parameters you need to estimate. It indicates how much independent information goes into a parameter estimate. In this vein, it's easy to see that you want a lot of information to go into parameter estimates to obtain more precise estimates and more powerful hypothesis tests. So, you want many df!

```
# generate a data sets with 3 IVs and y
x1 = rnorm(100,5,1)
x2 = rnorm(100,1,1)
x3 = rnorm(100,2,2)
y = 2*x1+1.5*x2+1.1*x3+rnorm(100) # sample size = 100
data = data.frame(y,x1,x2,x3)
# fit a linear model with 2 variables and find the df.
summary(lm(y~x1+x2,data = data)) # df = 100-3 (why 3 not 2?)
```

```
##
## Call:
## lm(formula = y ~ x1 + x2, data = data)
##
```

```
## Residuals:
##
      Min
                1Q Median
                                30
                                       Max
## -5.1770 -1.8477 -0.2214 1.7912 7.0916
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                 0.3079
                           1.2311
                                     0.250
                                              0.803
## x1
                 2.3772
                            0.2417
                                     9.837 3.02e-16 ***
## x2
                 1.8452
                            0.2345
                                     7.870 5.06e-12 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 2.556 on 97 degrees of freedom
## Multiple R-squared: 0.6253, Adjusted R-squared: 0.6176
## F-statistic: 80.94 on 2 and 97 DF, p-value: < 2.2e-16
```

Please note that the degree of freedom is 97, instead of 100-2 = 98. For a typical OLS regression, an intercept is estimated by default. How about if we only have 3 observations? Theoretically speaking, df = 3-3 = 0. As presented below, NAs were produced in 1m. Actually, a model with df = 0 is a just identified model with unique solution and the R^2 should be 1.

```
summary(lm(y~x1+x2,data = data[sample(100,3),])) # 3 observations
```

```
##
## Call:
## lm(formula = y \sim x1 + x2, data = data[sample(100, 3), ])
##
## Residuals:
## ALL 3 residuals are 0: no residual degrees of freedom!
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                   2.775
                                {\tt NaN}
                                         \mathtt{NaN}
                                                   NaN
## x1
                   1.413
                                NaN
                                         NaN
                                                   NaN
## x2
                   1.811
                                {\tt NaN}
                                         NaN
                                                   NaN
##
## Residual standard error: NaN on O degrees of freedom
## Multiple R-squared:
                             1, Adjusted R-squared:
## F-statistic: NaN on 2 and 0 DF, p-value: NA
summary(lm(y~x1+x2,data = data[sample(100,4),])) # 4 observations
##
## Call:
## lm(formula = y \sim x1 + x2, data = data[sample(100, 4), ])
##
## Residuals:
                 42
  -1.7637 3.2511 -0.7040 -0.7835
##
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
```

```
## (Intercept) -7.08236
                                              0.751
                          17.19098
                                    -0.412
## x1
                4.42213
                           4.04734
                                     1.093
                                              0.472
## x2
                           3.53052
               -0.01063
                                    -0.003
                                              0.998
##
## Residual standard error: 3.846 on 1 degrees of freedom
## Multiple R-squared: 0.689, Adjusted R-squared: 0.06689
## F-statistic: 1.108 on 2 and 1 DF, p-value: 0.5577
```

As $df \to 0$, the $R^2 \to 1$, even though it is not significant. The 1m function only works for df > 0. Nevertheless, we can solve the linear system mathematically when df = 0 (we cannot do that when df < 0).

```
library(matlib)

slice = data[sample(100,3),] # select the first 3 observations
A = as.matrix(slice[,2:3]) # IVs as matrix
b = slice$y # dependent variable
Solve(A, b) # solve the function
```

```
## x1 = 1.96005922
## x2 = 4.19579811
## 0 = 0.9158363
```

In statistics, over-fitting is "the production of an analysis that corresponds too closely or exactly to a particular set of data, and may therefore fail to fit additional data or predict future observations reliably". An over-fitted model is a statistical model that contains more parameters than can be justified by the data (i.e., small df). The essence of over-fitting is to have unknowingly extracted some of the residual variation (i.e., the noise) as if that variation represented underlying model structure.

How is this possible? In our case, we know how the data was generated according to $y = 2x_1 + 1.5x_2 + 1.1x_3 + \varepsilon$. If we measured all x_i , we will find a perfect solution. Let's include all three predictors:

```
summary(lm(y~x1+x2+x3,data))
```

```
##
## Call:
## lm(formula = y \sim x1 + x2 + x3, data = data)
##
## Residuals:
##
       Min
                1Q Median
                                 3Q
                                        Max
##
  -2.7058 -0.5030 -0.0408 0.4548
                                    2.6956
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.28883
                           0.48004
                                     -0.602
                                               0.549
## x1
                2.07023
                           0.09502
                                     21.788
                                              <2e-16 ***
                1.47369
                           0.09267
                                     15.903
                                              <2e-16 ***
## x2
                1.10295
                           0.04730
                                              <2e-16 ***
## x3
                                     23.319
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.9951 on 96 degrees of freedom
## Multiple R-squared: 0.9438, Adjusted R-squared: 0.942
## F-statistic: 537.2 on 3 and 96 DF, p-value: < 2.2e-16
```

The R^2 = is not 1, given the existence of $\varepsilon \sim \mathsf{Norm}(0,1)$. Otherwise, it is a deterministic models (in contrast to probabilistic). Let's generate y2 without the random error (ε) .

```
y2 = 2*x1+1.5*x2+1.1*x3
summary(lm(y2~x1+x2+x3))
## Warning in summary.lm(lm(y2 ~ x1 + x2 + x3)): essentially perfect fit: summary
## may be unreliable
##
## Call:
## lm(formula = y2 ~ x1 + x2 + x3)
## Residuals:
##
                     1Q
                            Median
                                            3Q
         Min
                                                     Max
## -1.967e-14 -4.798e-16 3.206e-16 1.006e-15 9.862e-15
##
## Coefficients:
##
               Estimate Std. Error
                                     t value Pr(>|t|)
## (Intercept) 0.000e+00 1.342e-15 0.000e+00
              2.000e+00 2.656e-16 7.530e+15
                                                <2e-16 ***
              1.500e+00 2.590e-16 5.791e+15
## x2
                                                <2e-16 ***
## x3
              1.100e+00 1.322e-16 8.320e+15
                                               <2e-16 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 2.782e-15 on 96 degrees of freedom
## Multiple R-squared:
                           1, Adjusted R-squared:
## F-statistic: 6.749e+31 on 3 and 96 DF, p-value: < 2.2e-16
```

Now, the R^2 is 1. Is this just-identified model (df = 0), a perfect model (deterministic), or over-fitted model? Let's show another $R^2 = 1$ example that it is an over-fitted model.

```
# create a categorical variable with 50 levels in the 'data'
data$v1 = as.factor(c(1:50,1:50))
summary(aov(x3~v1,data))
```

```
## v1 49 217.2 4.433 0.894 0.653 ## Residuals 50 248.0 4.961
```

The new variable is not associated with any x_i . Regress v1 on y:

```
# fit the first 50 observations
model1<-lm(y~v1,data[1:50,])
summary(model1)$r.squared</pre>
```

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

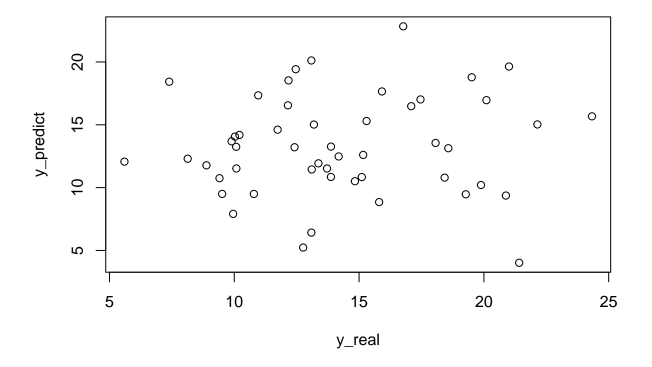
We can achieve $R^2 = 1$ without any information from x_i ! Is this a good model? Let's use "model1" to predict the rest of 50 ys in the data set.

```
# predict using other 50 observations (xs)
y_predict = predict(model1,data[51:100,])

# compare predicted and true values
y_real = y[51:100]
cor(y_real,y_predict)^2 #correlation between predicted and true values

## [1] 0.005262324

plot(y_real,y_predict)
```



Even though model is a perfect model for the first 50 observations, the model is not applicable to the second 50 observations. In this case, we say model is an over-fitted model.

