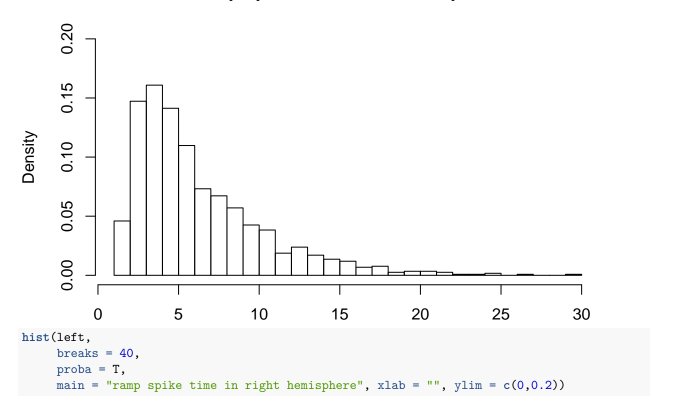
Statistics for Bioinformatics and eScience - Handin 2

Group 1
12/02/2019

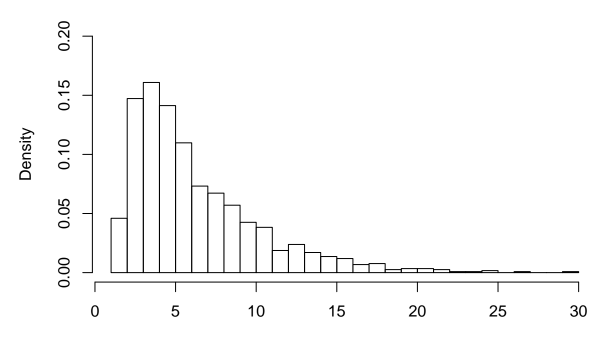
1. Brain cell dataset

4.1: plot separately histograms for the distribution of the ramp spike time variable in the left and right hemisphere

ramp spike time in left hemisphere

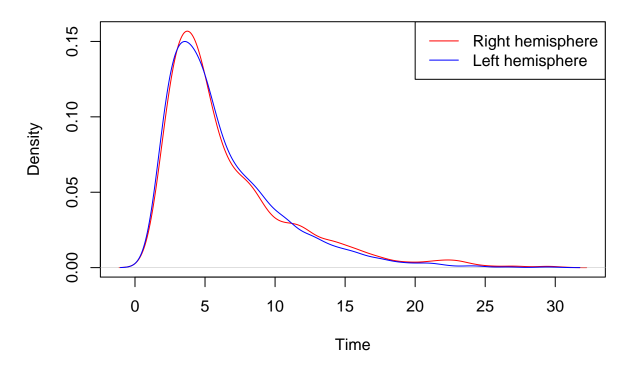


ramp spike time in right hemisphere



4.2: plot together kernel density estimations for the ramp spike time variable in the left and right hemisphere.

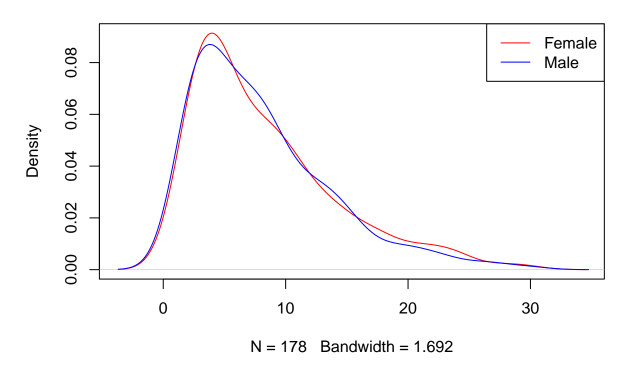
Kernel density estimation for the ramp spike time



4.3: plot together kernel density estimations for the ramp spike time variable for males and females.

```
female_raw <-
  cells$ef__peak_t_ramp[cells$donor__species == "Homo Sapiens" &
  cells$donor__sex == "Female"]
  female <- female_raw[!is.na(female_raw)]</pre>
  male raw <-
  cells$ef__peak_t_ramp[cells$donor__species == "Homo Sapiens" &
  cells$donor__sex == "Male"]
  male <- male_raw[!is.na(male_raw)]</pre>
plot(density(female),
     col = "red",
     main = "Kernel Density Estimations for males and females")
lines(density(male),
      col = "blue")
legend("topright",
       legend = c("Female", "Male"),
       col = c("red", "blue"),
       lty = 1)
```

