Week 1

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Ex 1 Blood types

In classical genetics, we recognize three alleles for the ABO gene that regulates ABO blood type: i, I^A and I^B . Any individual inherits a complete set of chromosomes from its two parents and thus the genotype of an individual is a pair of alleles (the order is not important).

1.1 Sample space

Describe the sample space related to the genotype of an individual

$$\Omega = \{\{i,i\},\{i,I^A\},\{i,I^B\},\{I^A,I^B\},\{I^A,I^A\},\{I^B,I^B\}\}$$

$$|\Omega| = 6 = 2 \times 3$$
 (not requested)

1.2 Probability for the offspring

Suppose both parents have genotype $iI^A = \{i, I^A\}$, compute the probability that the offspring will have genotype $ii = \{i, i\}$ and the probability that at least one of the two alleles will be i

We can represent the situation with a table, that we can generate in R as

```
## parent2
## parent1 i i^A
## i "i i" "i i^A"
## i^A "i^A i" "i^A i^A"
```

The possible genotypes for the offspring are,

$$ii, iI^A, I^AI^A$$

The probabilities of each cell in the table is 0.25 thus, the probabilities for the possible genotypes are,

$$P(ii) = 0.25$$
 $P(I^A I^A) = 0.25$ $P(iI^A) = 0.25 + 0.25 = 0.5$

Or similarly we can use the independence assumption to say that

$$P(ii) = P(parent 1 \text{ gives } i \text{ and parent 2 gives } i) =$$

$$P(\{\text{parent 1 gives } i\} \cap \{\text{ parent2 gives } i\}) =$$

$$P(\{\text{parent 1 gives } i\}) \times P(\{\text{ parent2 gives } i\}) = 0.5 \times 0.5$$

And so on for the other probabilities.

As you can see $P(ii) + P(iI^A) + P(I^AI^A) = 1$.

Now we can easily compute the required probabilities,

$$P(ii) = 0.25$$

and

$$P(\text{at least one alleles is }i) = P(\{ii\} \text{ or } \{iI^A\}) = P(\{ii\} \cup \{iI^A\})$$

$$= P(ii) + P(iI^A) = 0.75$$

Since the two events $\{ii\}$ and $\{iI^A\}$ are disjoint.

1.3 Two generations

- Parents with genotypes iI^A and iI^B generate offspring A
- Parents with genotypes I^AI^B generate offspring B
- Parents A and B generate individual C

Compute the probability of genotype of C being I^AI^B

$$P(\text{genotype of }C=I^AI^B)=$$

$$P(A\to C=I^A \text{ and }B\to C=I^B)+P(A\to C=I^B \text{ and }B\to C=I^A)$$

where we use the notation $K \to M = zzz$ to denote the events where parent K gives allele zzz to offspring M.

We proceed now to obtain the probabilities

$$P(A \to C = I^A \text{ and } B \to C = I^B)$$
 (first case)

$$P(A \to C = I^B \text{ and } B \to C = I^A)$$
 (second case)

Since the events of two different parents are independent, we have that

$$P(A \to C = I^A \text{ and } B \to C = I^B) = P(A \to C = I^A) \times P(B \to C = I^B)$$
 (first case)

To obtain for example $P(A \to C = I^A)$ we need to consider the possible genotypes for A, Individual A can have one of the four possible genotypes,

$$ii, iI^A, iI^B, I^AI^B$$

And

$$P(A=ii) = P(A=iI^A) = P(A=iI^B) = P(A=I^AI^B) = 0.25$$

Moreover we have that,

$$P(A \rightarrow C = I^{A}|A = ii) = 0$$

$$P(A \rightarrow C = I^{A}|A = iI^{A}) = 0.5$$

$$P(A \rightarrow C = I^{A}|A = iI^{B}) = 0$$

$$P(A \to C = I^A | A = I^A I^B) = 0.5$$

Thus using the chain rule (P(A, B) = P(A|B)P(B)) we obtain,

$$P(A \to C = I^A) = (0 \times 0.25 + 0.5 \times 0.25 + 0 \times 0.25 + 0.5 \times 0.25 = 0.25$$

