Math 475: Final Exam

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Problem 1

part a

Show the outputs of the EM algorithm are consistent with the given parameter equations To find the updated parameters (i.e. the maximized value) we fist need to find the Q-function (E step) then maximize the Q function by taking the derivative in regard to each of the parameters (M step). E step: Given the observed likelihood, we can compute the complete likelihood to be:

$$L(\theta|n_{k,i}) = \prod_{i=0}^{16} \frac{[z(\theta)^{(n_0)} t(\theta)^{n_i} p(\theta)^{n_i}]}{i!}$$

Given the likelihood equation, we can work out the log likelihood to be:

$$log[L(\theta|n_{k,i})] = \sum_{i=0}^{16} z_0 log(\alpha 1_{i=0}) +$$

$$(t_i)[log(\alpha 1_{i=0}) + log(\beta \mu^i e^{-\mu}) + log((1 - \alpha - \beta)\lambda^i e^{-\lambda})] +$$

$$(p_i)[log(\alpha 1_{i=0}) + log(\beta \mu^i e^{-\mu}) + log((1 - \alpha - \beta)\lambda^i e^{-\lambda})] - log(i!)$$

where y is the complete data set and z_0, t_i, p_i represent three different groups. These are further broken down into the zero, typical, and promiscuous groups. To find your Q-function, take the expectation of the log likelihood function:

$$Q(\theta|\theta^{(t)}) = n_{z,0}^{(t)} \sum_{i=0}^{16} log(\alpha 1_{i=0}) + n_{t,i}^{(t)} \sum_{i=0}^{16} log(\beta \mu^i exp(-\mu)) + n_{n,i}^{(t)} \sum_{i=0}^{16} log((1-\alpha-\beta)\lambda^i exp(\lambda))$$

M step: For the M step of the EM algorithm, we need to maximize the Q function in regard to each parameter then set it equal to zero.

When the derivative is set equal to zero, we find that the updated parameters equal to what we expected:

$$\begin{split} \alpha^{(t+1)} &= \frac{n_0 z_0(\theta^{(t)} t)}{N} \\ \beta^{(t+1)} &= \Sigma_{i=0}^{16} \frac{n_i t_i(\theta^{(t)})}{N} \\ \mu^{(t+1)} &= \frac{\Sigma_{i=0}^{16} i n_i t_i(\theta^{(t)})}{\Sigma_{i=0}^{16} n_i t_i(\theta^{(t)})} \\ \lambda^{(t+1)} &= \frac{\Sigma_{i=0}^{16} i n_i p_i(\theta^{(t)})}{\Sigma_{i=0}^{16} n_i p_i(\theta^{(t)})} \end{split}$$

part b and c

```
set.seed(475)
# initialize variables
data = data.frame(enc = 0:16, freq = c(379, 299, 222, 145, 109,
          95, 73, 59, 45, 30, 24, 12, 4, 2, 0, 1, 1))
N = sum(data$freq)
y = rep(data$enc, data$freq)
alpha = 0.5
beta = 0.8
mu = 2
lambda = 15
param = c(alpha, beta, mu, lambda)
param.guess = c(0.1, 0.2, 3, 4)
tol = 1e-10
tol.cur = 100
time = 0
i = 0:16
# functions
log.likelihood <- function(alpha, beta, mu, lambda, x) {</pre>
          1 = 0
          alpha = exp(alpha)/(1 + exp(alpha))
          beta = \exp(beta)/(1 + \exp(beta))
          mu = \exp(mu)/(1 + \exp(mu))
          lambda = \exp(lambda)/(1 + \exp(lambda))
          for (i in 1:length(x$enc)) {
                   e = x enc[i]
                   n = x freq[i]
                    if (e == 0) {
                              l = l + n * log(alpha + beta * exp(-mu) + (1 - alpha -
                                        beta) * exp(-lambda))
                             print(1)
                    } else {
                              1 = 1 + n * log(beta * (mu^e) * exp(-mu) + (1 - alpha - alph
                                        beta) * exp(-lambda) * lambda^e) - log(factorial(e))
                    }
          }
          return(1)
}
# EM Algorithm
while (tol.cur > tol) {
          pi = (beta * exp(-mu) * mu^i) + ((1 - alpha - beta) * exp(-lambda) *
                    lambda^i)
          pi[1] = pi[1] + alpha
          z.stat = alpha/(pi[1])
          t.stat = (beta * (mu^i) * exp(-mu))/pi
          p.stat = ((1 - alpha - beta) * exp(-lambda) * lambda^i)/pi
          alpha = (data$freq[1] * z.stat)/N
          beta = sum(data$freq * t.stat)/N
          mu = sum(i * data$freq * t.stat)/sum(data$freq * t.stat)
          lambda = sum(i * data$freq * p.stat)/sum(data$freq * p.stat)
          new.param = c(alpha, beta, mu, lambda)
          tol.cur = sum(abs(new.param - param))
```

```
param = new.param
    time = time + 1
}
param
## [1] 0.1221661 0.5625419 1.4674746 5.9388889
hist(rep(data\$enc, data\$freq), breaks = -0.5 + c(0:17), freq = F,
    main = "Histogram of Risky Sexual Encounters", xlab = "Encounters")
# hist(y,freq=F)
z = 0:16
prob = (beta * exp(-mu) * mu^z + (1 - alpha - beta) * exp(-lambda) *
    lambda^z)/(factorial(z))
prob[1] = prob[1] + alpha
for (i in 1:length(z)) {
    lines(c(z[i] - 0.1, z[i] - 0.1), c(0, prob[i]), lwd = 5,
        col = 1)
}
pois.hat = mean(y)
for (i in 1:length(z)) {
```