9. How to interpret regression coefficients (direction, magnitude/strength, significance, form)? How to interpret when predictors are categorical? How to compare regression coefficients?

```
# to generate a data set with both continuous and categorical variables
x1 = rnorm(1000,10,2)
x2 = rnorm(1000,7,3)+0.5*x1
x3 = rbinom(1000,1,0.3)
x4 = sample(1:3, size = 1000, replace = TRUE, prob = c(0.2,0.5,0.3))
```

```
#dummy coding for x4
x41 = ifelse(x4==1,1,0)
x42 = ifelse(x4==2,1,0)
x43 = ifelse(x4==3,1,0)

# dependence between x and y
y = -2.2*x1+2.2*x2+3.3*x3+4*x42+8*x43+rnorm(1000)

# create the data frame
data = data.frame(y,x1,x2,x3,x4,x41,x42,x43)
```

Let's fit an OLS regression using 1m and try to interpret the coefficients according to the direction (positive or negative), magnitude (weak or strong), form (linear or nonlinear), and significance (significant or not).

```
summary(m <- lm(y~x1+x2+x3+factor(x4)))</pre>
```

```
##
## Call:
## lm(formula = y \sim x1 + x2 + x3 + factor(x4))
##
## Residuals:
       Min
                  1Q
                      Median
                                    3Q
                                            Max
## -3.12756 -0.62884 -0.05932 0.65987
                                        3.06148
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
                                      0.421
                                              0.674
## (Intercept) 0.07978
                          0.18953
## x1
               -2.20835
                           0.01684 -131.134
                                              <2e-16 ***
## x2
               2.19645
                           0.01077 203.922
                                             <2e-16 ***
## x3
               3.23421
                           0.06811
                                     47.486
                                              <2e-16 ***
## factor(x4)2 3.99200
                           0.08237
                                     48.464
                                              <2e-16 ***
## factor(x4)3 7.98965
                           0.09243
                                     86.440
                                             <2e-16 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 1.001 on 994 degrees of freedom
## Multiple R-squared: 0.9825, Adjusted R-squared: 0.9824
## F-statistic: 1.116e+04 on 5 and 994 DF, p-value: < 2.2e-16
```

The coefficient for x_3 indicates the difference of y between $x_3 = 1$ and $x_3 = 0$. Then, what does the coefficient for $x_4 = 3$ mean? Does it indicate the difference between $x_4 = 3$ and $x_4 = 1$ or $x_4 = 3$?

```
#calculate the means for each category
aggregate(y~x4,data,mean)
```

```
## x4 y
## 1 1 5.253108
## 2 2 9.603733
## 3 3 13.298502
```

Using standardized coefficients βs for comparisons. It is easy to calculate βs using the lm.beta function.

```
library(lm.beta)
betas = lm.beta(m)
betas
##
## Call:
## lm(formula = y \sim x1 + x2 + x3 + factor(x4))
##
## Standardized Coefficients::
## (Intercept)
                                                  x3 factor(x4)2 factor(x4)3
                         x1
                                      x2
     0.0000000
               -0.5824554
                              0.9046067
                                           0.1997161
                                                       0.2644106
```

We see that the absolute value of raw coefficient of x1 is close to that of x2, however, it is smaller in terms of the standardized coefficients. So, can we say $|\beta_{x_2}| - |\beta_{x_1}|$ and x_2 shows a greater impact on y? We still need a significance test, which requires the SE. The full version of the **Variance Sum Law**:

$$Var(c_1b_1 + c_2b_2) = c_1^2 Var(b_1) + c_2^2 Var(b_2) + 2c_1c_2 Cov(b_1, b_2)$$

We can obtain the variance-covariance matrix using the vcov function in R:

```
vc = vcov(m) # variance-covariance matrix
vc
```

```
##
                 (Intercept)
                                                     x2
                                                                   xЗ
                                       x1
## (Intercept) 0.0359229446 -2.118473e-03 -8.013656e-04 -8.188163e-04
## x1
              -0.0021184735 2.835980e-04 -5.871701e-05 -6.086019e-05
## x2
               -0.0008013656 -5.871701e-05 1.160144e-04 8.038949e-07
## x3
              -0.0008188163 -6.086019e-05 8.038949e-07 4.638892e-03
## factor(x4)2 -0.0047362111 8.483126e-06 -1.402981e-05 -1.364965e-04
## factor(x4)3 -0.0048513610 2.936769e-06 -4.638961e-06 5.396930e-05
                factor(x4)2
                              factor(x4)3
## (Intercept) -4.736211e-03 -4.851361e-03
## x1
               8.483126e-06 2.936769e-06
              -1.402981e-05 -4.638961e-06
## x2
## x3
              -1.364965e-04 5.396930e-05
## factor(x4)2 6.785022e-03 4.859963e-03
## factor(x4)3 4.859963e-03 8.543299e-03
```

In this case, $SE_{b_{x_2}-b_{x_1}} = \sqrt{Var_{x_1} + Var_{x_2} + 2Cov(x_1, x_2)}$. Therefore, the standard error of the difference could be calculated as below:

```
# the difference between the two unstandardized coefficients
diff = abs(m$coefficients['x2'])-abs(m$coefficients['x1'])
var_x1 = vc[2,2] # variance
var_x2 = vc[3,3] # variance
cov_x1x2 = vc[3,2] # covariance
se_diff = sqrt(var_x1+var_x2+2*cov_x1x2) # standard error of the difference
z = diff/se_diff # z score
names(z) = NULL
z
```

```
## [1] -0.708288
```

It is less than 1.96, so it is not significant. We can use the following code to obtain the p value:

```
pvalue = pnorm(-abs(z))
pvalue
```

[1] 0.2393832

We can do something even more interesting, such as to test whether $b_{x_1} > 2.2$.

```
diff=abs(m$coefficients['x1'])-2.2
var_x1 = vc[2,2]
var_2.2 = 0
cov_x12.2 = 0
se_diff = sqrt(var_x1+var_2.2+2*cov_x12.2)
z = diff/se_diff
names(z) = NULL
pvalue = pnorm(-abs(z))
c(z,pvalue)
```

[1] 0.4955377 0.3101103

Whether $b_{x_4=3} > 2b_{x_4=2}$?

```
diff=abs(m$coefficients['factor(x4)3']-2*m$coefficients['factor(x4)2'])
var_x43 = vc[6,6]
var_x42 = vc[5,5]
cov_x4 = vc[5,6]
se_diff = sqrt(var_x43+4*var_x42+4*cov_x4)
z = diff/se_diff
names(z) = NULL
pvalue = pnorm(-abs(z))
c(z,pvalue)
```

[1] 0.02404922 0.49040668

Will standardization of variables change the variance-covariance matrix?

```
summary(m1<-lm(y~x1+x2,data = data))</pre>
```

```
##
## Call:
## lm(formula = y \sim x1 + x2, data = data)
## Residuals:
               1Q Median
##
      Min
                               3Q
                                      Max
## -7.7747 -2.1571 -0.2651 2.3975 8.4230
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.93884 0.58630 8.424
                                            <2e-16 ***
              -2.16880 0.05509 -39.370
## x1
                                            <2e-16 ***
```

```
## x2
               2.19897
                          0.03528 62.331 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 3.278 on 997 degrees of freedom
## Multiple R-squared: 0.8116, Adjusted R-squared: 0.8113
## F-statistic: 2148 on 2 and 997 DF, p-value: < 2.2e-16
# standardized all variables
y_s = (data$y-mean(data$y))/sd(data$y)
x1_s = (data$x1-mean(data$x1))/sd(data$x1)
x2_s = (data$x2-mean(data$x2))/sd(data$x2)
summary(m2<-lm( y_s ~ x1_s+x2_s-1)) # without intercept</pre>
##
## Call:
## lm(formula = y_s ~ x1_s + x2_s - 1)
##
## Residuals:
##
                 1Q Median
       Min
                                   3Q
## -1.03040 -0.28588 -0.03513 0.31775 1.11631
##
## Coefficients:
##
       Estimate Std. Error t value Pr(>|t|)
## x1_s -0.57203 0.01452 -39.39
                                   <2e-16 ***
## x2 s 0.90565
                   0.01452
                             62.36
                                     <2e-16 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 0.4342 on 998 degrees of freedom
## Multiple R-squared: 0.8116, Adjusted R-squared: 0.8113
## F-statistic: 2150 on 2 and 998 DF, p-value: < 2.2e-16
lm.beta(m2)
##
## Call:
## lm(formula = y_s ~ x1_s + x2_s - 1)
##
## Standardized Coefficients::
##
        x1 s
                   x2 s
## -0.5720259 0.9056471
```

The coefficients in the regression model based on the standardized variables (m2) are equal to the standardized coefficients. Check the variance-covariance matrices:

```
vcov(m1)
```

```
## (Intercept) x1 x2
## (Intercept) 0.343748920 -0.0228019769 -0.0086871959
## x1 -0.022801977 0.0030346850 -0.0006298252
## x2 -0.008687196 -0.0006298252 0.0012445918
```

vcov(m2)

```
## x1_s x2_s
## x1_s 2.108966e-04 -6.834699e-05
## x2_s -6.834699e-05 2.108966e-04
```

They are different. Any implications?