For individual B we have that,

$$P(B = I^A I^A) = P(I^B I^B) = 0.25$$

and

$$P(B = I^A I^B) = 0.5$$

and moreover,

$$P(B \to C = I^B | B = I^A I^A) = 0$$

 $P(B \to C = I^B | B = I^A I^B) = 0.5$
 $P(B \to C = I^B | B = I^B I^B) = 1$

Thus now using again the chain rule,

$$P(B \to C = I^B) = 0 \times 0.25 + 0.5 \times 0.5 + 1 \times 0.25 = 0.5$$

We can now obtain from equation (first case) that,

$$P(A \to C = I^A \text{ and } B \to C = I^B) = 0.25 \times 0.5 = 0.125$$

Similarly we need to obtain,

$$P(A \to C = I^B \text{ and } B \to C = I^A) = P(A \to C = I^B) \times P(B \to C = I^A)$$
 (second case)

The same reasoning will lead us to,

$$P(A \rightarrow C = I^B) = 0.25$$

and

$$P(B \to C = I^A) = 0.5$$

Thus,

$$P(A \rightarrow C = I^B \text{ and } B \rightarrow C = I^A) = 0.125$$

And finally

$$P(\text{genotype of }C=I^AI^B)=$$

$$P(A\to C=I^A\text{ and }B\to C=I^B)+P(A\to C=I^B\text{ and }B\to C=I^A)=$$

$$0.125+0.125=0.25$$

1.4 Phenotypes

Which are the possible phenotypes for individual C?

The possible genotypes are,

$$\{iI^A, iI^B, I^AI^B, I^AI^A, I^BI^B\} = \Omega \setminus \{ii\}$$

Then the possible phenotypes are:

- Type B from iI^B or I^BI^B
- Type A from iI^A or I^AI^A
- type AB from I^AI^B

The only phenotypes that individual C can not be is **type 0**.

Can you describe the distribution of the phenotype of C?

We have that,

$$P(\text{phenotype } C = \text{Type AB}) = P(C = I^A I^B) = 0.25$$

Moreover it is obvious that,

$$P(\text{phenotype } C = \text{Type A}) = P(\text{phenotype } C = \text{Type B})$$

since the problem is symmetric. And since the sum of the probabilities has to be 1, we obtain that,

$$P(\text{phenotype } C = \text{Type A}) = (1 - 0.25)/2 = 0.375$$

and

$$P(\text{phenotype } C = \text{Type B}) = (1 - 0.125)/2 = 0.375$$

1.5 Conditional probability

Ex 2 Some simulations

2.1 Random genotype

How can we simulate a random individual genotype?

We use the sample function in R, and we represent genotypes with vectors.

```
randomGenotype <- function(alleles = c("i", "A", "B")){
  sample(alleles, size = 2, replace = T)
}
randomGenotype()</pre>
```

```
## [1] "A" "A"
```

2.2 Random offspring and phenotype

```
randomOffspring <- function( parent1, parent2){
  c( sample(parent1, size = 1), sample(parent2, size = 1))
}</pre>
```

```
p1 <- randomGenotype()</pre>
p2 <- randomGenotype()</pre>
randomOffspring(p1, p2)
## [1] "A" "B"
genotype2phenotype <- function(genotype){</pre>
  if (all(genotype == "i")){
  return("Type 0")
  }else if ("i" %in% genotype){
      return(paste("Type", genotype[genotype != "i"] ))
  }else if (genotype[1] == genotype[2]){
    return(paste("Type", genotype[1]))
  }else{
    return("Type AB")
  }
}
Lets test it,
genotype2phenotype(c("i","i"))
## [1] "Type O"
genotype2phenotype(c("A","i"))
## [1] "Type A"
genotype2phenotype(c("i","B"))
## [1] "Type B"
genotype2phenotype(c("A","A"))
## [1] "Type A"
genotype2phenotype(c("B","B"))
## [1] "Type B"
genotype2phenotype(c("A","B"))
## [1] "Type AB"
Or in a more compact way,
#generate all possible genotypes
possible <- expand.grid(c("i","A","B"), c("i","A","B") )</pre>
possible phen <- apply (possible, MARGIN = 1, FUN = genotype2phenotype)
possible
    Var1 Var2
##
                  phen
## 1
        i
             i Type O
## 2
        Α
             i Type A
## 3
             i Type B
        В
## 4
      i
            A Type A
## 5
             A Type A
        Α
```

```
## 6 B A Type AB
## 7 i B Type B
## 8 A B Type AB
## 9 B B Type B
```