Histogram of Risky Sexual Encounters

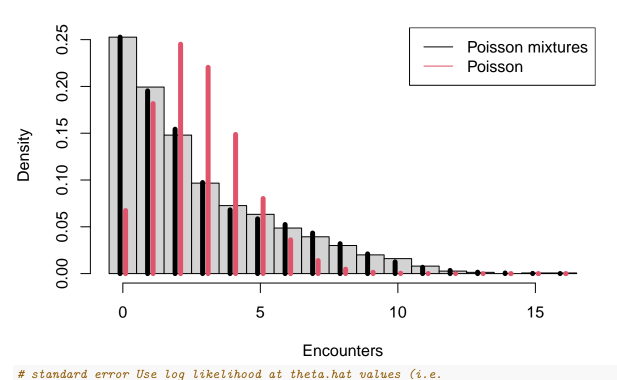
lines(c(z[i] + 0.1, z[i] + 0.1), c(0, dpois(z[i], pois.hat)),

legend("topright", c("Poisson mixtures", "Poisson"), lty = 1,

lwd = 5, col = 2)

col = 1:2)

}



```
# new parameter values)
set.seed(1234)
data = data.frame(enc = 0:16, freq = c(379, 299, 222, 145, 109,
    95, 73, 59, 45, 30, 24, 12, 4, 2, 0, 1, 1))
tol = 1e-10
B = 1000
result.boot <- NULL
alpha <- 1
beta <- 2
mu <- 4
lambda <- 10
param <- c(alpha, beta, mu, lambda)
for (j in 1:B) {
    # randomize samples
    data.boot <- rmultinom(1, sum(data$freq), prob = data$freq/length(data$freq))</pre>
    # set initial values
    tol.cur <- 100
    N <- sum(data.boot)</pre>
    i = c(0:16)
    # loop
    while (tol.cur > tol) {
        pi = (beta * exp(-mu) * mu^i) + ((1 - alpha - beta) *
            exp(-lambda) * lambda^i)
        pi[1] = pi[1] + alpha
        z.stat = alpha/(pi[1])
        t.stat = (beta * (mu^i) * exp(-mu))/pi
        p.stat = ((1 - alpha - beta) * exp(-lambda) * (lambda^i))/pi
        alpha = (data.boot[1] * z.stat)/N
        beta = sum(data.boot * t.stat)/N
        mu = sum(i * data.boot * t.stat)/sum(data.boot * t.stat)
        lambda = sum(i * data.boot * p.stat)/sum(data.boot *
            p.stat)
        new.param = c(alpha, beta, mu, lambda)
        tol.cur = sum(abs(new.param - param))
        param = new.param
    }
    result.boot <- rbind(result.boot, param)</pre>
}
result.boot[B, ]
## [1] 0.1365674 0.5705522 1.6564927 6.2467511
cov(result.boot) #covariance matrix to show standard error
##
                 [,1]
                                [,2]
                                             [,3]
## [1,] 0.0003955050 -0.0001790008 0.0015697232 0.001484594
## [2,] -0.0001790008  0.0004392937  0.0001291111  0.001400945
## [3,] 0.0015697232 0.0001291111 0.0121965277 0.013091738
## [4,] 0.0014845938 0.0014009455 0.0130917380 0.038778808
# pairwise correlation
cor(result.boot)
```

```
## [1,] 1.000000 -0.42943889 0.71470853 0.3790831
## [2,] -0.4294389 1.0000000 0.05577864 0.3394271
## [3,] 0.7147085 0.05577864 1.0000000 0.6019799
## [4,] 0.3790831 0.33942709 0.60197988 1.0000000
```

part a: Metropolis and M-H Algorithm

We are seeking to estimate $\alpha, \beta, \mu, \lambda$ using the Metropolis Algorithm.