10. Why the I.I.D. assumption is essential?

A regression model could be write as a combination of two parts: random part + systematic part.

$$Y_i \sim \mathsf{Norm}(\mu_i, \sigma)$$

 $\mu_i = X_i \beta_1 + \beta_0$

I.I.D refers to independent and identical distribution.

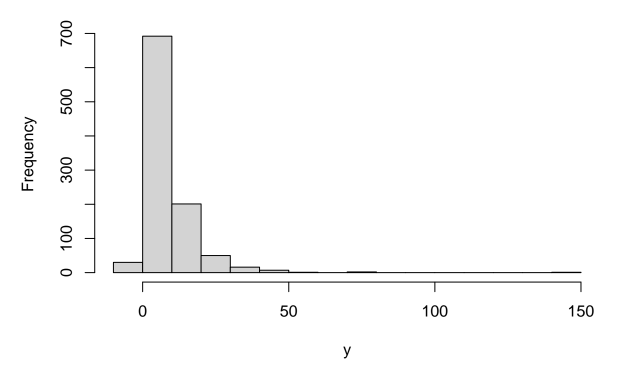
- Each draw from the bag must be independent. This means that the value you get on one draw does not depend in any way on other draws. Not repeated measures.
- Each observation is drawing from the same bag. In other words, you are drawing values from the same distribution. The shape of this distribution does not matter, even though some people will claim that it must be normally distributed. However, regardless of the shape, all observations must be drawing from an identically shaped distribution.

11. What if residuals are normally distributed, but the dependent variable is not? Is this possible?

Let's create a variable y that is determined by two skewed variables and a normally distributed random error.

```
x1 = rexp(1000,1.5)
x2 = rweibull(1000,0.5)
y = 7*x1+2*x2+rnorm(1000)
hist(y)
```

Histogram of y



summary(lm(y~x1+x2))

```
##
## Call:
  lm(formula = y \sim x1 + x2)
##
##
## Residuals:
##
        Min
                  1Q
                       Median
                                     3Q
                                             Max
  -2.65615 -0.67830
                      0.01426
                               0.59607
                                         2.89016
##
##
##
   Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
   (Intercept) 0.020544
                                               0.654
##
                           0.045881
                                      0.448
## x1
               7.001456
                           0.046065 151.990
                                              <2e-16 ***
                           0.007674 261.478
## x2
               2.006464
                                              <2e-16 ***
## ---
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 0.9783 on 997 degrees of freedom
## Multiple R-squared: 0.9891, Adjusted R-squared: 0.989
## F-statistic: 4.507e+04 on 2 and 997 DF, p-value: < 2.2e-16
```

As you can see, the dependent variable is skewed while the OLS regression estimated the coefficients correctly. The assumption is not about the shape of the dependent variable but how the random error of generating the dependent variable (the random part in the generation process).

12. Why effect size is important? Is it true that larger coefficients indicate larger effect sizes? How to measure the unique effect of an independent variable on the dependent variable without any confounding effects of other independent variables (or can stepwise regression models solve the problem)?

```
summary(m)
##
## Call:
## lm(formula = y \sim x1 + x2 + x3 + factor(x4))
##
## Residuals:
##
                  1Q
                      Median
                                    3Q
       Min
                                             Max
  -3.12756 -0.62884 -0.05932 0.65987
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.07978
                           0.18953
                                      0.421
                                               0.674
               -2.20835
                           0.01684 -131.134
                                              <2e-16 ***
## x1
                           0.01077
## x2
                2.19645
                                    203.922
                                              <2e-16 ***
                3.23421
                           0.06811
                                     47.486
## x3
                                              <2e-16 ***
## factor(x4)2 3.99200
                           0.08237
                                     48.464
                                               <2e-16 ***
## factor(x4)3 7.98965
                           0.09243
                                     86.440
                                              <2e-16 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 1.001 on 994 degrees of freedom
## Multiple R-squared: 0.9825, Adjusted R-squared: 0.9824
## F-statistic: 1.116e+04 on 5 and 994 DF, p-value: < 2.2e-16
betas
##
## Call:
## lm(formula = y \sim x1 + x2 + x3 + factor(x4))
##
## Standardized Coefficients::
  (Intercept)
                                                 x3 factor(x4)2 factor(x4)3
                        x1
                                    x2
     0.0000000 -0.5824554
                             0.9046067
                                         0.1997161
##
                                                      0.2644106
                                                                  0.4714295
```

There are many effect size measures. β could be used as effect size, but it is really hard to interpret. In regression models, R^2 is most intuitive (variance explained), but it is a overall measure.

Squared Semi-partial correlation (sr^2) tells us how much of the unique contribution of an independent variable to the total variation in dependent variable. In other words, it explains increment in R-square (ΔR^2) when an independent variable is added.

```
a = summary(lm(y~x3+factor(x4),data))$r.squared #0.18
b = summary(lm(y~x1+x3+factor(x4),data))$r.squared #0.27
deta = b-a
deta
```

```
## [1] 0.08340707
```

Therefore, x_1 increased R^2 by 0.0834071?

```
c = summary(lm(y~x1+x2+x3+factor(x4),data))$r.squared #0.98 deta2 = c-b deta2
```

[1] 0.7321488

Therefore, x_2 increased R^2 by 0.7321488? Why is this method problematic? Let's change the input order...

```
cor.test(data$x1,data$x2)
```

```
##
## Pearson's product-moment correlation
##
## data: data$x1 and data$x2
## t = 10.822, df = 998, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.2674585 0.3784676
## sample estimates:
## cor
## 0.3240781

b_ = summary(lm(y~x2+x3+factor(x4),data))$r.squared
b -a # variance x2 explained</pre>
```

```
## [1] 0.5127946
```

```
c - b_ # variance x1 explained
```

```
## [1] 0.3027613
```

It's a different story, which suggests that x_1 increased R^2 by 0.3027613, while x_2 R^2 by 0.5127946. The correct way to calculate the **Squared Semi-partial correlation** (sr^2) is to use the "one-less" approach:

- fit the full model with all predictors and calculate the overall R_f^2ull
- remove one predictor (x_i) from the full model and then fit the "one-less" model
- calculate the $R^2_{(i)}$ for the "one-less" model and $\Delta R^2_i = R^2_f u l l R^2_{(i)}$

```
# total
ovall = summary(lm(y~x1+x2+x3+factor(x4),data))$r.squared

# one less by removing x1 from m
ovall - summary(lm(y~x2+x3+factor(x4),data))$r.squared
```

```
## [1] 0.3027613
```

```
# one less by removing x2 from m
ovall - summary(lm(y~x1+x3+factor(x4),data))$r.squared

## [1] 0.7321488

# one less by removing x3 from m
ovall - summary(lm(y~x1+x2+factor(x4),data))$r.squared
```

[1] 0.03970017

```
# one less by removing x4 from m
ovall - summary(lm(y~x1+x2+x3,data))$r.squared
```

[1] 0.132892

In fact, x_1 can explain 0.3027613 variance. Why is the sum of all R^2s larger than 100%?

13. Is it possible to estimate the main effects from a regression model with interaction terms?

Let's fit a model with interaction:

```
summary(full <- lm(y~x1*x3+x2,data))</pre>
```

```
##
## lm(formula = y \sim x1 * x3 + x2, data = data)
##
## Residuals:
               1Q Median
      Min
                               3Q
                                      Max
## -7.0299 -1.5767 -0.1398 2.3720 6.7235
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.91352
                          0.62005
                                    7.924 6.12e-15 ***
## x1
              -2.26550
                          0.05881 -38.521 < 2e-16 ***
## x3
               1.43199
                          1.02276
                                    1.400 0.1618
               2.19767
## x2
                          0.03152 69.723 < 2e-16 ***
## x1:x3
               0.17119
                          0.09934
                                    1.723
                                           0.0852 .
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 2.928 on 995 degrees of freedom
## Multiple R-squared: 0.8501, Adjusted R-squared: 0.8495
## F-statistic: 1410 on 4 and 995 DF, p-value: < 2.2e-16
```

Can we say the main effect of x1 is -2.27?