2.3

```
Approximate from simulation the probabilities of Ex 1.2

parent1 <- parent2 <- c("i", "A")

N <- 10000

## or you can use sapply or a for.
offsprings <- t(replicate(N, randomOffspring(parent1, parent2)))

pii_est <- nrow(offsprings[ offsprings[,1] == "i" & offsprings[,2] == "i", ]) / N

pii_est

## [1] 0.2567

pi_est <- nrow(offsprings[ offsprings[,1] == "i" | offsprings[,2] == "i", ]) / N

pi_est

## [1] 0.758

2.4

Code the two generation scenario and sample phenotype of C
```

```
twoGenC <- function(){
A <- randomOffspring(c("i","A"), c("i","B"))
B <- randomOffspring(c("A","B"), c("A","B"))

C <- randomOffspring(A, B)
return(list(A = A, B = B, C = C))
}

twoGenC()</pre>
```

```
## $A
## [1] "A" "i"
##
## $B
## [1] "A" "B"
##
##
## $C
## [1] "i" "A"
Sample 1000 phenotypes of C,
phenotypesC <- replicate(1000, genotype2phenotype(twoGenC()$C))
table(phenotypesC)</pre>
```

phenotypesC

```
Type A Type AB Type B
##
       365
               239
                       396
table(phenotypesC) / 1000
## phenotypesC
   Type A Type AB Type B
    0.365
            0.239
                     0.396
Increase the number of sample
N <- 100000
phenotypesC <- replicate(N, genotype2phenotype(twoGenC()$C))</pre>
table(phenotypesC)
## phenotypesC
  Type A Type AB Type B
     37138
             25173
                     37689
table(phenotypesC) / N
## phenotypesC
## Type A Type AB Type B
## 0.37138 0.25173 0.37689
```

Ex 3 DNA sequences

Let

$$\Omega = \{A,B,C,G\}$$

The sample space of a single DNA nucleotide type.

3.1

Describe the sample space of DNA sequences of length 2

$$\Theta = \Omega \times \Omega = \Omega^2 = \{(A, A), (A, B), \dots, (A, G), (B, A), (B, B), \dots (G, G)\}$$

3.2

Describe the event of observing the motif ACG in a DNA sequence of length 5

$$E = \{ACG **, *ACG*, **ACG\}$$

where * indicates whatever nucleotide.

Compute the probability of this event

Observe that the events ACG * *, *ACG*, * *ACG are disjoints. Thus,

$$P(E) = P(ACG * *) + P(*ACG *) + P(* * ACG)$$

Moreover,

$$P(ACG * *) = P(*ACG*) = P(* * ACG) = \left(\frac{1}{4}\right)^3$$

since the events in different positions are independent.

Thus,

$$P(E) = 3\left(\frac{1}{4}\right)^3$$

 $3 * (0.25)^3$

[1] 0.046875

3.3

Suppose we observe from a given position in the genome. We keep observing until we find the nucleotide G. Describe the sample space of the observed DNA sequence. What is the probability that the DNA sequence observed before observing G has length 10?

$$S = \{G, *G, **G, ***G, \ldots\}$$

 $P(\text{length sequence before observing G} = 10) = (1 - 0.25)^{10}(0.25)$

 $(1-0.25)^{\circ}(10) * (0.25)$

[1] 0.01407838

3.4

Now we remove the independence assumption. Suppose that the probability of observing A,T,C or G given the previous nucleotide being C, is respectively 0.2, 0.2, 0.5, 0.1. In general in different locations of the DNA sequence, this conditional probabilities will be different and this fact is important for biological sequence detecting.