Because the proposal distribution is symmetric, it can be canceled out in the ratio calculation, and we can then focus on the ratio of the target distribution with theta star and the previous theta value.

part 2: MCMH

```
# Metropolis Samples (Symmetric Proposal Distribution)
set.seed(575)
# initialize variables and constants
data = data.frame(enc = 0:16, freq = c(379, 299, 222, 145, 109,
           95, 73, 59, 45, 30, 24, 12, 4, 2, 0, 1, 1))
N = sum(data\$freq)
y = rep(data$enc, data$freq)
reject <- 0
alpha = rep(0, N)
beta = rep(0, N)
mu = rep(0, N)
lambda = rep(0, N)
sigma = 2
# functions
log.likelihood <- function(alpha, beta, mu, lambda, x) {</pre>
           1 = 0
            # reparameterization
            alpha = exp(alpha)/(1 + exp(alpha))
           beta = \exp(beta)/(1 + \exp(beta))
           mu = exp(mu)
           lambda = exp(lambda)
           for (i in 1:length(x$enc)) {
                       e = x enc[i]
                      n = x freq[i]
                       if (e == 0) {
                                   l = l + n * log(alpha + beta * exp(-mu) + (1 - alpha -
                                              beta) * exp(-lambda))
                       } else {
                                   1 = 1 + n * log(beta * (mu^e) * exp(-mu) + (1 - alpha - alph
                                              beta) * exp(-lambda) * (lambda^e)) - n * log(factorial(e))
           }
           return(1)
pi_i <- function(alpha, beta, mu, lambda, i) {</pre>
            if (i == 0)
                       return(alpha + beta * exp(-mu) + (1 - alpha - beta) *
                                   exp(-lambda)) else return(beta * mu^i * exp(-mu) + (1 - alpha - beta) *
                       lambda^i * exp(-lambda))
LogL <- function(alpha, beta, mu, lambda, df) {</pre>
           sum <- 0
```

```
alpha = exp(alpha)/(1 + exp(alpha))
    beta = \exp(beta)/(1 + \exp(beta))
   mu = exp(mu)
   lambda = exp(lambda)
   for (i in length(df)) {
        sum <- sum + df[i] * (log(pi_i(alpha, beta, mu, lambda,</pre>
            i)) - log(factorial(i)))
   return(sum)
}
# likelihood <- function{alpha,beta,mu,lambda}{ } #initialize</pre>
# sample values i.e chains alpha[1] = rnorm(1,3,sigma)
# beta[1] = rnorm(1,3,sigma) mu[1] = rnorm(1,3,sigma)
# lambda[1] = rnorm(1,3,siqma) for(i in 2:N){ #sample from
# proposal distribution (symmetrical) alpha.star <-</pre>
\# rnorm(1, mean = alpha[i-1], sd = sigma) beta.star <-
\# rnorm(1, mean = beta[i-1], sd = sigma) mu.star <-
\# rnorm(1, mean = mu[i-1], sd = sigma) \ lambda.star <-
\# rnorm(1, mean = lambda[i-1], sd = siqma)
# print(c(alpha.star,beta.star,mu.star,lambda.star)) num <-</pre>
# LogL(alpha.star,beta.star,mu.star,lambda.star,data$freq)
# dem <-
\# LogL(alpha[i-1],beta[i-1],mu[i-1],lambda[i-1],data\$freq)
\# ratio <- exp(num - dem) accept.prob <- min(1, ratio) U =
# runif(1) if(U < accept.prob){ alpha[i] <- alpha.star</pre>
# beta[i] <- beta.star mu[i] <- mu.star lambda[i] <-</pre>
\# lambda.star \} else{ alpha[i] <- alpha[i-1] beta[i] <-}
\# = reject + 1 \} alpha = exp(alpha)/(1+exp(alpha)) beta =
\# exp(beta)/(1+exp(beta)) mu = exp(mu) lambda = exp(lambda)
c(mean(alpha), mean(beta), mean(mu), mean(lambda))
## [1] 0 0 0 0
print("Rejection Rate:")
## [1] "Rejection Rate:"
100 * (reject/N)
## [1] 0
```

part a:

From the given data for the clinical trial, we can see from the box-and-whisker plot that the Hormone group with the censored time has the most patients out of the four groups. Not only that, but the means are more spread apart within the hormone group (difference of 13.68) compared to the control group (difference of 7.2). Based on the combined Normal QQ plot, the Control Group with Censored time most closely follows a normal distribution ("3" from list.id legend). The two from the hormone group of Recurrence and Censor times (0 and 1, respectively), do not closely align with a normal distribution. ## part b: Given the likelihood and prior, we can use those to calculate the conditional distributions of θ and τ . To do this we will first need to calculate the joint probability density function:

$$P(y|\theta,\tau) \propto L(\theta,\tau|y) f(\theta,\tau)$$

$$\pi(\theta,\tau|y) \propto \pi(\theta,\tau,y) = L(\theta,\tau|y) f(\theta,\tau)$$

$$P(\theta,\tau|y) = \theta^{\sum \delta_i^c + \sum \delta_i^H + 2a} \tau^{\sum \delta_i^H + b} exp(-\theta(\sum x_i^C + c) - \tau \theta(\sum x_i^H + d))$$

With this we can calculate the conditional distributions for the parameters:

$$P(\tau|\theta, y) = \tau^{\sum \delta_i^H + b} exp(-\tau\theta(\sum x_i^H + d))$$

$$\propto Gamma(\tau|\sum \delta_i^H + b + 1, \sum x_i^C \sum x_i^H + c + d)$$

$$P(\theta|\tau, y) = \theta^{\sum \delta_i^c + \sum \delta_i^H + a} exp(-\theta(\sum x_i^C + c) - \tau\theta(\sum x_i^H + d))$$

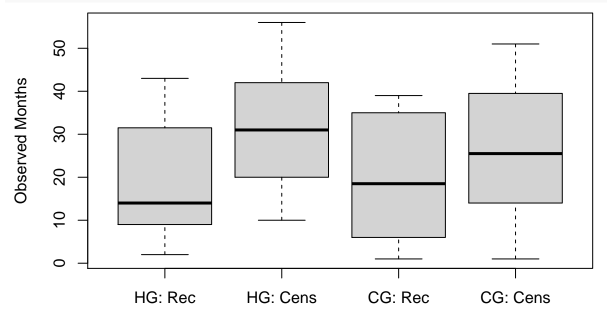
$$\propto Gamma(\theta|\sum \delta_i^c + \sum \delta_i^H + a + 1, \sum x_i^C + \tau \sum x_i^H + c + d\tau)$$

```
set.seed(12345)
library(ggplot2)
library(Rmisc)
```

Loading required package: lattice

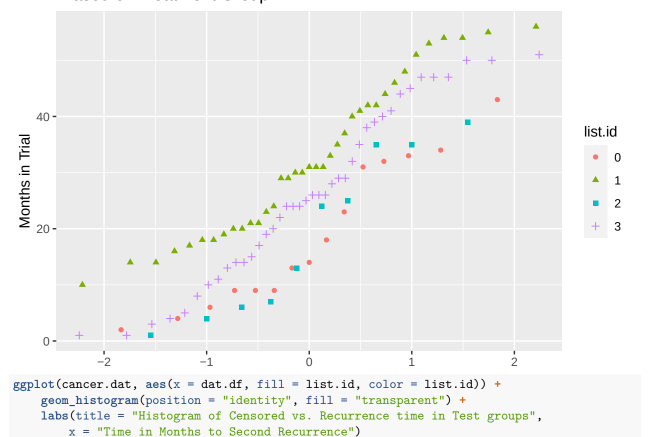
Loading required package: plyr

```
# initialize data sets
hormone.rec <- c(2, 4, 6, 9, 9, 9, 13, 14, 18, 23, 31, 32, 33,
    34, 43)
lab.h.r <- rep(0, length(hormone.rec))</pre>
hormone.cens \leftarrow c(10, 14, 14, 16, 17, 18, 18, 19, 20, 20, 21,
    21, 23, 24, 29, 29, 30, 30, 31, 31, 31, 33, 35, 37, 40, 41,
    42, 42, 44, 46, 48, 51, 53, 54, 54, 55, 56)
lab.h.c <- rep(1, length(hormone.cens))</pre>
control.rec = c(1, 4, 6, 7, 13, 24, 25, 35, 35, 39)
lab.c.r <- rep(2, length(control.rec))</pre>
control.cens = c(1, 1, 3, 4, 5, 8, 10, 11, 13, 14, 14, 15, 17,
    19, 20, 22, 24, 24, 24, 25, 26, 26, 26, 28, 29, 29, 32, 35,
    38, 39, 40, 41, 44, 45, 47, 47, 47, 50, 50, 51)
lab.c.c <- rep(3, length(control.cens))</pre>
list.id <- c(lab.h.r, lab.h.c, lab.c.r, lab.c.c)</pre>
dat.df <- c(hormone.rec, hormone.cens, control.rec, control.cens)</pre>
cancer.dat <- data.frame(dat.df, list.id)</pre>
cancer.dat$list.id <- as.factor(cancer.dat$list.id)</pre>
# part a: Plots and quantiles
```