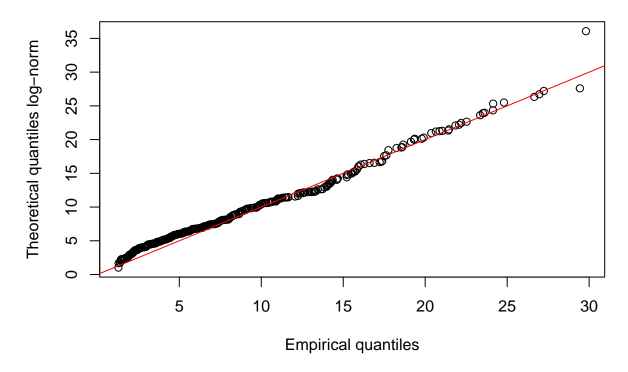
Kernel Density Estimations for males and females



 $\mathbf{Ex} \ \mathbf{5}$

5.1: Q-Q plot of ramp spike time distribution for humans, between empirical quantiles and theoretical ones (dlnorm distribution)

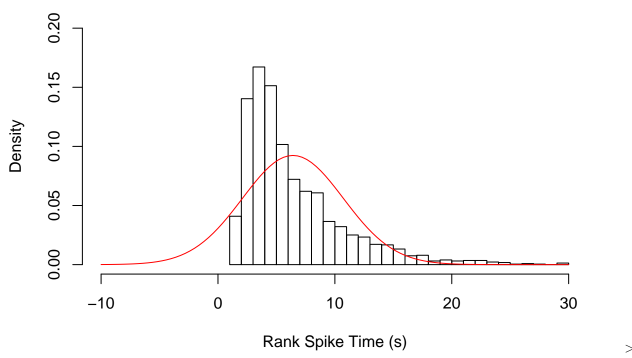
Q-Q Plot, ramp spike time d. and log-normal d.



5.2: Plot the Gaussian density on top of the histogram of the ramp spike time with better parameter values

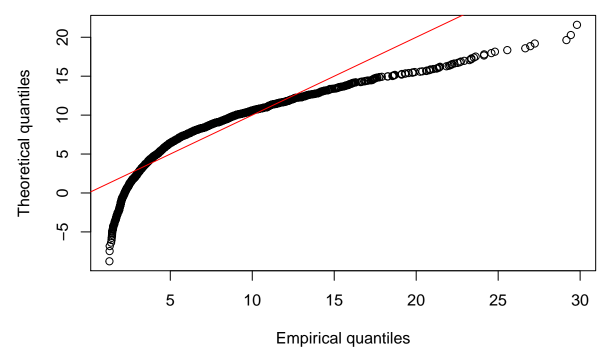
```
allcells <- cells$ef__peak_t_ramp[!is.na(cells$ef__peak_t_ramp)]
hist(allcells,
    breaks = 40,
    ylim = c(0, 0.2),
    xlim = c(-10, 30),
    probability = TRUE,
    main = "Rank Spike Time frequencies for all cells and d.norm estimation",
    xlab = "Rank Spike Time (s)")
curve(dnorm(x, mean = 6.41, sd = 4.32),
    col = "red",
    add = TRUE)</pre>
```

Rank Spike Time frequencies for all cells and d.norm estimation



Our data daesnt have a left tail and the peak of the histogram is much larger than the curve of the norm distribution.

5.3: Q-Q plot for the ramp spike time against the Gaussian distribution with these new parameter values



We can visually observe that our data don't follow a Gaussian distribution since several quantiles in the tails don't match in the diagonal.

2. Empirical mean and variance

6.1: Sample n values of $X \sim Bi(parameter\ size=100,\ prob=0.3)$ for $n=10,\ 100,\ 1000,\ 1000.$

```
n = 100
               # n. of experiments (size or i.e. nb of coin tosses)
p = 0.3
               # p. of success (i.e. head)
EM = c()
EV = c()
Esd = c()
rep = c()
for (x in c(10, 100, 1000, 10000)){
                                           \# x = nb of sampling repetitions
  sample <- rbinom(x, size = n, prob = p)</pre>
  EM <- c(EM, mean(sample))</pre>
                                            # empirical mean
  EV <- c(EV, var(sample))</pre>
                                            # empirical variance
  Esd <- c(Esd, sd(sample))
                                            # empirical standard deviation
  rep <- c(rep, x)
                                            # n. of sampling repetition
mean <- n * p
                                      # true or theoretical mean
var <- n * p * (1-p)
                                      # true or theoretical variance
sd <- sqrt(var)</pre>
                                      # true or theoretical standard deviation
data.frame(EM, mean, EV, var, Esd, sd,
           row.names = rep)
```