Suppose we observe C at a given location, what is the probability of observing the motif CG immediately after?

P(observing CG starting from position 2|observe C in position 1) =?

Lets introduce three extended random variable D_1 , D_2 , D_3 , indicating the nucleotide in position 1,2 and 3.

$$D_1, D_2, D_3$$

We want to compute,

$$P(D_2 = C, D_3 = G|D_1 = C) = P(D_3 = G|D_2 = C, D_1 = C)P(D_2 = C|D_1 = C) = 0.1 \times 0.5 = 0.05$$

Suppose now in a different region of the DNA sequence, the conditional probabilities of A, T, C, G given C become 0.1, 0.2, 0.3, 0.4, then what is in that region the probability of observing CG given that we observed C in the previous location?

Similarly,

$$P(D_2 = C, D_3 = G|D_1 = C) = P(D_3 = G|D_2 = C, D_1 = C)P(D_2 = C|D_1 = C) = 0.4 \times 0.3 = 0.12$$

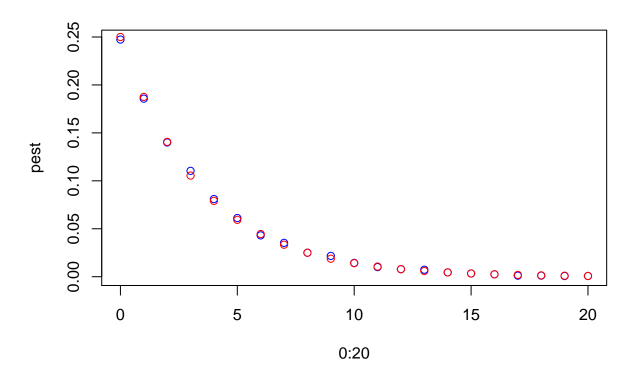
Ex 4 Some simulations

4.1

})

```
simDNA <- function(length = 1, nucleotides = c("A", "B", "C", "G")){</pre>
  sample(nucleotides, size = length, replace = TRUE)
}
estMotifProb <- function(length = 5, motif = c("A", "C", "G"), N = 1000){
  S<-t(replicate(N, simDNA(length)))</pre>
  sum(apply(S, MARGIN = 1, FUN = function(dna){
    length(grep(pattern = paste0(motif,collapse = ""),
                 x = paste0(dna, collapse = ""), value = FALSE)) > 0
  } )) / N
}
estMotifProb(5, c("A", "C", "G"), 1000)
## [1] 0.041
4.2
tryBefore <- function(before = "G", max = Inf){</pre>
 x \leftarrow simDNA(1)
 k <- 0
  while (x != before \& k <= max ){
    x \leftarrow simDNA(1)
    k <- k + 1
  }
  return(k)
N <- 100000
sum(replicate(N, tryBefore("G", 11)) == 10) / N
## [1] 0.01394
4.3
N <- 10000
##estimation as before
pest <- sapply(0:20, function(i){</pre>
  sum(replicate(N, tryBefore("G", i + 1)) == i) / N
})
## real prob
preal <- sapply(1:21, function(i){</pre>
  return(0.75^(i-1) * 0.25)
```

```
plot(0:20,pest, col = "blue")
points(0:20, preal, col = "red")
```



Ex 5 ISI data set

We load the data

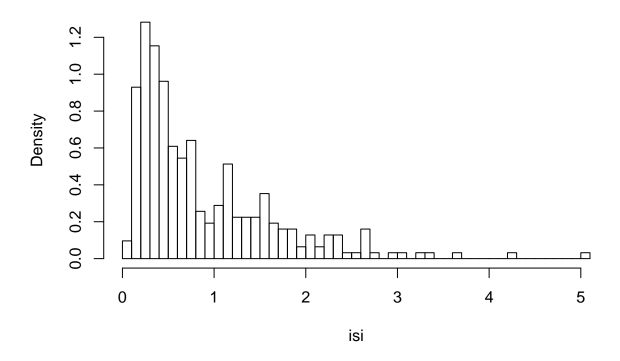
```
isidata <- read.table("neuronspikes.txt", col.names = "isi")
isi <- isidata$isi ##here we have the vector of observations</pre>
```