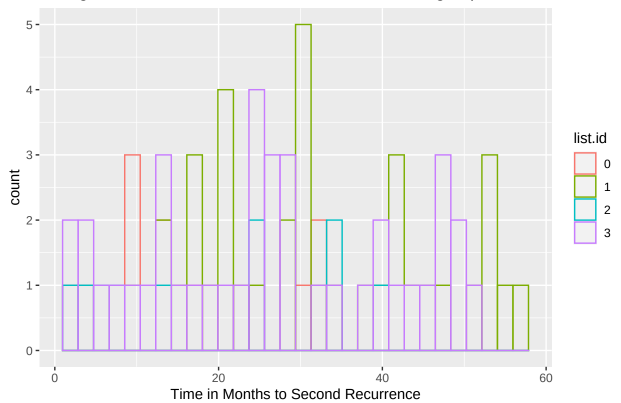
Patient Group

Months in Trial Based on Treatment Group



`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.

Histogram of Censored vs. Recurrence time in Test groups



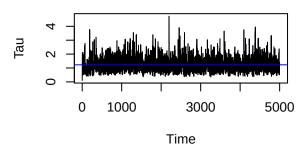
```
# part c: Gibbs Sampler Initialize and prepare recurrence and
# censored data based on patient group
delta.h <- c(rep(1, length(hormone.rec)), rep(0, length(hormone.cens)))</pre>
delta.c <- c(rep(1, length(control.rec)), rep(0, length(control.cens)))</pre>
dat.hormone <- data.frame(x = c(hormone.rec, hormone.cens), delta.h)</pre>
dat.control <- data.frame(x = c(control.rec, control.cens), delta.c)</pre>
a = 3
b = 1
c = 60
d = 120
# FUNCTIONS
gibbs <- function(dat.control, dat.hormone, n, burn, hyperparameter) {</pre>
    mat <- matrix(ncol = 2, nrow = n)</pre>
    tau <- 0.1
    theta <- 0.1
    mat[1, ] <- c(theta, tau)</pre>
    a <- hyperparameter[1]
    b <- hyperparameter[2]</pre>
    c <- hyperparameter[3]</pre>
    d <- hyperparameter[4]</pre>
    for (i in 2:n) {
        tau <- mat[i - 1, 2]
        one.tau <- a + 1 + sum(dat.control$delta.c) + sum(dat.hormone$delta.h)
        two.tau <- sum(dat.control$x) + (tau * sum(dat.hormone$x)) +
             c + (d * tau)
        mat[i, 1] <- rgamma(1, one.tau, two.tau)</pre>
```

```
theta <- mat[i, 1]</pre>
        one.theta <- b + 1 + sum(dat.hormone$delta.h)</pre>
        two.theta <- theta * sum(dat.hormone$x) + (theta * d)</pre>
        mat[i, 2] <- rgamma(1, one.theta, two.theta)</pre>
    burn <- burn + 1
    return(mat[burn:n, ])
}
gibby <- gibbs(dat.control, dat.hormone, 1000, 100, c(3, 1, 60,
    120))
colMeans(gibby)
## [1] 0.009277404 1.253663033
gibbies <- gibbs(dat.control, dat.hormone, 5000, 0, c(3, 1, 60,
    120))
colMeans(gibbies)
## [1] 0.009430104 1.227765240
# Convergence diagnostics
par(mfrow = c(2, 2))
plot(ts(gibbies[, 1]), main = "Theta from 10000 Iterations",
    ylab = "Tau")
abline(h = mean(gibbies[, 1]), col = "red")
plot(ts(gibbies[, 2]), main = "Tau from 10000 Iterations", ylab = "Tau")
abline(h = mean(gibbies[, 2]), col = "blue")
hist(gibbies[, 1], 40, main = "Theta from 10000 Iterations",
    xlab = "Theta Estimates")
hist(gibbies[, 2], 40, main = "Tau from 10000 Iterations", xlab = "Tau Estimates")
```

