## **Lab 1 Probability Distributions**

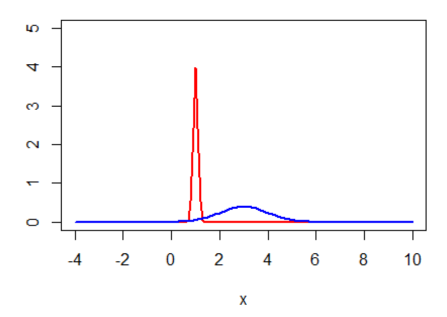
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#### **Table of Contents**

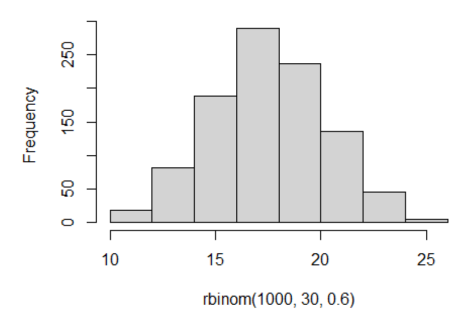
#### Lab 1 Lab Manual Exercise

copy and paste your work by following each example from the lab manual for this exercise



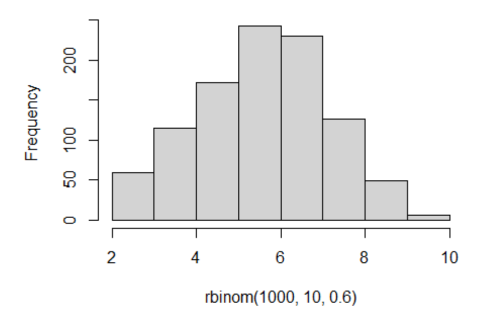
```
# # Function Syntax
# function_name <- function(arg_1, arg_2, ...) {</pre>
     Function body
#
# }
# Calculate the 60th %ile of the standard normal.
qnorm(0.6,0,1)
## [1] 0.2533471
# Calculate the probability that a value lies below 0.8 in the standard
normal distribution
pnorm(0.8,0,1)
## [1] 0.7881446
# Draw 1000 samples of 30 coin tosses with p(heads) = 0.6 # and plot the
distribution
# Syntax: rbinom (# observations, # trials per observation, probability of
success )
hist(rbinom(1000,30,0.6))
```

# Histogram of rbinom(1000, 30, 0.6)

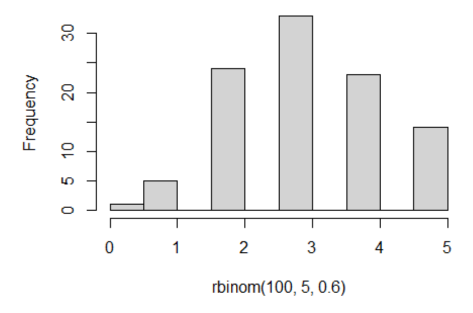


# Do the above with only 10 trials per observation
hist(rbinom(1000,10,0.6))

# Histogram of rbinom(1000, 10, 0.6)

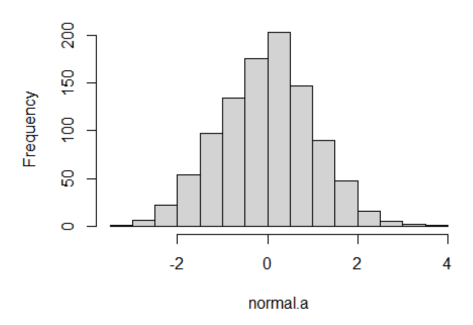


## Histogram of rbinom(100, 5, 0.6)



```
# Transformations between probability distributions
# generate 1000 trials from a normal distribution
normal.a <- rnorm( n=1000, mean=0, sd=1 )
hist( normal.a )</pre>
```

#### Histogram of normal.a

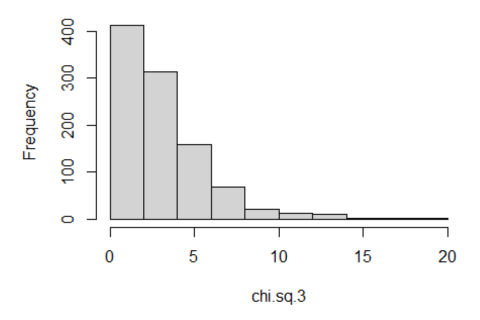


```
#next, we generate a chi-square distribution with 3 #degrees of freedom:
normal.b <- rnorm( n=1000 )  # another set of normally distributed data
normal.c <- rnorm( n=1000 )  # and another!

# Take the SUM of SQUARES of the above 3 normally distributed variables a, b,
and c
chi.sq.3 <- (normal.a)^2 + (normal.b)^2 + (normal.c)^2

# and the resulting chi.sq.3 variable should contain 1000 observations that
follow a chi-square distribution with 3 degrees of freedom. You can use the
hist() function to have a look at these observations yourself
hist(chi.sq.3)</pre>
```