5.1

Plotting the histogram

```
hist(isi, breaks = 50, probability = TRUE)
```

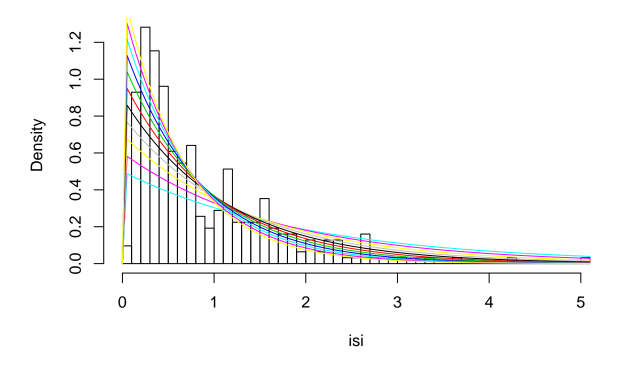
Histogram of isi



5.2

```
hist(isi, breaks = 50, probability = TRUE)
for (rate in seq(0.5, 1.5, 0.1)){
   curve(dexp(x, rate = rate), add = TRUE, col = rate*10)
}
```

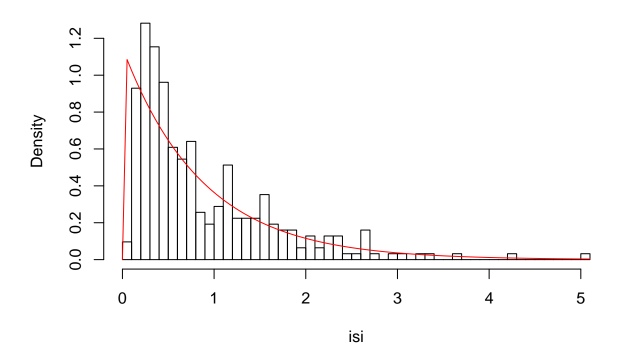
Histogram of isi



Maybe a nice parameter can be $\lambda=1.15$

```
hist(isi, breaks = 50, probability = TRUE)
curve(dexp(x, rate = 1.15), col = "red", add = TRUE)
```

Histogram of isi



5.3

```
rate <- 1.15

pexp(1, rate = rate)

## [1] 0.6833632

pexp(1.5, rate = rate) - pexp(0.5, rate = rate)

## [1] 0.3845318</pre>
```

Ex 6 Brain cell types data set

We load the brain cell data set,

```
cells <- read.csv("cell_types.csv", na.strings = "")</pre>
```

6.1

We use the parameter na.strings = "" to specify which string (in this case the empty strig) is used as a placeholder for the NA value.

6.2

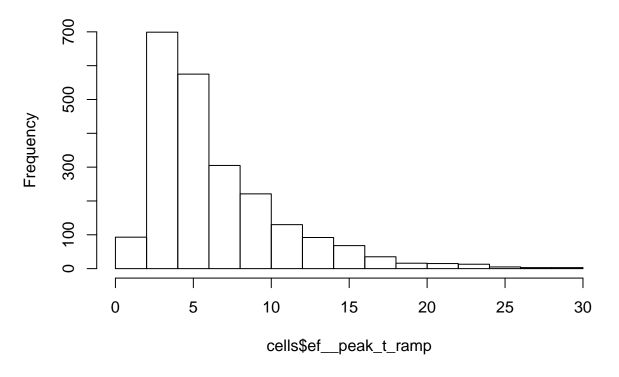
To find which observations are from human donors or mice, firts of all we locate the appropriate variable in the data, and secondly we check which are the possible values.