Theta from 10000 Iterations

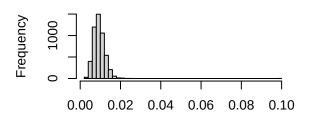
0.00 3000 5000 Time

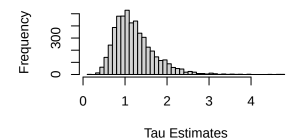
Tau from 10000 Iterations



Theta from 10000 Iterations

Tau from 10000 Iterations





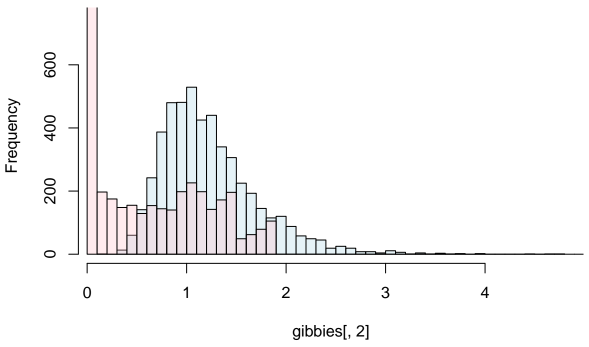
Theta Estimates

```
##
                             Theta
                                           Tau
## Marginal Mean
                      0.009430104 1.227765240
## Standard Deviation 0.002968410 0.484634152
## 95% CI upper
                      0.009347805 1.214328860
## 95% CI lower
                      0.009512402 1.241201620
# part e: graphing the prior and posterior distributions
# Posterior is the theta and tau estimates from above with
# the prior being a gamma distribution
par(mfrow = c(1, 1))
c1 <- rgb(173, 216, 230, max = 255, alpha = 80, names = "lt.blue")
c2 <- rgb(255, 192, 203, max = 255, alpha = 80, names = "lt.pink")
theta.pos <- hist(gibbies[, 1], breaks = 40, plot = FALSE)</pre>
tau.pos <- hist(gibbies[, 2], breaks = 40, plot = FALSE)</pre>
```

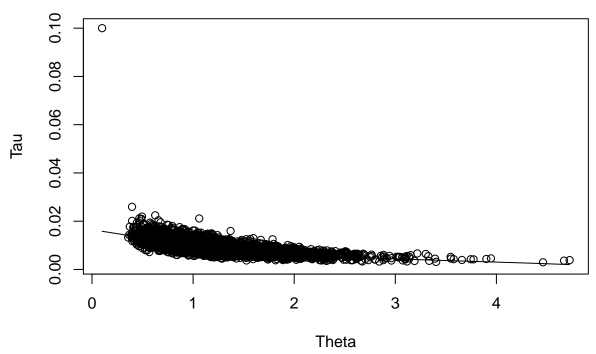
```
theta.prior <- hist(dexp(quantile(gibbies[, 1])), breaks = 40,
    plot = FALSE)
tau.prior <- hist(dgamma(quantile(gibbies[, 2]), shape = 1 +
    1 + sum(dat.hormone$delta.h), rate = gibbies[, 1] * sum(dat.hormone$x) +
    (gibbies[, 1] * 120)), breaks = 120, plot = FALSE)

plot(tau.pos, col = c1, main = "Histogram of Tau Posterior and Prior",
    ylim = range(1:750))
plot(tau.prior, col = c2, add = TRUE)</pre>
```

Histogram of Tau Posterior and Prior



Theta ~ Tau



cor(gibbies[, 2], gibbies[, 1])

```
## [1] -0.6475986
```

```
# inverse correlation, indicating that there is not
# part g: sensitivity analysis
hyp.half <- gibbs(dat.control, dat.hormone, 10000, 2500, c(1.5,
    2, 30, 60))
half.mean <- colMeans(hyp.half)
half.sd <- c(sd(hyp.half[, 1]), sd(hyp.half[, 2]))
half.ci.theta <- CI(hyp.half[, 1], ci = 0.95)
half.ci.tau \leftarrow CI(hyp.half[, 2], ci = 0.95)
hyp.dou <- gibbs(dat.control, dat.hormone, 10000, 2500, c(6,
    0.5, 120, 240))
dou.mean <- colMeans(hyp.dou)</pre>
dou.sd <- c(sd(hyp.dou[, 1]), sd(hyp.dou[, 2]))</pre>
dou.ci.theta \leftarrow CI(hyp.dou[, 1], ci = 0.95)
dou.ci.tau \leftarrow CI(hyp.dou[, 2], ci = 0.95)
# Summary statistics based on hyperparameters
hyp.table <- matrix(c(half.mean[1], half.mean[2], dou.mean[1],
    dou.mean[2], half.sd[1], half.sd[2], dou.sd[1], dou.sd[2],
    half.ci.theta[3], half.ci.tau[3], dou.ci.theta[3], dou.ci.tau[3],
    half.ci.theta[1], half.ci.tau[1], dou.ci.theta[1], dou.ci.tau[1]),
    ncol = 4, byrow = TRUE)
colnames(hyp.table) <- c("Theta, Halved", "Tau, Halved", "Theta, Doubled",</pre>
    "Tau Doubled")
rownames(hyp.table) <- c("Marginal Mean", "Standard Deviation",</pre>
```

```
"CI 95% Lower", "CI 95% Upper")
as.table(hyp.table)
```

part f and g:

