# Assignment 2

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#### & Theoratical exercises

### Exercise 2.1 (Additional exercises 1.3)

"+/-": positive/negative results; "disease/!disease": have/don't have cancer

**a**)

```
P(+) = P(+ | disease) * P(disease) + P(+ | !disease) * P (!disease)
= 0.95 * 0.004 + (1-0.95) * (1-0.004)
= 0.054
```

The probability P(+) calculated here is called **belief** in statistics, representing the possibility of a certain event happening, whilst the probability P(disease | +) calculated in the additional exercises is called **conditional probability** illustrating the possibility of a certain event happening based on a **revised model** tunned by new information (an event has already happened).

b)

```
P(disease | +) = P(disease) * P(+ | disease) / P(+)
= 0.95 * 0.004 / 0.054 = 38 / 536
= 0.071
```

**c**)

(1) Yes, they are dependant because

```
P(disease AND +) = P(disease) * P(+ | disease)
= 0.004 * 0.95
```

is no the same as

$$P(diease) * P(+) = 0.004 * 0.054$$

(2) Yes, a test result was positive increase the risk of having cancer because

$$P(disease | +) == 0.071 > P(disease) == 0.004$$

#### Exercise 2.2

a)

Let X denote the random variable 'outcome of two 8-sided die-rolls'; Let  $x_i$  (where i = 1, 2, ..., 64) denote the possible outcomes

Because multiple rolls do not influence each other, we have

1)

$$x_1 == (1, 1); x_2 == (1, 2); x_3 == (1, 3); x_4 == (1, 4); x_5 == (1, 5); x_6 == (1, 6); x_7 == (1, 7); x_8 == (1, 8);$$

$$x_9 == (2, 1); x_{10} == (2, 2); x_{11} == (2, 3); x_{12} == (2, 4); x_{13} == (2, 5); x_{14} == (2, 6); x_{15} == (2, 7); x_{16} == (2, 8);$$

. . .

$$x_{57} == (8, 1); x_{58} == (8, 2); x_{59} == (8, 3); x_{60} == (8, 4); x_{61} == (8, 5); x_{62} == (8, 6); x_{63} == (8, 7); x_{64} == (8, 8);$$

2) 
$$P(X=x_1) == P(X=x_2) == \dots == P(X=x_{64}) == 1/64$$

b)

Let Y denote the random variable 'number of 1's in two 8-sided die rolls'; Let  $y_j$  (where j = 1, 2, 3) denote the possible outcomes

j	Уј	P(Y=y <sub>j</sub> )
1	0	49/64
2	1	7/32
3	2	1/64

**c**)

$$E(Y) = 0 * 49/64 + 1 * 7/32 + 2 * 1/64 = 0.25 = 1/4$$

d)

$$Var(Y) = 0^2 * 49/64 + 1^2 * 7/32 + 2^2 * 1/64 - E(Y)^2 = 7/32$$

**e**)

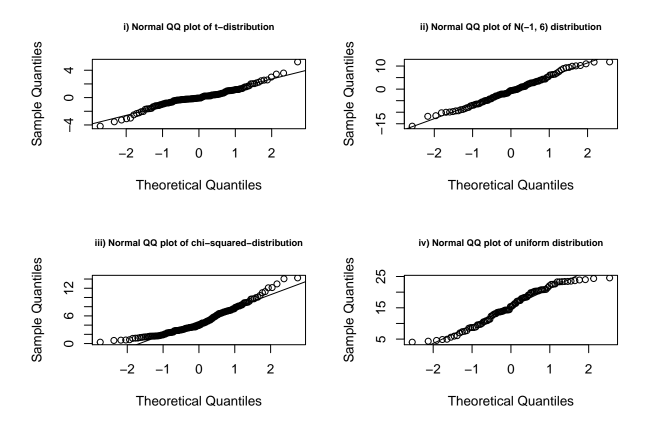
Since the sample size is much greater than 30, according to the **central limit theorem**, the random variables  $\mathbf{X_i}$  approximately have a normal distribution with an expectation  $\mu$  of 1/4 and a standard deviation  $\sigma$  of  $\sqrt{7/32}$ , which is