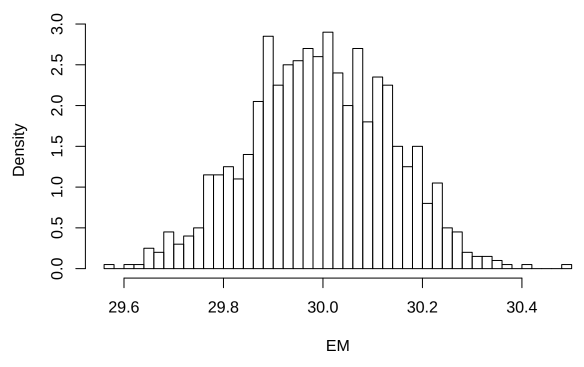
EM mean EV var Esd sd ## 10 30.2000 30 9.955556 21 3.155243 4.582576

```
## 100 29.4500 30 20.775253 21 4.557988 4.582576
## 1000 29.8450 30 19.296271 21 4.392752 4.582576
## 10000 29.9306 30 21.212305 21 4.605682 4.582576
```

6.2: Write a function to repeat the sampling and estimation for n = 1000 and plot.

```
samp_est_f <- function(nb_rep){</pre>
 n = 100
  p = 0.3
  EM = c()
  EV = c()
  Esd = c()
  for (i in 1:1000){
    sample <- rbinom(nb_rep, size = n, prob = p)</pre>
    EM <- c(EM, mean(sample)) # that's an easy way to "append" values to a vector
  }
  hist (EM,
       probability = TRUE,
       breaks = 50)
                               # histogram of the distribution of the empirical means
  data = list()
                               # list to store the output of the function
  data$EM <- EM
                                # means distributions for 6.4
  data$sample <- sample</pre>
                               # the last sample generated
                                # n. of repetition
  data$rep <- nb_rep
  data$var_EM <- var(EM)</pre>
                              # (emp.) variance of empirical/sample means
                                # (emp.) standard deviation of the empirical/sample means
  data$sd_EM <- sd(EM)</pre>
  cat("rep:", data$rep, # cat() to print numbers and strings in the same line
      ", var_EM:", data$var_EM,
      ", sd_EM:", data$sd_EM)
  return(data)
data <- samp_est_f(1000)</pre>
```

Histogram of EM



rep: 1000 , var_EM: 0.01999034 , sd_EM: 0.1413872

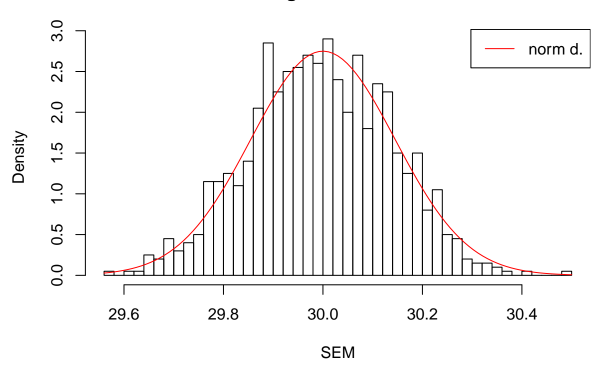
6.3: Compute the standard error of the mean

```
SEM_f <- function(x){
    SEM <- (sd(data$sample) / sqrt(length(data$sample) - 1))
    return(SEM)
}
data$SEM <- SEM_f(1000)
cat("sd_EM:", data$sd_EM, ", SEM:", data$SEM)

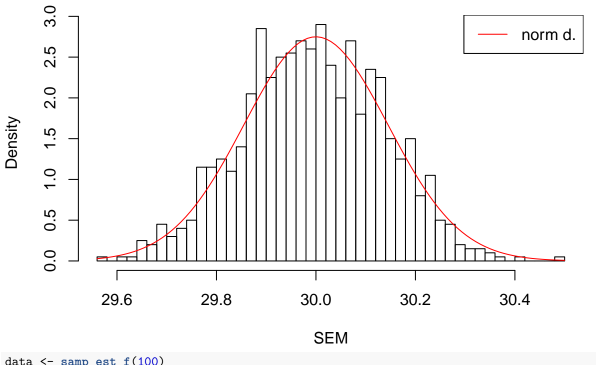
## sd_EM: 0.1413872 , SEM: 0.1451042

# SEM = ESTIMATOR of the error between the sample means and the population mean
# sd_EM = standard deviation of the sample means</pre>
```

6.4: draw the density of Gaussian distribution with parameters mean = 100 \times 0.3 and sd equal to the sem



check the result for different values of x:



```
data <- samp_est_f(100)
data$SEM <- SEM_f(100)

cat("rep:", data$rep,
        ", var_EM:", data$var_EM,
        ", sd_EM:", data$sd_EM)

## rep: 1000 , var_EM: 0.01999034 , sd_EM: 0.1413872

cat("sd_EM:", data$sd_EM, ", SEM:", data$SEM)

## sd_EM: 0.1413872 , SEM: 0.1451042

plot_f(100)</pre>
```