Lets list all the variables,

colnames(cells)

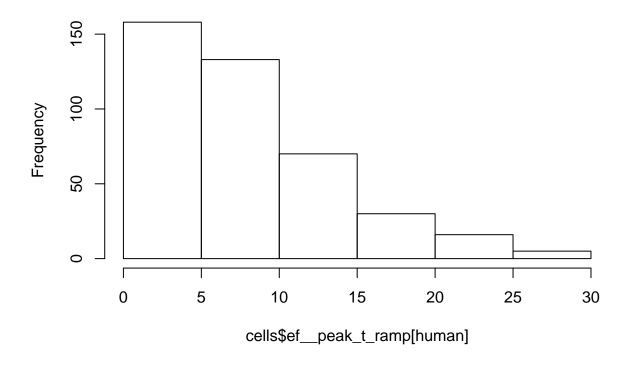
```
[1] "line_name"
##
    [2] "specimen_id"
##
##
    [3] "specimen__name"
       "specimen_hemisphere"
##
    [4]
##
   [5] "structure__id"
##
   [6] "structure__name"
##
    [7] "structure_acronym"
##
       "structure_parent__id"
   [9] "structure_parent__acronym"
## [10] "structure__layer"
  [11]
        "nr__max_euclidean_distance"
## [12] "nr__number_stems"
## [13] "nr__number_bifurcations"
## [14] "nr__average_contraction"
## [15] "nr__average_parent_daughter_ratio"
## [16] "nr__reconstruction_type"
## [17] "nrwkf id"
## [18] "erwkf__id"
## [19] "ef__fast_trough_v_long_square"
## [20] "ef_upstroke_downstroke_ratio_long_square"
## [21] "ef__adaptation"
## [22] "ef__f_i_curve_slope"
## [23] "ef__threshold_i_long_square"
## [24] "ef__tau"
## [25] "ef__avg_isi"
  [26] "ef_avg_firing_rate"
       "ef__ri"
  [27]
  [28]
       "ef__peak_t_ramp"
## [29] "ef__vrest"
## [30]
       "si__height"
## [31] "si__width"
## [32] "si path"
## [33] "csl x"
       "csl__y"
## [34]
## [35] "csl__z"
  [36] "csl__normalized_depth"
## [37] "cell_reporter_status"
## [38] "m__glif"
## [39] "m__biophys"
## [40] "m_biophys_perisomatic"
## [41] "m_biophys_all_active"
## [42] "tag__apical"
## [43] "tag__dendrite_type"
  [44] "morph_thumb_path"
  [45] "ephys_thumb_path"
## [46] "ephys_inst_thresh_thumb_path"
```

```
## [47] "donor__age"
## [48] "donor__sex"
## [49] "donor__disease_state"
## [50] "donor__race"
## [51] "donor_years_of_seizure_history"
## [52] "donor__species"
## [53] "donor id"
## [54] "donor__name"
we are interested in donor_species
Since string are converted to factors ina data.frame (thanks to read.csv function), we can check the levels,
levels(cells$donor_species)
## [1] "Homo Sapiens" "Mus musculus"
So we can extract the observations from humans and from mice,
human <- cells$donor species == levels(cells$donor species)[1]
mouse <- cells$donor_species == levels(cells$donor_species)[2]</pre>
We check how many they are,
sapply( list(human = human, mouse = mouse), sum)
## human mouse
     413 1920
The proportion of human cells in the data set is,
sum(human) / nrow(cells)
## [1] 0.1770253
6.3
Histogram for the ramp spike time,
hist(cells$ef__peak_t_ramp)
```



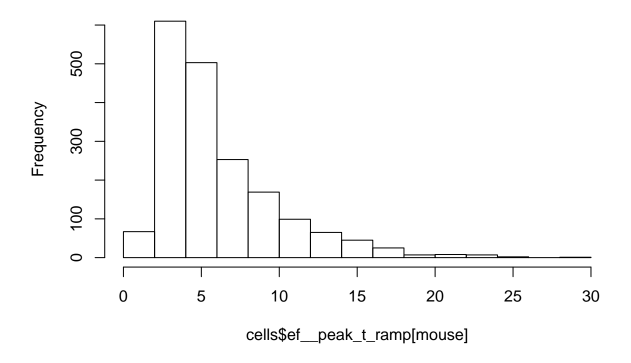
Separately for human cells,

hist(cells\$ef__peak_t_ramp[human])