When $x_3 = 0$:

```
summary(lm(y~x1*x3+x2,data[data$x3==0,]))
##
## Call:
## lm(formula = y \sim x1 * x3 + x2, data = data[data$x3 == 0, ])
##
## Residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
## -6.4804 -1.6330 -0.1946 2.5007
                                   6.7126
## Coefficients: (2 not defined because of singularities)
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.99334
                           0.64466
                                     7.746 3.47e-14 ***
               -2.26027
                           0.06017 -37.564 < 2e-16 ***
## x1
## x3
                     NA
                                NA
                                        NA
                                                 NA
                2.18670
                           0.03853
                                            < 2e-16 ***
## x2
                                    56.747
## x1:x3
                     NA
                                NA
                                        NA
                                                 NA
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 2.95 on 679 degrees of freedom
## Multiple R-squared: 0.8439, Adjusted R-squared: 0.8435
## F-statistic: 1836 on 2 and 679 DF, p-value: < 2.2e-16
When x_3 = 1,
summary(lm(y~x1*x3+x2,data[data$x3==1,]))
## Call:
## lm(formula = y \sim x1 * x3 + x2, data = data[data$x3 == 1,])
## Residuals:
##
      Min
                1Q Median
## -6.9305 -1.4577 -0.1049 1.9112
                                   6.1695
##
## Coefficients: (2 not defined because of singularities)
              Estimate Std. Error t value Pr(>|t|)
                                     6.874 3.36e-11 ***
## (Intercept) 6.19746
                           0.90158
## x1
               -2.10749
                           0.08594 -24.523 < 2e-16 ***
## x3
                     NΑ
                                NA
                                        NA
                                                 NΑ
## x2
                2.22094
                           0.05480
                                    40.526
                                            < 2e-16 ***
## x1:x3
                     NA
                                NA
                                        NA
                                                 NA
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 2.882 on 315 degrees of freedom
## Multiple R-squared: 0.8478, Adjusted R-squared: 0.8468
## F-statistic: 877.4 on 2 and 315 DF, p-value: < 2.2e-16
```

It demonstrates that it is not the main effect but the effect when $x_3 = 0$. The main effects could be obtained in the following ways (the real value is in between):

```
# margins based on the full model
library(margins)
margins(full)
##
       x1
             x3
                   x2
   -2.211 3.146 2.198
# fit a model without interaction term
summary(lm(y~x1+x2+x3,data))
##
## Call:
## lm(formula = y \sim x1 + x2 + x3, data = data)
##
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -6.9394 -1.6071 -0.0995 2.3721 6.6387
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.34488
                        0.52548
                                    8.268 4.31e-16 ***
               -2.21015
                          0.04932 -44.816 < 2e-16 ***
## x1
## x2
               2.19922
                          0.03154 69.731 < 2e-16 ***
               3.16074
                          0.19930 15.859 < 2e-16 ***
## x3
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 2.931 on 996 degrees of freedom
## Multiple R-squared: 0.8496, Adjusted R-squared: 0.8492
## F-statistic: 1876 on 3 and 996 DF, p-value: < 2.2e-16
```

The marginal effect represents the difference of (two) predictions for an (infinitesimal) change in x (the focal term). The average marginal effect represents the average slope of that predictor. In other words: the average marginal effects is one value per parameter (term), thus it can be considered as an **adjusted regression coefficient**, while predicted values usually predict the average outcome for different values of x - you usually don't have just one coefficient in the latter case that represents the overall effect of x.

IV. Regression (GLM)

14. How to deal with non-normally distributed dependent variables? How to interpret the coefficients?

As mentioned in section III, there are many regression models in addition to OLS regression. Most regression models could be expressed as the combination of a random plus a systematic part to generate the dependent variable. As we presented in Question 10, the dependent variable in OLS regression is generated according to a normal distribution and the *mean* varies with other predictors systematically. In this sense, as long as we know the generating process (or underlying distribution) of the dependent variable, we can model the data in a regression form. We call them Generalized Regression Models (GLMs). Some commonly used models include:

- Logistic Regression (DV is a binary variable, which is generated from a Bernoulli distribution)
- Poisson Regression (DV is a count variable, which is generated from a Poisson distribution)
- Negative Binomial (DV is a over-dispersed count variable, which is generated from a Negative Binomial distribution)
- Beta Regression (DV is a percentage/proportion variable, which is generated from a Beta distribution)

Logistic Regression

$$Y_i \sim \mathsf{Bern}(\pi_i)$$

$$\mathsf{logit}(\pi_i) = \log(\frac{\pi_i}{1-\pi_i}) = X_i\beta_1 + \beta_0$$

In logistic regression, the dependent variable Y_i is generated from a Bernoulli distribution with a single parameter π_i , which is the probability of occurrence (the event of interest). The **link function** is logit (not log). It connects the predictors X_i to the parameter π_i .

 β_0 is the log odds of the event of interest, when $X_i = 0$, e^{β_0} is the odds. $\beta_1 = \log(odds_{x+1} - \log(odds_x))$, or $e^{\beta_1} = \frac{odds_{x+1}}{odds_x}$. It reflects the change in log odds.

```
# load the admission data for an example
mydata <- read.csv("https://stats.idre.ucla.edu/stat/data/binary.csv")
head(mydata)</pre>
```

```
##
     admit gre gpa rank
## 1
         0 380 3.61
## 2
         1 660 3.67
                         3
## 3
         1 800 4.00
## 4
         1 640 3.19
                         4
## 5
         0 520 2.93
                         4
         1 760 3.00
```

The glm function in R can fit most generalized linear models. For logistic regression, we set the family = "binomial":

```
library(jtools)
mylogit <- glm(admit ~ gre + gpa + factor(rank), data = mydata, family = "binomial")
jtools::summ(mylogit)</pre>
```

Observations	400
Dependent variable	admit
Type	Generalized linear model
Family	binomial
Link	logit

$\chi^{2}(5)$	41.46
Pseudo-R ² (Cragg-Uhler)	0.14
Pseudo-R ² (McFadden)	0.08
AIC	470.52
BIC	494.47

	Est.	S.E.	z val.	p
(Intercept)	-3.99	1.14	-3.50	0.00
gre	0.00	0.00	2.07	0.04
gpa	0.80	0.33	2.42	0.02
factor(rank)2	-0.68	0.32	-2.13	0.03
factor(rank)3	-1.34	0.35	-3.88	0.00
factor(rank)4	-1.55	0.42	-3.71	0.00

Standard errors: MLE

The jtools::summ function are extremely useful when you want to obtain the *Pseudo-R*². Since the relationships in logistic regression are non-linear. Usually, they are less intuitive to be interpreted. We can use the margins and ggpredict to estimate the predicted probabilities, which will help us to interpret the results.

```
library(margins)
library(ggeffects)

margins(mylogit, variables = "rank")

## rank2 rank3 rank4
## -0.1566 -0.2872 -0.3212

ggpredict(mylogit, terms = "rank")

## # Predicted probabilities of admit
```

```
##
## rank | Predicted |
                             95% CI
##
## 1
               0.51 | [0.38, 0.64]
## 2
               0.35 | [0.27, 0.43]
               0.22 | [0.15, 0.30]
## 3
               0.18 | [0.11, 0.30]
## 4
## Adjusted for:
## * gre = 580.00
## * gpa =
           3.39
```

For example, the marginal effects report that compare to rank 1, rank2's probability to be admitted is 15.7% lower.

Poisson Regression

$$Y_i \sim \mathsf{Pois}(\lambda_i)$$

$$\log(\lambda_i) = X_i \beta_1 + \beta_0$$

 β_0 is the logged average of Y, when $X_i = 0$. e^{β_0} is the average of Y. $\beta_1 = \log(\lambda_{x+1}) - \log(\lambda_x)$, or $e^{\beta_1} = \frac{\lambda_{x+1}}{\lambda_x}$.

head(warpbreaks)

##		breaks	wool	tension
##	1	26	Α	L
##	2	30	Α	L
##	3	54	Α	L
##	4	25	Α	L
##	5	70	Α	L
##	6	52	Α	L

mypois <-glm(breaks ~ wool+tension, data = warpbreaks,family = poisson)
summ(mypois)</pre>

Observations	54
Dependent variable	breaks
Type	Generalized linear model
Family	poisson
Link	\log

$\chi^2(3)$	86.98
Pseudo-R ² (Cragg-Uhler)	0.80
Pseudo-R ² (McFadden)	0.15
AIC	493.06
BIC	501.01

	Est.	S.E.	z val.	p
(Intercept)	3.69	0.05	81.30	0.00
woolB	-0.21	0.05	-3.99	0.00
tensionM	-0.32	0.06	-5.33	0.00
tensionH	-0.52	0.06	-8.11	0.00

Standard errors: MLE

Negative Binomial:

$$Y_i \sim \mathsf{NegBin}(\mu_i, r)$$

$$\log(\mu_i) = X_i \beta_1 + \beta_0$$

Both Poisson regression and Negative Binomial regression are models for count data. Therefore, the interpretation of coefficients of Negative Binomial is similar to that of Poisson regression. The difference is that E(Y) = Var(Y) in Poisson distribution, while $E(Y) \neq Var(Y)$ in Negative Binomial distribution. This

is governed by the dispersion parameter r. For large reciprocal dispersion parameter r, $Var(Y) \approx E(Y)$, NegBin \rightarrow Pois. The Negative Binomial distribution is especially useful to model highly skewed data.