$$N(1/4, \frac{7/32}{n})$$

## & R-exercises

#### Exercise 2.3

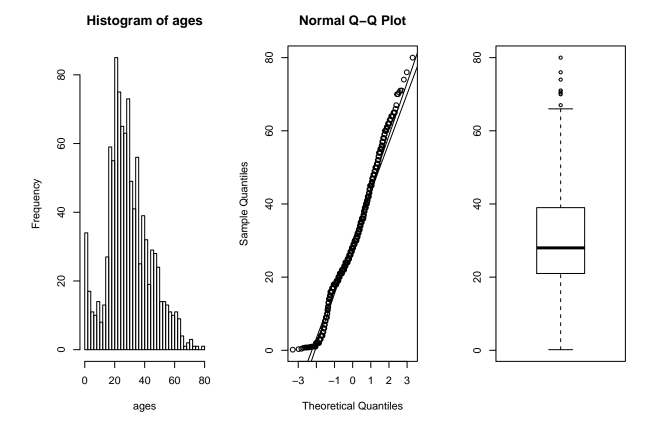
a)



- i) The left part of the plot is below the QQ-line and the right part is above the QQ-line. Therefore, we conclude that both the left tail and right tail of the t-distribution is heavier than that of the standard normal distribution;
- ii) As the top-right figure demonstrates, because N(-1, 6) and the standard normal are in the same distribution are in the same **location-scale family**, the QQ-line of the N(-1, 6) distribution adhere to the QQ-line of the standard distribution very well;
- iii) Both the left part and the right part of the plot are above the QQ-line. Therefore, we conclude that the left tail of the chi-squared-distribution is lighter than that of the standard normal distribution, whilst the right tail of the chi-squared-distribution is heavier than that of the standard normal distribution;
- iv) The left part of the plot is above the QQ-line and the right part is below the QQ-line. Therefore, we conclude that the left tail of the uniform distribution is heavier than that of the standard normal distribution, whilst the right tail of the uniform distribution is lighter than that of the standard normal distribution.

b)

#### (i) titanic3.csv dataset:

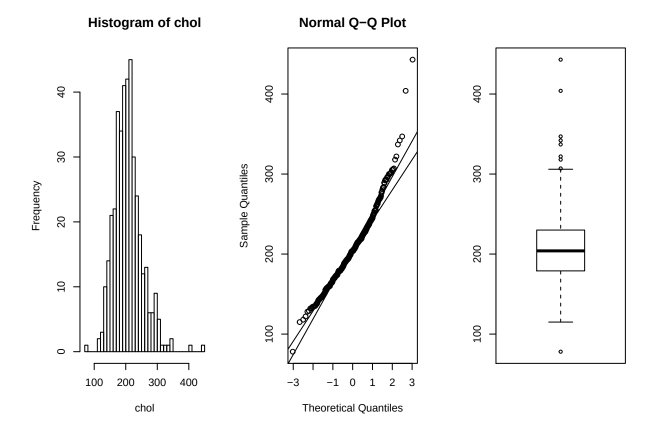


- (1) According to the plots above, we conclude that the answer for the first question is **Normality cannot** be excluded for the following reasons:
  - a. the histogram demonstrates a roughly bell shape;
  - b. as the normal Q-Q plot shows, although the right tail is heavier and right tail is lighter, the major quantiles adhere to the Q-Q line and  $\mathbf{y} = \mu \mathbf{x} + \sigma^2$  family;
  - c. as we can see from the boxplot, excluding outliers, the density of the date, ranging from 1st quartile to 3rd quartile, lies in the middle of the range, as well as the median value. Thus, the distribution is **approximately symmetric.**

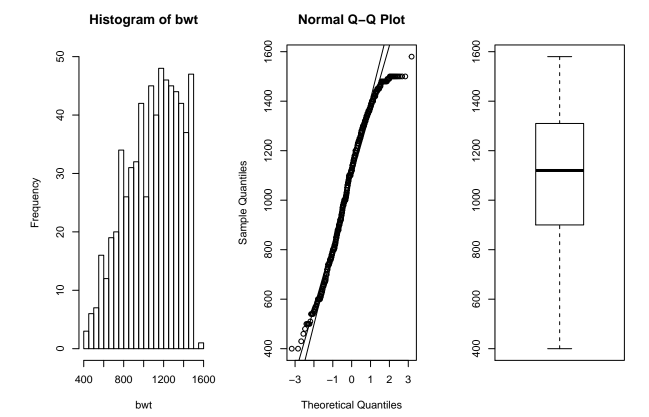
#### (2) Peculiarities:

- a. as the histogram shows, the left part of the distribution is not only heavier but also has an abnormal decreasing trend;
- b. the location of this distribution is dragged left (left-skewed) because of too many children.