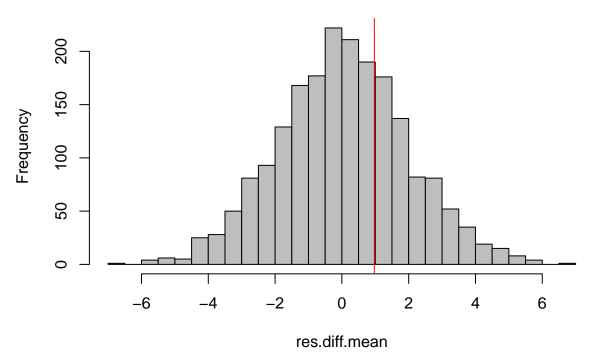
Based on the summary statistics produced in part d, I would say that the recurrence times are not statistically significant from the control group. This would mean that the clinical trial was not a huge success because the drug has very little affect on combating recurring episodes of breast cancer. We can see from the linear regression graph and the correlation output (-0.6475986) that the two parameters are inversely related.

For the hyperparameters, we can see from the table of summary statistics, that changing the hyperparameters affects our tau term more than the theta term. This can be seen in the 95% CI. This would be important to understand where these numbers come from and how to better estimate them before completing another clinical trial, because the hyperparameters could affect whether the drug is (statistically) effective.

```
set.seed(475)
# part b: mean recurrence time
m.r.h <- mean(hormone.rec)</pre>
m.r.c <- mean(control.rec)</pre>
cat("Mean Recurrence Time for Hormone Group", m.r.h, "\n")
## Mean Recurrence Time for Hormone Group 18.66667
cat("Mean Recurrence Time for Control Group", m.r.c, "\n")
## Mean Recurrence Time for Control Group 18.9
tau.est <- gibbies[, 2]</pre>
theta.est <- gibbies[, 1]</pre>
better \leftarrow c(0.032429, 0.76652)
hormone.estimate <- rexp(length(theta.est), rate = mean(tau.est) *
    mean(theta.est))
control.estimate <- rexp(length(theta.est), rate = mean(theta.est))</pre>
cat("Mean Recurrence Time for Hormone Group", mean(hormone.estimate),
## Mean Recurrence Time for Hormone Group 87.97597
cat("Mean Recurrence Time for Control Group", mean(control.estimate),
## Mean Recurrence Time for Control Group 104.0848
# part c:
B = 1000 #Bootstrap Iteration
alpha <- 0.05
means.hormone <- rep(0, B)
means.control <- rep(0, B)
for (i in 1:B) {
    means.hormone[i] <- mean(sample(hormone.estimate, size = 1:length(hormone.estimate),</pre>
        replace = TRUE))
    means.control[i] <- mean(sample(control.estimate, size = 1:length(control.estimate),</pre>
        replace = TRUE))
}
j \leftarrow (alpha/2) * B
k <- (1 - (alpha/2)) * B
cat("95% CI for Hormone Group:", c(means.hormone[j], means.hormone[k]),
   "\n")
## 95% CI for Hormone Group: 123.3696 103.0919
cat("95% CI for Control Group:", c(means.control[j], means.control[k]),
   "\n")
## 95% CI for Control Group: 68.94879 234.4083
# part d: Permutation Tests Use permutation tests to show
# that there is no difference between the mean recurrence
# times Null hypothesis: the mean recurrence time between the
# two groups is equal
P <- 2000 #number of permutations
```

```
est <- c(control.estimate, hormone.estimate)</pre>
id <- c(rep(0, length(control.estimate)), rep(1, length(hormone.estimate)))</pre>
df <- data.frame(est, id)</pre>
theta.h \leftarrow mean(df[df$id == 1, 1])
theta.c \leftarrow mean(df[df$id == 0, 1])
res.diff.mean <- rep(0, P)</pre>
for (p in 1:P) {
    perm <- sample(nrow(df))</pre>
    dat <- transform(df, id = id[perm])</pre>
    theta.h <- mean(dat[dat$id == 1, "est"])</pre>
    theta.c <- mean(dat[dat$id == 0, "est"])</pre>
    res.diff.mean[p] <- theta.h - theta.c</pre>
}
obs.diff.mean <- theta.h - theta.c
hist(res.diff.mean, breaks = 25, col = "gray", main = "Permutation Test for Difference of the Means")
abline(v = obs.diff.mean, col = "red")
```

Permutation Test for Difference of the Means



```
quantile(res.diff.mean, probs = c(0.025, 0.975))

## 2.5% 97.5%
## -3.800114 3.945974
obs.diff.mean
```