Unfortunately, we cannot use glm to fit a Negative Binomial model. Instead, we need to use glm.nb function from the MASS package:

```
library(MASS)
library(haven) # to read stata data
dat <- read_stata("https://stats.idre.ucla.edu/stat/stata/dae/nb_data.dta")</pre>
head(dat)
## # A tibble: 6 x 5
##
        id
               gender math daysabs prog
##
     <dbl>
            <dbl+1bl> <dbl>
                               <dbl> <dbl>
## 1
      1001 2 [male]
                          63
                                    4
                                          2
## 2
      1002 2 [male]
                          27
                                    4
                                          2
                                    2
      1003 1 [female]
                          20
```

2

2

2

3

3

13

hist(dat\$daysabs)

1004 1 [female]

1005 1 [female]

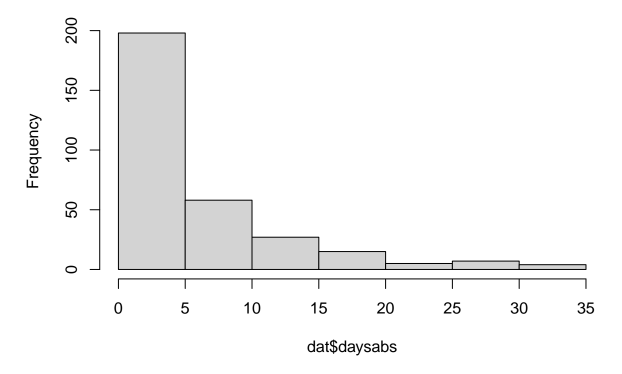
1006 1 [female]

16

2

71

Histogram of dat\$daysabs



```
summ(mynegbin <- glm.nb(daysabs ~ math + prog, data = dat))</pre>
```

Observations	314
Dependent variable	daysabs
Type	Generalized linear model
Family	Negative $Binomial(1.0222)$
Link	\log

$\chi^2()$	0.17	0.03	1741.52	1756.51
Pseudo-R ² (Cragg-Uhler)	0.17	0.03	1741.52	1756.51
Pseudo-R ² (McFadden)		0.03	1741.52	1756.51
AIC	0.17	0.03	1741.52	1756.51
BIC	0.17	0.03	1741.52	1756.51

	Est.	S.E.	z val.	р
(Intercept)	3.49	0.23	15.48	0.00
math	-0.01	0.00	-2.72	0.01
prog	-0.68	0.10	-6.98	0.00

Standard errors: MLE

The theta parameter estimated above is the dispersion parameter. Note that R parameterizes this differently from SAS, Stata, and SPSS. The R parameter (theta) is equal to the inverse of the dispersion parameter estimated in these other software packages. Thus, the theta value of 1.022 seen here is equivalent to the 0.978 because 1/1.022 = 0.978.

Beta Regression

$$Y_i \sim \mathsf{Beta}(\mu_i, \phi)$$

$$\mathsf{logit}(\mu_i) = \log(\frac{\mu_i}{1 - \mu_i}) = X_i \beta_1 + \beta_0$$

Beta regression also uses logit as the link function. The distribution ranges from 0 to 1 (0 and 1 are not usually excluded). Therefore, it is useful to model percent and proportion data. We use **betareg** to fit the model. The parameter ϕ is a precision parameter: the higher ϕ the lower the variance for given mean μ .

```
library(betareg)
data("FoodExpenditure", package = "betareg")
head(FoodExpenditure)
```

```
# food/income is a proportion
mybeta <- betareg(I(food/income) ~ income + persons, data = FoodExpenditure)
summary(mybeta)</pre>
```

```
## Call:
## betareg(formula = I(food/income) ~ income + persons, data = FoodExpenditure)
## Standardized weighted residuals 2:
##
                1Q Median
                                3Q
                                       Max
  -2.7818 -0.4445 0.2024
                           0.6852 1.8755
##
## Coefficients (mean model with logit link):
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.622548
                           0.223854 -2.781 0.005418 **
## income
               -0.012299
                           0.003036
                                    -4.052 5.09e-05 ***
                                      3.352 0.000802 ***
                0.118462
                           0.035341
## persons
##
## Phi coefficients (precision model with identity link):
         Estimate Std. Error z value Pr(>|z|)
##
## (phi)
            35.61
                        8.08
                               4.407 1.05e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Type of estimator: ML (maximum likelihood)
## Log-likelihood: 45.33 on 4 Df
## Pseudo R-squared: 0.3878
## Number of iterations: 28 (BFGS) + 4 (Fisher scoring)
margins(mybeta)
## Average marginal effects
## betareg(formula = I(food/income) ~ income + persons, data = FoodExpenditure)
      income persons
   -0.00248 0.02389
##
```

Therefore, one more person leads to 2.4% increase in food consumption given the income (food/income).

15. Can we model the variance (instead of mean) of the dependent variable? E.g., the variance of the salary of elder people is smaller than the variance of the salary of young people, given equal salary.

Recall the expression for normal regression. We simply change the systematic part by including σ :

$$Y_i \sim \mathsf{Norm}(\mu, \sigma_i)$$

 $\sigma_i = X_i \beta_1 + \beta_0$

In fact, we can model both μ_i and σ_i simultaneously. We can fit the models using the gamlss (Generalized Additive Models for Location, Scale and Shape) package in R (see more example here).

```
library(gamlss)
head(mtcars)
```

```
##
                     mpg cyl disp hp drat
                                             wt qsec vs am gear carb
## Mazda RX4
                           6 160 110 3.90 2.620 16.46
                                                       0
                    21.0
## Mazda RX4 Wag
                    21.0
                           6 160 110 3.90 2.875 17.02
## Datsun 710
                    22.8
                           4 108 93 3.85 2.320 18.61
                                                                    1
## Hornet 4 Drive
                    21.4
                           6
                              258 110 3.08 3.215 19.44
                                                       1
                                                                    1
                                                                    2
## Hornet Sportabout 18.7
                           8 360 175 3.15 3.440 17.02 0
                                                         0
                           6 225 105 2.76 3.460 20.22 1 0
## Valiant
                    18.1
```

Let's fit an OLS regression model:

```
summ(lm(mpg~hp+wt,mtcars),model.info = FALSE, model.fit = FALSE)
```

	Est.	S.E.	t val.	p
(Intercept)	37.23	1.60	23.28	0.00
hp	-0.03	0.01	-3.52	0.00
wt	-3.88	0.63	-6.13	0.00

Standard errors: OLS

Model σ using gamlss. There are two formulas, one for the *mean* and another for the *standard deviation*. Set the first part as in lm. If you really believe that the *mean* is not related to other variables, ~1 indicates that only the intercept is included in the model (thus is a constant). The second part is sigma.formula, using =~ to specify the predictors. You can specify the family, which means you can go beyond Normal distributions. Here, family = NO(), indicating a Normal distribution.

```
## Family: c("NO", "Normal")
##
## Call: gamlss(formula = mpg ~ 1, sigma.formula = ~hp + wt,
##
      family = NO(), data = mtcars, control = gamlss.control(trace = FALSE))
##
## Fitting method: RS()
##
## Mu link function: identity
## Mu Coefficients:
##
            Estimate Std. Error t value Pr(>|t|)
## (Intercept) 15.0982
                        0.3599
                               41.95 <2e-16 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Sigma link function: log
## Sigma Coefficients:
```

```
##
           Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3.356603 0.355505 9.442 3.37e-10 ***
          ## wt
           0.060041 0.146335 0.410 0.684713
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## -----
## No. of observations in the fit: 32
## Degrees of Freedom for the fit: 4
      Residual Deg. of Freedom: 28
##
                  at cycle: 13
##
## Global Deviance:
                 192.8026
##
           AIC:
                 200.8026
##
           SBC:
                  206.6656
summary(myglass2)
## Family: c("NO", "Normal")
##
## Call: gamlss(formula = mpg ~ hp + wt, sigma.formula = ~hp +
     wt, family = NO(), data = mtcars, control = gamlss.control(trace = FALSE))
##
##
## Fitting method: RS()
## -----
## Mu link function: identity
## Mu Coefficients:
          Estimate Std. Error t value Pr(>|t|)
## (Intercept) 35.97612 2.47576 14.531 5.40e-14 ***
                  0.00713 -3.901 0.000604 ***
## hp
          -0.02782
## wt
          -3.69296
                  0.79155 -4.666 8.13e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Sigma link function: log
## Sigma Coefficients:
           Estimate Std. Error t value Pr(>|t|)
## (Intercept) 1.224105 0.545910 2.242 0.0337 *
## hp
          -0.003228
                   0.002775 -1.163 0.2553
           0.039559 0.224216 0.176 0.8613
## wt
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## -----
## No. of observations in the fit: 32
## Degrees of Freedom for the fit: 6
     Residual Deg. of Freedom: 26
##
                  at cycle: 8
```

```
## Global Deviance: 146.998
## AIC: 158.998
## SBC: 167.7924
```

Another (maybe more intuitive/flexible) way is to use Bayes approach. It is easy to set the model using the brms packages in R. The code turns out to be very slow. Please use at your own risk.

```
# library(brms)
# myBayes <- brm(bf(mpg ~ hp + wt, sigma ~ hp+wt),
# data = mtcars,
# family = gaussian)
# summary(myBayes)</pre>
```

16. What is wrong with using Z-statistics (and associated p-values) of the coefficient of a multiplicative term to test for a statistical interaction in nonlinear models with categorical dependent variables (e.g., logistic regression)?