#### (ii) diabetes.csv dataset:



- (1) According to the plots above, we conclude that the answer for the first question is **Normality cannot** be excluded for the following reasons:
  - a. the histogram demonstrates a roughly bell shape;
  - b. as the normal Q-Q plot shows, although the right tail is lighter and right tail is heavier, the major quantiles adhere to the Q-Q line and  $\mathbf{y} = \mu \mathbf{x} + \sigma^2$  family;
  - c. as we can see from the boxplot, excluding outliers, the density of the date, ranging from 1st quartile to 3rd quartile, lies in the middle of the range, as well as the median value. Thus, the distribution is **approximately symmetric.**
- (2) **Peculiarities:** As we can see from the histogram, there are some "holes" which represent absences of some certain cholesterol values.
- (iii) vlbw.csv dataset:



- (1) according to the plots above, we conclude that the answer for the first question is **Obviously not** from a normal distribution" for the following reasons:
  - a. the shape of the histogram is by no means a **bell shape**;
  - b. as the normal Q-Q plot shows, although the major quantiles adhere to the Q-Q line and  $y = \mu x + \sigma^2$  family, the left tail and, especially, the right tail horribly deviate from two lines;
  - c. as we can see from the boxplot, the distribution is quite **left-skewed** and therefore it is not **symmetric** whatsoever.
- (2) **Peculiarities:** Compared to the previous two datasets, this distribution demonstrates a more **centralized pattern** and there are **no outliers**.

#### Exercise 2.4

**a**)

As the diagram illustrates, when the number of trails becomes larger, the absolute difference of two die rolls converge to the theoretically expected value of the absolute difference 1.9444.

b)

(1)

```
source("function02.txt")
num_trails = 1000
outcomes = diffdice(num_trails)

E = 0
for (x_i in 1:6) {
    # calculate the probability of P(X=xi);
    P_xi = length(outcomes[outcomes==x_i]) / num_trails
    # calculate the approximate expectation which is the weighted average of x_i;
    E = E + x_i * P_xi
    E
}
```

#	outcome	frequency
1	0	168
2	1	260
3	2	219
4	3	190
5	4	112
6	5	51

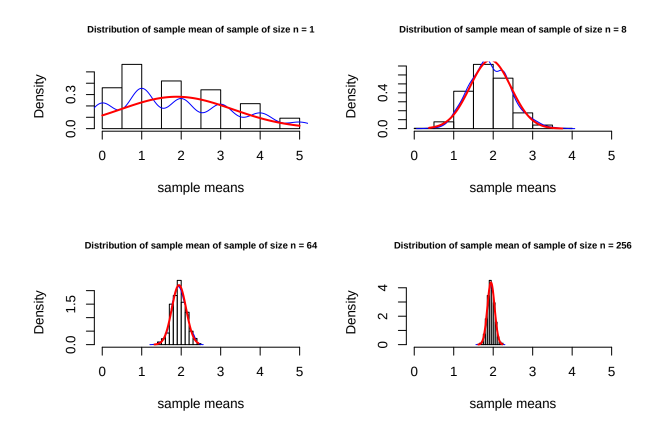
Using the function diffdice, we found an approximate value of expectation of the random variable 'the absolute difference of two die rolls': 1.971

(2)

```
P_3 = length(outcomes[outcomes==3]) / num_trails
```

The probability of the event 'the absolute difference of two die rolls is 3' is 0.19

**c**)



d)

The blue lines in the above figures represent empirical distributions and the red lines are the normal distribution. As these plots demonstrate, those two lines gradually merge together as the sample size become larger. In other words, the distribution of the sample means steadily move toward a normal distribution.