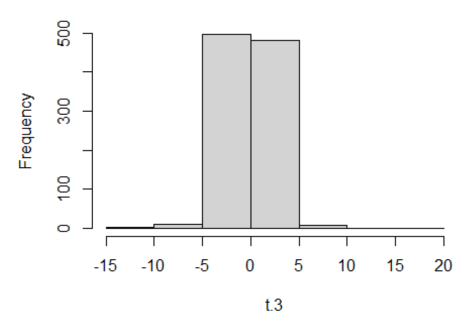
## Histogram of chi.sq.3



```
## Now how do we get to a t-distribution from Normal and chi-sq
distributions?
# First, take a scaled chi-sq by dividing it by the degrees of freedom
scaled.chi.sq.3 <- chi.sq.3 / 3
# Then take a normally distributed variable and divide them by the square
root of the scaled chi-sq variable to get a t-distribution with the same
degrees of freedom

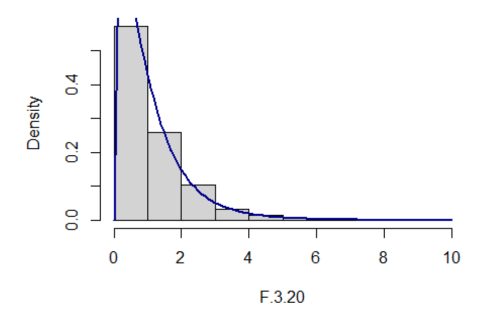
normal.d <- rnorm( n=1000 )  # yet another #set of normally
distributed data
t.3 <- normal.d / sqrt( scaled.chi.sq.3 ) # divide by #square root of scaled
chi-square to get t
hist (t.3)</pre>
```

### Histogram of t.3



```
## To get to an F distribution, take the ratio between two scaled chi-sq
distributions.
# F distribution with 3 and 20 degrees of freedom:
# first take two chi-sq variables, with 3 dof and 20 dof respectively, and
take the ratio:
chi.sq.20 <- rchisq( 1000, 20)
                                               # generate chi square data
with df = 20...
scaled.chi.sq.20 <- chi.sq.20 / 20</pre>
                                               # scale #the chi square
variable...
F.3.20 <- scaled.chi.sq.3 / scaled.chi.sq.20 # take the ratio of the two
chi squares...
hist( F.3.20, freq = FALSE)
                                                             # ... and draw a
picture
curve(df(x, 3, 20),
      col="darkblue", lwd=2, add=TRUE, yaxt="n")
```

#### Histogram of F.3.20



## The curve above confirms this looks similar if you use the R built-in function df (just like dnorm, but for the F distribution)

#### Lab 1 Generalization exercises

use the code from above to attempt to solve the extra things we ask you do for this assignment

```
# Q1 Plot a normal distribution with mean = 2, s.d. = 0.4

# Q2 Calculate the 85th %ile of the above distribution.

# Q3 Calculate the probability that a value lies in between 1 and 2 given the above distribution

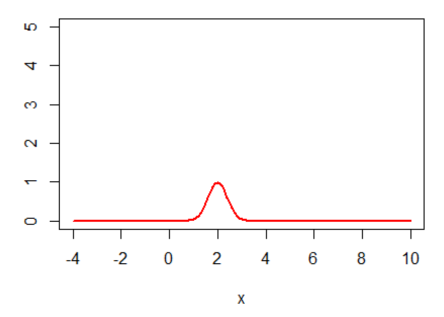
# Q4 Plot a simulated t-distribution with 5 degrees of freedom.

# Q5 Plot a chi-sq distribution with 5 degrees of freedom.
```

### Lab 1 Written answer question

The following is the code for Q1

```
x <- seq(-4, 10, 0.1)
plot(x, dnorm(x, mean = 2, sd = 0.4), type = "1",
    ylim = c(0, 5), ylab = "", lwd = 2, col = "red")</pre>
```



The following is the code for Q2

```
qnorm(0.85, mean=2, sd=0.4)
## [1] 2.414573
```

The following is the code for Q3

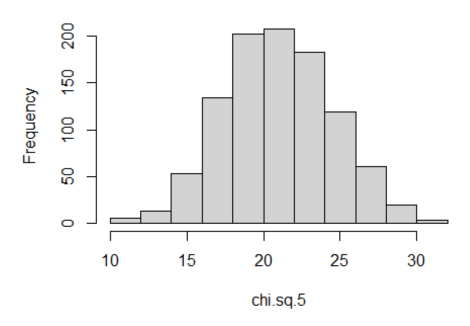
```
a <- pnorm(1, 2, 0.4)
b <- pnorm(2, 2, 0.4)
prob.between.1.and.2 <- b - a
print(prob.between.1.and.2)
## [1] 0.4937903</pre>
```

The following is the code for Q4

```
normal.a <- rnorm( n=1000, mean=2, sd=0.4 )
normal.b <- rnorm( n=1000 , mean=2, sd=0.4 )
normal.c <- rnorm( n=1000 , mean=2, sd=0.4 )
normal.d <- rnorm(n =1000 , mean=2, sd=0.4)
normal.e <- rnorm (n = 1000 , mean=2, sd=0.4)
chi.sq.5 <- (normal.a)^2 + (normal.b)^2 + (normal.c)^2 + (normal.d)^2 +</pre>
```

```
(normal.e)^2
hist(chi.sq.5)
```

# Histogram of chi.sq.5



```
scaled.chi.sq.5 <- chi.sq.5 / 5
normal.f <- rnorm(n = 1000 , mean=2, sd=0.4)
t.5 <- normal.f / sqrt( scaled.chi.sq.5 )
hist (t.5)</pre>
```

# Histogram of t.5