In linear models, the interaction term tests the difference between the two slopes. However, interactions in non-linear models (e.g., logistic regression) could be more complicated. Critically, in binary probit and logit, the equality of regression coefficients across groups does not imply that the marginal effects of a predictor on the probability are equal (Long & Mustillo, 2021).

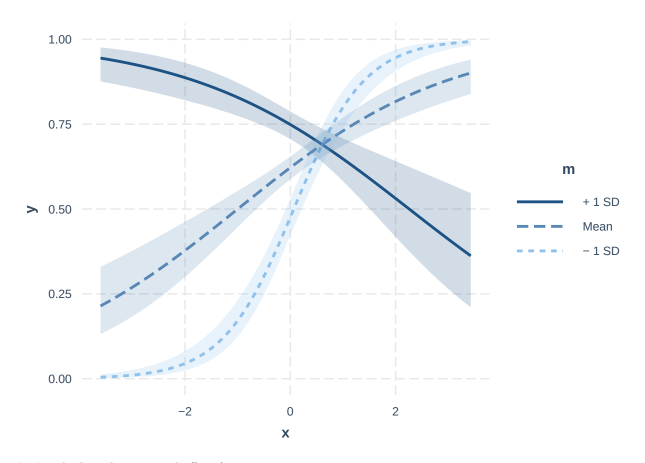
```
# generate a data set with binary y
x <- rnorm(1000)
m <- rnorm(1000)
prob <- binomial(link = "logit")$linkinv(.25 + .3*x + .3*m + -.5*(x*m) + rnorm(1000))
y <- rep(0, 1000)
y[prob >= .5] <- 1
summ(logit_fit <- glm(y ~ x * m, family = binomial), model.info = FALSE, model.fit = FALSE)</pre>
```

	Est.	S.E.	z val.	р
(Intercept)	0.50	0.07	6.93	0.00
x	0.49	0.08	6.12	0.00
m	0.60	0.09	7.08	0.00
x:m	-1.00	0.11	-9.47	0.00

Standard errors: MLE

The interaction term x:m is significant. However, due to the non-linear (log) transformation, the slope differs at different values for x, thus, the **marginal effect** or **association** (in terms of probabilities) is not constant across values of x. Let's plot out the interaction effect:

```
library(interactions)
interact_plot(logit_fit, pred = x, modx = m, interval = T)
```



Let's calculate the marginal effect for x:

```
summary(ef <- margins(logit_fit))</pre>
```

```
## factor AME SE z p lower upper

## m 0.1082 0.0124 8.7119 0.0000 0.0838 0.1325

## x 0.0902 0.0120 7.5301 0.0000 0.0667 0.1137
```

On average, a unit-change in x changes the predicted probability that the outcome equals 1 by 0.09 (see here).

It might be less intuitive to interpret average marginal effects, in particular for non-Gaussian models, because it is harder to understand an average effect where we actually have varying effects across the range of the focal term. Instead, it would be better to look at predictions at different values of the focal term(s), which is what ggeffects returns by default:

```
ggpredict(logit_fit, "x")
```

```
## # Predicted probabilities of y
##
## x | Predicted | 95% CI
## ------
## -4 | 0.18 | [0.10, 0.30]
## -3 | 0.27 | [0.18, 0.38]
## -2 | 0.38 | [0.30, 0.46]
```

```
## -1 |
              0.50 \mid [0.45, 0.55]
##
    0 1
              0.62 | [0.59, 0.65]
              0.73 | [0.69, 0.77]
##
    1 l
              0.82 | [0.76, 0.86]
##
##
    4 |
              0.92 | [0.86, 0.96]
##
## Adjusted for:
## * m = -0.01
```

##

The non-linear relationship makes the interaction effect vary. We can estimate the marginal effects (of x) based on different levels of the moderator (m):

```
summary(margins(logit_fit,at = list(m=c(-2.5,0,2.5)),variables = "x"))
```

```
##
                       AME
                                SE
    factor
                                                            upper
                 m
                                         z
                                                    lower
                                                р
##
         x - 2.5000
                    0.3259 0.0100 32.4295 0.0000
                                                   0.3062
                                                           0.3456
##
            0.0000 0.1092 0.0163 6.6823 0.0000 0.0771
##
            2.5000 -0.2242 0.0231 -9.6912 0.0000 -0.2695 -0.1788
```

Note: The range of observed values of m is [-3.39, 3.16]

Or we can formally conduct the Johnson-Neyman intervals and simple slopes analysis. The "classic" way of probing an interaction effect is to calculate the slope of the focal predictor at different values of the moderator. When the moderator is binary, this is especially informative, e.g., what is the slope for men vs. women? But you can also arbitrarily choose points for continuous moderators.

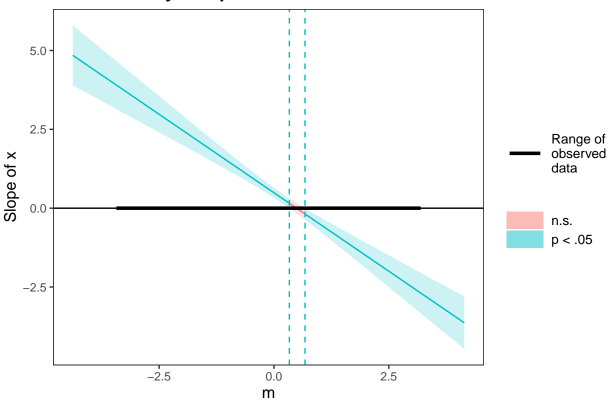
With that said, the more statistically rigorous way to explore these effects is to find the **Johnson-Neyman** interval, which tells you the range of values of the moderator in which the slope of the predictor is significant vs. nonsignificant at a specified alpha level.

The sim_slopes function will by default find the **Johnson-Neyman interval** and tell you the predictor's slope at specified values of the moderator; by default either both values of binary predictors or the mean and the mean +/- one standard deviation for continuous moderators.

```
sim_slopes(logit_fit,pred = x, modx = m, jnplot = TRUE)

## JOHNSON-NEYMAN INTERVAL
##
## When m is OUTSIDE the interval [0.33, 0.68], the slope of x is p < .05.</pre>
```

Johnson-Neyman plot



```
## SIMPLE SLOPES ANALYSIS
## Slope of x when m = -0.996597103 (-1 SD):
##
##
    Est. S.E. z val.
## -----
    1.48
         0.15
                 10.16 0.00
##
## Slope of x when m = -0.009456456 (Mean):
##
##
    Est.
          S.E. z val.
    0.50 0.08
               6.22 0.00
##
##
## Slope of x when m = 0.977684190 (+ 1 SD):
##
         S.E.
               z val.
##
     Est.
## -----
               -4.23 0.00
    -0.48
         0.11
```

V. Regression (Causal Inference)

17. Everyone knows that correlation is not causation, when regression coefficients could be interpreted as causal effects? Is including the lagged explanatory variable a way to go?

Regression models have two different purposes. One is for description, another is for causal inference. For the purpose of description, we emphasize the overall fit (e.g., R^2). For the purpose of causal inference, we place more emphasis on the "accurate" estimation of a focal variable on the dependent variable (i.e., the treatment effect). So, a natural question is whether it is possible to estimate the treatment effect even though the overall model fit is not so good (e.g., relatively low R^2).