## Appendix

```
# 2.3 a
par(mfrow=c(2,2))
# i)
a = rt(150, 4)
qqnorm(a, main = "i) Normal QQ plot of t-distribution")
abline(mean(a), sd(a))
# ii)
b = rnorm(100, -1, 6)
qqnorm(b, main = "ii) Normal QQ plot of N(-1, 6) distribution")
abline(mean(b), sd(b))
# iii)
c = rchisq(170, 5)
```

```
qqnorm(c, main = "iii) Normal QQ plot of chi-squared-distribution")
abline(mean(c), sd(c))
# iv)
d = runif(90, 4, 25)
qqnorm(d, main = "iv) Normal QQ plot of uniform distribution")
abline(mean(d), sd(d))
# 2.3 b
par(mfrow=c(1,3))
# (i)
titanic = read.csv("titanic3.csv")
ages = titanic$age
sd_titanic=sd(ages, na.rm=TRUE)
mean_titanic=summary(ages)[4]
# Add legends later
hist(ages, breaks = 50)
qqnorm(ages)
qqline(ages)
abline(as.numeric(mean_titanic), sd_titanic)
boxplot(ages)
# (ii)
diabetes = read.csv("diabetes.csv")
chol = diabetes$chol
sd chol=sd(chol, na.rm=TRUE)
mean_chol=summary(chol)[4]
hist(chol, breaks =50)
qqnorm(chol)
qqline(chol)
abline(as.numeric(mean_chol), sd_chol)
boxplot(chol)
# (iii)
vlbw = read.csv("vlbw.csv")
bwt = vlbw$bwt
sd_bwt=sd(bwt, na.rm=TRUE)
mean_bwt=summary(bwt)[4]
hist(bwt, breaks = 30, ylim = c(0, 50))
qqnorm(bwt)
qqline(bwt)
abline(as.numeric(mean_bwt), sd_bwt)
boxplot(bwt)
# Exercise 2.4
# a)
dice = (1: 6)
results = c()
```

```
for (j in 1:1000) {
  mean_diff = 0
  for(i in 1:j) {
    trail = sample(6, 2, replace = TRUE)
    diff = abs(trail[1]-trail[2])
    mean_diff = mean_diff + diff
 mean diff = mean diff/j
  results[j] <- mean_diff
plot(results, type="l", ylim=c(0, 5), xlim=c(0,1100), xlab = "Number of trails", ylab = "the absolute d
abline(1.9444, 0) # later, add 1.9444 labbel
# c)
samples_of_1 = numeric(1000)
samples_of_8 = numeric(1000)
samples_of_64 = numeric(1000)
samples_of_256 = numeric(1000)
for (i in 1:1000) {
  samples_of_1[i] <- mean(diffdice(1))</pre>
  samples_of_8[i] <- mean(diffdice(8))</pre>
  samples_of_64[i] <- mean(diffdice(64))</pre>
  samples_of_256[i] <- mean(diffdice(256))</pre>
}
par(mfrow=c(2,2))
legend( 0,0, legend=c("imperical distribution", "normal distribution"), col=c("blue", "red"), lty=1:2,
hist(samples_of_1, prob = T, xlim = c(0,5), main = "Distribution of sample mean of sample of size n = 1
lines(density(samples_of_1), col = "blue")
x <- seq(min(samples_of_1), max(samples_of_1), by=.001)
y <- dnorm(x, mean=mean(samples_of_1), sd=sd(samples_of_1))
lines(x, y, type="l", lwd=2, col = "red")
hist(samples_of_8, prob = T, xlim = c(0,5), main = "Distribution of sample mean of sample of size n = 8
lines(density(samples_of_8), col = "blue")
x <- seq(min(samples_of_8), max(samples_of_8), by=.001)
y <- dnorm(x, mean=mean(samples_of_8), sd=sd(samples_of_8))
lines(x, y, type="1", lwd=2, col = "red")
hist(samples_of_64, prob = T, xlim = c(0,5), main = "Distribution of sample mean of sample of size n = "
lines(density(samples_of_64), col = "blue")
x <- seq(min(samples_of_64), max(samples_of_64), by=.001)
y <- dnorm(x, mean=mean(samples_of_64), sd=sd(samples_of_64))
lines(x, y, type="l", lwd=2, col = "red")
hist(samples_of_256, prob = T, xlim = c(0,5), main = "Distribution of sample mean of sample of size n =
lines(density(samples_of_256), col = "blue")
x \leftarrow seq(min(samples_of_256), max(samples_of_256), by=.001)
y <- dnorm(x, mean=mean(samples_of_256), sd=sd(samples_of_256))
lines(x, y, type="l", lwd=2, col = "red")
```