and mice,

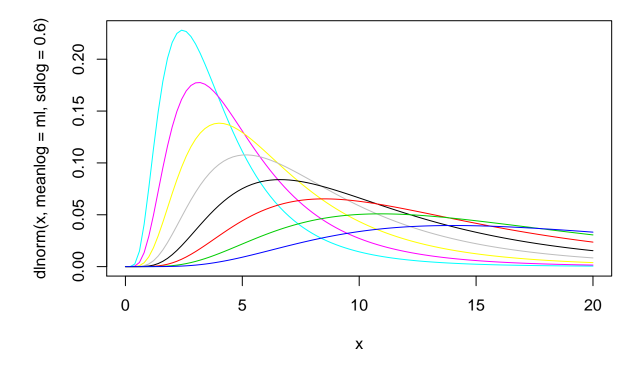
hist(cells\$ef__peak_t_ramp[mouse])



6.4

Plotting log-normal densities,

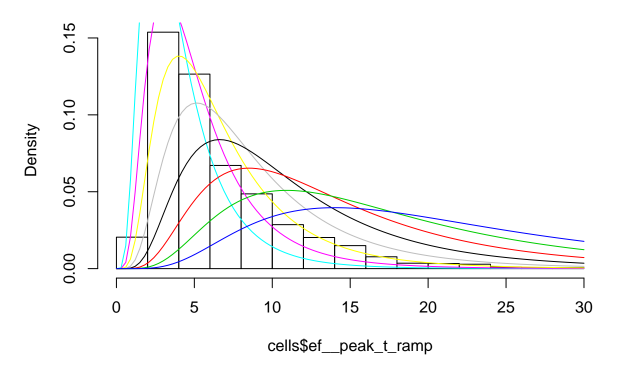
```
firstplot <- F
for (ml in seq(1.25, 3, 0.25)){
   curve(dlnorm(x,meanlog = ml, sdlog = 0.6),
        from = 0, to = 20, add = firstplot, col = ml*4)
   firstplot <- TRUE
}</pre>
```



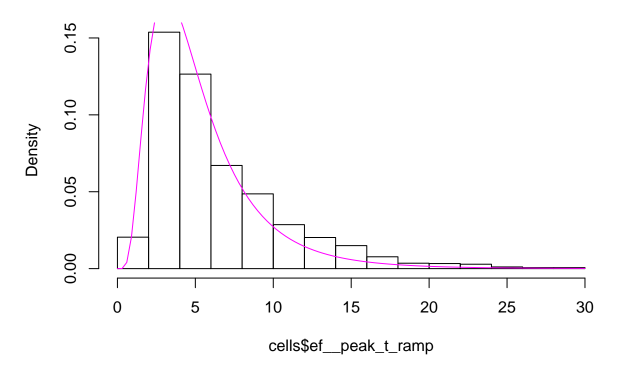
6.5

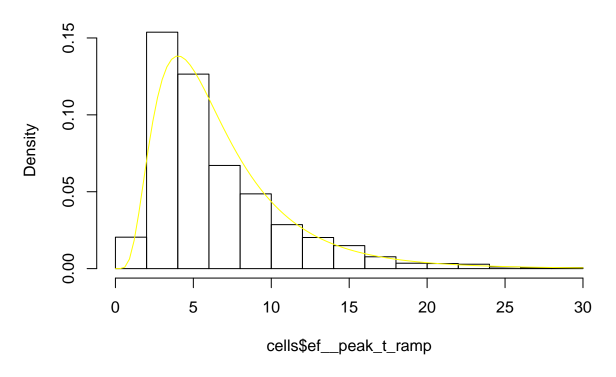
Plot densities on top of histogram,

```
hist(cells$ef__peak_t_ramp, freq = FALSE)
for (ml in seq(1.25, 3, 0.25)){
  curve(dlnorm(x,meanlog = ml, sdlog = 0.6),
      add = TRUE, col = ml*4 )
}
```



Probably meanlog = 1.5 or 1.75





6.6

Only on top of the human observations,