Example 1 (independent): Let's create three random variables (x1, x2, Treatment). They are independent to each other. The focal variable is "Treatment", while the dependent variable is created based on x1 and Treatment.

```
x1 = rnorm(1000,2,1)
x2 = rnorm(1000,3,1)
Treatment = rnorm(1000,5,2)
y = 2*x1+3*Treatment+rnorm(1000)
```

We can run a regression model including all three variables as predictors of y.

```
library(jtools)
summ(lm(y~x1+x2+Treatment),model.info = FALSE, model.fit = FALSE)
```

	Est.	S.E.	t val.	p
(Intercept)	0.05	0.15	0.37	0.71
x1	2.00	0.03	62.64	0.00
x2	-0.01	0.03	-0.44	0.66
Treatment	2.99	0.02	179.15	0.00

Standard errors: OLS

For sure, the R^2 is close to 1, and the coefficients are close to the true values. How about if only regress Treatment on y (imaging that we have no way to observe or measure x1)?

```
summ(lm(y~Treatment),model.info = FALSE, model.fit = FALSE)
```

	Est.	S.E.	t val.	p
(Intercept)	4.12	0.20	20.91	0.00
Treatment	2.99	0.04	80.57	0.00

Standard errors: OLS

The coefficient of Treatment on v remains accurate, when other predictors are independent to Treatment.

Example 2 (dependent): Let's create three random variables. This time, x1 is positively correlated with x2. Actually x1 leads to x2 $(x_1 \rightarrow x_2)$. Treatment is negatively dependent on x2 $(x_2 \rightarrow \text{Treatment})$.

```
x1 = rnorm(1000,2,1)
x2 = 3.1*x1 + rnorm(1000)
Treatment = -0.9*x2 + rnorm(1000)
y = 2*x1+3*Treatment+rnorm(1000)
summ(lm(y~Treatment+x1+x2),model.info = FALSE, model.fit = FALSE)
```

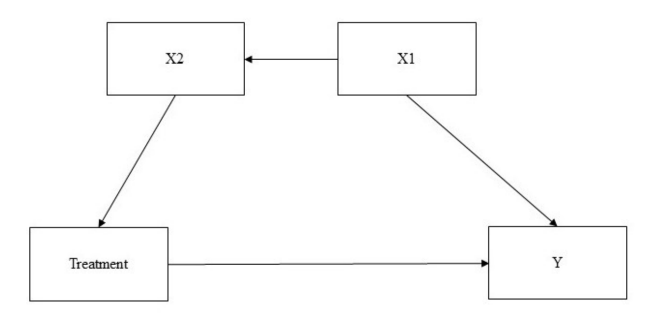


Figure 1: Causal Graph.

	Est.	S.E.	t val.	p
(Intercept)	-0.04	0.07	-0.50	0.61
Treatment	3.01	0.03	97.87	0.00
x1	1.94	0.10	19.21	0.00
x2	0.03	0.04	0.69	0.49

Standard errors: OLS

If we measured all predictors (x1, x2, and Treatment) and included them correctly in the regression model, the estimated coefficient of Treatment should be accurate as presented above. If we miss both x1 and x2, the estimated effect of Treatment will be biased:

summ(lm(y~Treatment),model.info = FALSE, model.fit = FALSE)

	Est.	S.E.	t val.	p
(Intercept)	0.76	0.09	8.42	0.00
Treatment	2.42	0.01	175.19	0.00

Standard errors: OLS

But, how to correct the bias? Should we control for both x1 and x2? The answer is "not necessary".

```
m0 <- lm(y~Treatment+x2)
m1 <- lm(y~Treatment+x1)
export_summs(m0,m1)</pre>
```

Both models can estimate the treatment effect correctly (However, the estimation of x_s is biased). Why? This is rooted in the so-called **Backdoor Criterion** in causal inferences: Given an ordered pair of variables (X, Y) in a directed acyclic graph G, a set of variables Z satisfies the backdoor criterion relative to (X, Y) if no node in Z is a descendant of X, and Z blocks every path between X and Y that contains an arrow into X.

	Model 1	Model 2
(Intercept)	0.26 **	-0.03
	(0.08)	(0.07)
Treatment	3.01 ***	2.99 ***
	(0.04)	(0.02)
x2	0.61 ***	
	(0.03)	
x1		1.99 ***
		(0.07)
N	1000	1000
R2	0.98	0.98
*** p < 0.00	1; ** p < 0.01; *	p < 0.05.

Question:

• Why we also consider demographics (age/gender) as control variables? (exogenous variables without any arrow directed to them)

Example 3 (lagged variable): If we don't know either x1 or x2, is a lagged variable of the Treatment sufficient to identify the causal effect? It works in some special conditions only!

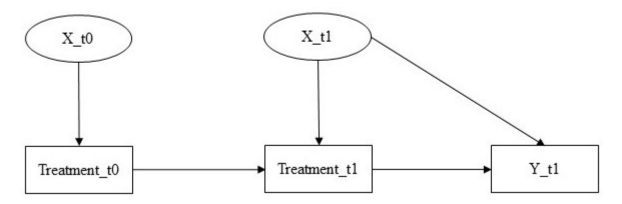


Figure 2: Another Causal Graph.

Note that x_t0 and x_t1 are unobservable.

```
# two independent confounding variables
x_t0 = rnorm(1000,1,1.5)
x_t1 = rnorm(1000,2,1.5)
```

```
# treatment and its lagged variable [simultaneous influences from x on treatment]
Treatment_t0 = 1.5*x_t0 + rnorm(1000,3,1.2)
Treatment_t1 = 2*Treatment_t0 + 1.5*x_t1 + rnorm(1000) # lagged var. should be auto-correlated
# define y
y = 2*Treatment_t1 + 3*x_t1 + rnorm(1000) # the treatment effect is 2
summ(lm(y~Treatment_t1+Treatment_t0), model.info = FALSE, model.fit = FALSE)
```

	Est.	S.E.	t val.	р
(Intercept)	1.00	0.16	6.30	0.00
$Treatment_t1$	3.66	0.03	135.15	0.00
$Treatment_t0$	-3.30	0.06	-54.68	0.00

Standard errors: OLS

No! The estimated effect of Treatment_t01 is bias. Notice that x_t0 actually is an **instrumental variable** of $x_t1 -> y$. We need to use two stage least square regression (or IV regression) to estimate the treatment effect.

```
library(ivreg)
summary(ivreg(y~Treatment_t1|Treatment_t0))
```

```
##
## Call:
## ivreg(formula = y ~ Treatment_t1 | Treatment_t0)
##
## Residuals:
##
         Min
                          Median
                                         30
                                                  Max
                    10
  -16.03945
             -3.24714
                         0.06349
                                   3.12610
                                            16.77861
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                 6.04855
                            0.37102
                                      16.30
## Treatment_t1 2.01155
                            0.02848
                                      70.62
                                               <2e-16 ***
##
## Diagnostic tests:
                    df1 df2 statistic p-value
## Weak instruments
                      1 998
                                 4231
                                      <2e-16 ***
## Wu-Hausman
                      1 997
                                 2990
                                       <2e-16 ***
## Sargan
                                           NA
                      0
                         NA
                                   NA
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 4.642 on 998 degrees of freedom
## Multiple R-Squared: 0.8901, Adjusted R-squared: 0.89
## Wald test: 4987 on 1 and 998 DF, p-value: < 2.2e-16
```

The coefficient is very close to the true value 2. So, is this the solution? No, because it is very unlikely x_t0 is not correlated in x_t1 . And it is likely that x_t0 can influence y directly (cross-lagged influence).

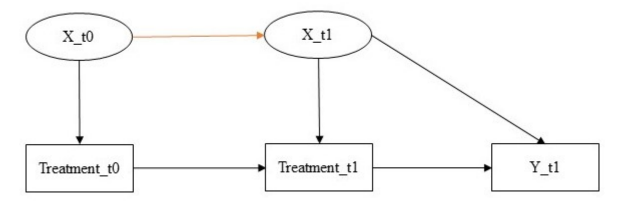


Figure 3: Revised Causal Graph.

```
# confounding variables
x_t0 = rnorm(1000,1,1.5)
x_t1 = 1.8*x_t0+rnorm(1000) #x_t0->x_t1

# treatment and its lagged variable [simultaneous influences from x on treatment]
treatment_t0 = 1.5*x_t0 + rnorm(1000,3,1.2)
treatment_t1 = 2*treatment_t0 + 1.5*x_t1 + rnorm(1000) # lagged var. should be auto-correlated
# define y
y = 2*treatment_t1 + 3*x_t1 + rnorm(1000)
summ(lm(y~treatment_t1+treatment_t0), model.info = FALSE, model.fit = FALSE)
```

	Est.	S.E.	t val.	p
(Intercept)	-0.76	0.16	-4.73	0.00
$treatment_t1$	3.69	0.03	136.95	0.00
$treatment_t0$	-2.99	0.09	-31.65	0.00

Standard errors: OLS

summary(ivreg(y~treatment_t1|treatment_t0))

```
##
## ivreg(formula = y ~ treatment_t1 | treatment_t0)
##
## Residuals:
##
                 1Q Median
       Min
                                   ЗQ
                                           Max
## -9.03205 -2.14537 -0.03731 2.13634 9.85717
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) -3.83535 0.15756 -24.34
                                            <2e-16 ***
## treatment_t1 2.80415
                           0.01095 256.20
                                            <2e-16 ***
## Diagnostic tests:
```

Both 1m and ivreg are incorrect! How about if x does not change over time, i.e., $x_t0 = x_t1$.

```
# confounding variables
x = rnorm(1000,3,1.5)

# treatment and its lagged variable [simultaneous influences from x on treatment]
treatment_t0 = 1.5*x + rnorm(1000,3,1.2)
treatment_t1 = 2*treatment_t0 + 1.5*x + rnorm(1000) # lagged var. should be auto-correlated
# define y
y = 2*treatment_t1 + 3*x + rnorm(1000)
summ(lm(y~treatment_t1+treatment_t0), model.info = FALSE, model.fit = FALSE)
```

	Est.	S.E.	t val.	p
(Intercept)	-1.17	0.17	-6.67	0.00
$treatment_t1$	3.00	0.04	78.35	0.00
$treatment_t0$	-1.23	0.11	-11.38	0.00

Standard errors: OLS

Does not work!