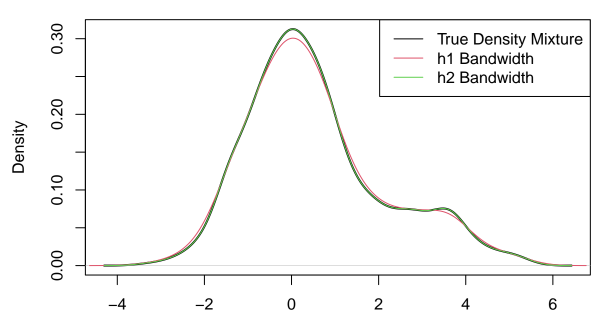
[1] 0.9722439

```
set.seed(1000)
n <- 1000  #sample size
comp <- sample(1:2, prob = c(0.8, 0.2), size = n, replace = TRUE)
mu = c(0, 3)
stan.dev = sqrt(c(1, 1))
samp <- rnorm(n, mean = mu[comp], sd = stan.dev[comp])

h1 <- 1.06 * (n^(-1/5)) * sd(samp)
h2 <- 0.9 * (n^(-1/5)) * min(sd(samp), (IQR(samp))/1.34)

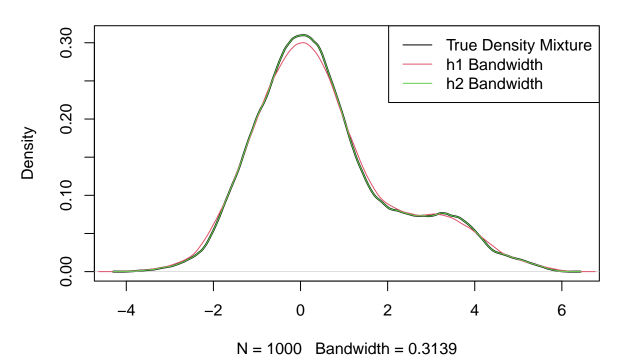
plot(density(samp), col = 1, main = "Gaussian Kernel Density",
    lwd = 2)  #true density estimate
lines(density(samp, bw = h1), col = 2)
lines(density(samp, bw = h2), col = 3)
legend("topright", c("True Density Mixture", "h1 Bandwidth",
    "h2 Bandwidth"), lty = 1, col = 1:3)</pre>
```

Gaussian Kernel Density

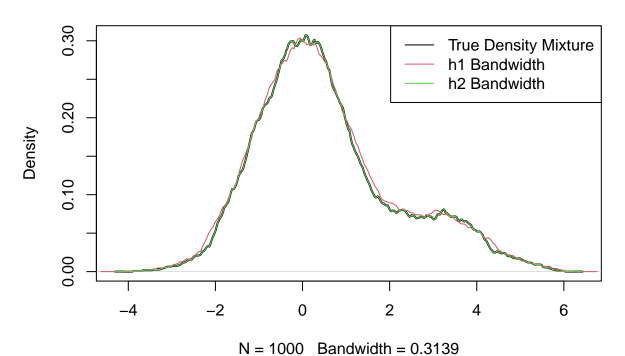


N = 1000 Bandwidth = 0.3139

Epanechnikov Kernel Density

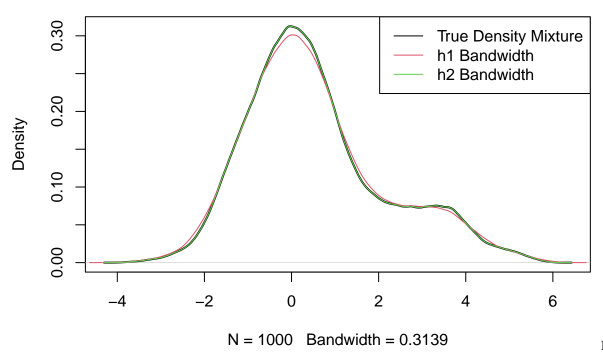


Rectangular Kernel Density



```
plot(density(samp, kernel = "triangular"), col = 1, main = "Triangular Kernel Density",
    lwd = 2) #true density estimate
lines(density(samp, bw = h1, kernel = "triangular"), col = 2)
lines(density(samp, bw = h2, kernel = "triangular"), col = 3)
legend("topright", c("True Density Mixture", "h1 Bandwidth",
    "h2 Bandwidth"), lty = 1, col = 1:3)
```

Triangular Kernel Density



on the plots from above: The h2 bandwidth is a better smoothing parameter compared to using the h1 bandwidth because the h2 bandwidth plot fits over the true density mixture plots. Moreover, this shows that while changing the kernel type will change the shape of the mixture plot, the h2 bandwidth still fits over the true density mixture better than the h1 bandwidth plot. The h1 bandwidth plot is hardly affected when changing the kernel.