18. Is a random experiment always better than other methods to identify causality? Why or why not?

- Plausibility and ethics: sex manipulation?
- Experiments also have assumptions: Randomization rules out all confounding variables
- Compliance
- Heterogeneous effects (lack of external validity):

Suppose the treatment effect (conditions) varies across gender (gender*cond).

```
cond = sample(c(0,1),1000,replace = T)
gender = sample(c(0,1),1000,replace = T)
y = 2*cond -1*gender*cond + rnorm(1000)

pp = data.frame(y,cond,gender)
# a random experiment based on population
table(pp$cond)/1000
```

```
##
##
       0
              1
## 0.516 0.484
table(pp$gender)/1000
##
##
       0
## 0.484 0.516
summ(lm(y~cond,data = pp),model.info = FALSE, model.fit = FALSE)
                                            Est.
                                                   S.E.
                                                         t val.
                                                                   р
                                                         -0.70
                               (Intercept)
                                            -0.03
                                                   0.05
                                                                 0.48
                               cond
                                             1.53
                                                   0.07
                                                         23.04
                                                                 0.00
```

It does not influence the estimation of the treatment effect if we have a representative sample. However, it does not work with a biased sample.

Standard errors: OLS

```
# a random experiment from a biased sample
male = pp[pp$gender==1,]
female = pp[pp$gender==0,]
ps = rbind(male[sample(nrow(male),400),],female[sample(nrow(female),100),])
table(ps$cond)/500
```

```
## 0 1
## 0.52 0.48
```

```
table(ps$gender)/500
```

```
## 0 1
## 0.2 0.8

summ(lm(y~cond,data = ps),model.info = FALSE, model.fit = FALSE)
```

##

##

	Est.	S.E.	t val.	р
(Intercept)	-0.11	0.07	-1.63	0.10
cond	1.22	0.09	13.01	0.00

Standard errors: OLS

19. Is it always better to control more variables than less? Should we remove non-significant variables from the regression?

Collider: $x_1 \to x \leftarrow y$. Don't control for colliders. Controlling for colliders creates spurious correlations.

```
x1 = rnorm(1000)
x2 = rnorm(1000)
y = 3.5*x2 + rnorm(1000)

x = 3*x1+2*y+rnorm(1000) # x is a collider

m0 <- lm(y~x1+x2) # correct (x1 does not influence y)
m1 <- lm(y~x1+x2+x) # wrong (x1 influences y conditioning on x)
export_summs(m0,m1)</pre>
```

	Model 1	Model 2
(Intercept)	-0.00	-0.01
	(0.03)	(0.01)
x1	-0.04	-1.21 ***
	(0.03)	(0.02)
x2	3.52 ***	0.73 ***
	(0.03)	(0.05)
X		0.39 ***
		(0.01)
N	1000	1000
R2	0.92	0.98

*** p < 0.001; ** p < 0.01; * p < 0.05.

x is significantly related to y but should not be included in the regression. It makes the irrelevant variable x1 now related to y significantly.

A related question, should we include non-significant variables? If x is an irrelevant variable (to y):

```
x1 = rnorm(1000)
x2 = rnorm(1000)
# if x is really irrelevant but correlated with x1 and x2
x = 2.5*x1+3.5*x2+rnorm(1000)
y = 2.5*x1 +3.5*x2 + rnorm(1000)

m0 <- lm(y~x1+x2+x)
m1 <- lm(y~x1+x2)
export_summs(m0,m1)</pre>
```

The example demonstrated that if x is really not related to y (not caused by sampling or estimation in models), there are just some minor differences with or without including x. It is always safe the include all variables, because usually we don't know whether the insignificant variables are really non-relevant to y.

Nevertheless, there are some (minor) drawbacks by including really non-relevant variables: it will decrease df, increase SE! Not parsimony! If x is NOT correlated with x1 and x2, SEs will not change much (why?).

	Model 1	Model 2
(Intercept)	-0.06	-0.06
	(0.03)	(0.03)
x1	2.47 ***	2.43 ***
	(0.08)	(0.03)
x2	3.54 ***	3.48 ***
	(0.11)	(0.03)
X	-0.02	
	(0.03)	
N	1000	1000
R2	0.95	0.95

*** p < 0.001; ** p < 0.01; * p < 0.05.

One more example, x is indeed related to y, however, the effect size is small and thus it might be nonsignficant in the regression.

```
x1 = rnorm(1000,3,1)
x2 = rnorm(1000,5,2)
x = 3*x1+2*x2+5*rnorm(1000)
y = 2*x1 + 3*x2 + 0.02*x+ rnorm(1000,6,2.5)

m0 <- lm(y~x1+x2+x) # x is not significant but it is relevant
m1 <- lm(y~x1+x2) # removing the variable will bias the estimation
export_summs(m0,m1)</pre>
```

6.26 ***	o o a distrib
	6.24 ***
(0.32)	(0.32)
1.82 ***	1.88 ***
(0.09)	(0.08)
3.05 ***	3.09 ***
(0.05)	(0.04)
0.02	
(0.01)	
1000	1000
0.87	0.86
	1.82 *** (0.09) 3.05 *** (0.05) 0.02 (0.01)

*** p < 0.001; ** p < 0.01; * p < 0.05.

20. Should we control for mediators to estimate the treatment effect?

- direct effects
- indirect effects
- total effects = direct + indirect effects

```
x = rnorm(1000,5,1)
m = 2*x+3*rnorm(1000)
y = 3*x+5*m+7*rnorm(1000,6,3)

# the traditional way of mediation test:
m0 <- lm(y~x)
m1 <- lm(y~x+m)
export_summs(m0,m1)</pre>
```

The tradition way of mediation test is to compare the regression models with/without the mediator (i.e., m0, m1). Without mediator in m0, the coefficient of x is 12.9, while it is 2.97 in m1, indicating the existence of partially mediation. This method is not precise, without estimation of the size and test of significance. It would be better to estimate using the SEM approach.

	Model 1	Model 2	
(Intercept)	41.16 ***	40.08 ***	
	(4.29)	(3.46)	
X	12.90 ***	2.97 ***	
	(0.84)	(0.80)	
m		5.08 ***	
		(0.22)	
N	1000	1000	
R2	0.19	0.47	

*** p < 0.001; ** p < 0.01; * p < 0.05.

```
## lavaan 0.6-9 ended normally after 30 iterations
##
     Estimator
##
                                                        ML
##
     Optimization method
                                                    NLMINB
##
     Number of model parameters
##
##
     Number of observations
                                                      1000
##
## Model Test User Model:
##
##
     Test statistic
                                                     0.000
##
     Degrees of freedom
                                                         0
##
## Parameter Estimates:
##
```

##	Standard errors					Sootstrap
##	Number of requested bootstrap draws					
##	Number of successful bootstrap draws					1000
##						
##	Regress	ions:				
##			Estimate	Std.Err	z-value	P(> z)
##	m ~					
##	x	(a)	1.956	0.089	21.926	0.000
##	у ~					
##	m	(b)	5.079	0.218	23.246	0.000
##	x	(c)	2.972	0.816	3.642	0.000
##						
##	Variance	es:				
##			Estimate	Std.Err	z-value	P(> z)
##	.m		9.295	0.423	21.965	0.000
##	· y		443.487	21.301	20.820	0.000
##						
##	${\tt Defined}$	${\tt Parameters}$:			
##			Estimate	Std.Err	z-value	P(> z)
##	ab		9.932	0.593	16.754	0.000
##	tota	al	12.904	0.813	15.871	0.000

The total effect is 12.9, which is consistent with m0. This is the overall treatment effect. ab is the indirect effect (the effect of x on y via m). It is possible that x could influence y via many mechanisms. Unless, you're interested in a particular mechanism, we estimate the treatment effect without controlling for mediators.