Computational Statistics - STAT 575 - HW #4

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Problem 1

Use Metropolis Hasting algorithm to generate $Y \sim \text{GAM}(\alpha, 1)$, where $\alpha > 1$. Note α need not to be an integer. Consider the proposal distribution g, which is the density of GAM(a, b), where $a = |\alpha|$ and $b = a/\alpha$.

Part (a)

Implement your accept-reject algorithm and Metropolis-hastings algorithms to get a sample of 10000 from $Y \sim \text{GAM}(\alpha = 2.5, 1)$.

Acceptance-rejection sampler

We implement the density and sampler functions, respectively dgamma1 and rgamma1.

Suppose $X \sim \text{GAM}(\alpha, \beta)$ where α is the shape parameter and β is the rate parameter. Then, X has a density given by

$$f_X(x) = \frac{\beta^{\alpha}}{\Gamma(\alpha)} x^{\alpha - 1} e^{-\beta x}.$$

Suppose $Y \sim \text{GAM}(\alpha, 1)$, $\alpha \geq 1$. This is hard to sample from. However, $Z \sim \text{GAM}(a, b)$ where $a = \lfloor \alpha \rfloor$ and $b = a/\alpha$, is easy to sample from, since

$$Z = \sum_{i=1}^{a} X_i \sim \text{GAM}(a, b)$$

where $X_i \sim \text{EXP}(b)$.

We may thus use acceptance-rejection sampling to sample $z \sim f_Z$ and then $u \sim U(0,1)$ and accept z as a realization from f_Y if $u \leq \frac{f_Y(z)}{cf_Z(z)}$ where, optimally,

$$c = \max_{y} \left\{ \frac{f_Y(y)}{f_Z(y)} \right\},\,$$

but in general c must only satisfy $f_Y(y)/cf_Z(y) \leq 1$.

Note that we may use their respective kernels instead, since the ratio of the kernels is proportional to $f_Y(y)/f_Z(y)$. The ratio of the kernels is $h(y) = y^{\alpha-a}e^{(b-1)x}$. Thus, we seek $c = \max_y h(y)$ which may be obtained by solving for the input y^* that maximizes $\log h$, which is the stationary point

$$\left. \frac{d}{dy} \log h(y) \right|_{y=y^*} = 0,$$

and then substituting that into the ratio and simplifying, yielding the result

$$c = h(y^*) = (\alpha/e)^{\alpha - \lfloor \alpha \rfloor}.$$

Observe that if α is an integer, then c = 1 and h(y) = 1, in which case the target distribution and the candidate distribution are identical.

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We implement the acceptance-rejection algorithm with the following code:

```
# The density function for random variates in the family
# GAM(shape=alpha, rate=1)
dgamma1 <- function(x, alpha) {</pre>
    dgamma(x, shape = alpha, rate = 1)
}
# An acceptance-rejection sampling procedure for random variates in the family
# GAM(shape=alpha, rate=1)
accept.reject.rgamma1 <- function(n, alpha) {</pre>
    a <- floor(alpha)
    rate <- a/alpha
    q \leftarrow (alpha/exp(1))^(a - alpha)
    ys <- vector(length = n)
    for (i in 1:n) {
        repeat {
             y <- sum(rexp(a, rate)) # draw candidate
             if (runif(1) \le q * y^(alpha - a) * exp(y * a/alpha - y)) {
                 ys[i] <- y
                 break
             }
        }
    }
    ys
}
```

We sample from GAM(2.5, 1) with the the acceptance-rejection method with:

```
alpha <- 2.5
m <- 10000
accept.reject.samp <- accept.reject.rgamma1(m, alpha)</pre>
```

Metropolis-Hastings algorithm

Here is our implementation of the Metropolis-Hastings algorithm:

```
# A sampling procedure for random variates in the family
# GAM(shape=alpha,rate=1) using Metropolis-Hastings algorithm
metro.hast.rgamma1 <- function(n, alpha, burn = 0) {
    a <- floor(alpha)
    rate <- a/alpha

# density for random variates in the family GAM(shape=alpha,rate=1)
    f <- function(x) {
        dgamma(x, shape = alpha, rate = 1)
    }
    g <- function(x) {
        dgamma(x, shape = a, rate = rate)
    }

m <- n + burn
    ys <- vector(length = m)
    ys[1] <- sum(rexp(a, rate))</pre>
```

```
for (i in 2:m) {
    v <- sum(rexp(a, rate)) # draw from g
    u <- ys[i - 1]
    R <- f(v) * g(u)/(f(u) * g(v))
    if (runif(1) <= R) {
        ys[i] <- v
    } else {
        ys[i] <- u
    }
}
ys[(burn + 1):m]
}</pre>
```

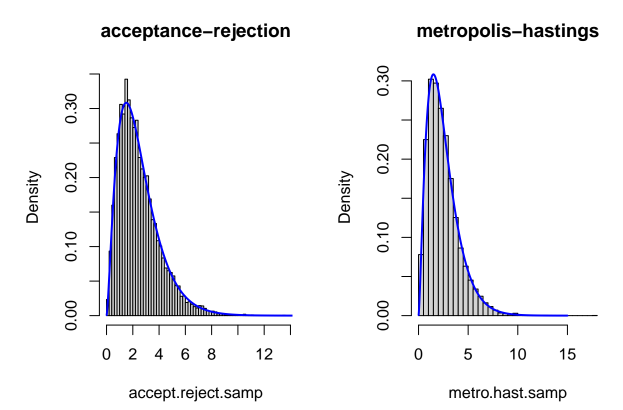
We sample from the Metro-Hastings algorithm with the following code:

```
metro.hast.samp <- metro.hast.rgamma1(m, alpha)
```

Part (b)

Check on mixing and convergence using plots. Run multiple chain and compute the Gelman-Rubin statistics. You may pick any reasonable burn-in.

We plot the histograms with:



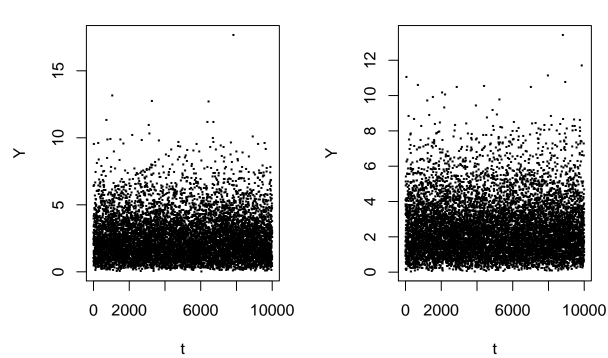
Both samplers seem to be compatible with the density.

We plot the sample path of the Metropolis-Hastings and acceptance-rejection samplers with:

```
par(mfrow = c(1, 2))
plot(metro.hast.samp, pch = ".", xlab = "t", ylab = "Y", main = "metropolis-hastings")
plot(accept.reject.samp, pch = ".", xlab = "t", ylab = "Y", main = "acceptance-rejection")
```

metropolis-hastings

acceptance-rejection



Both of these look good, as neither remain at or near the same value for many iterations. They also both quickly move away from their initial values.

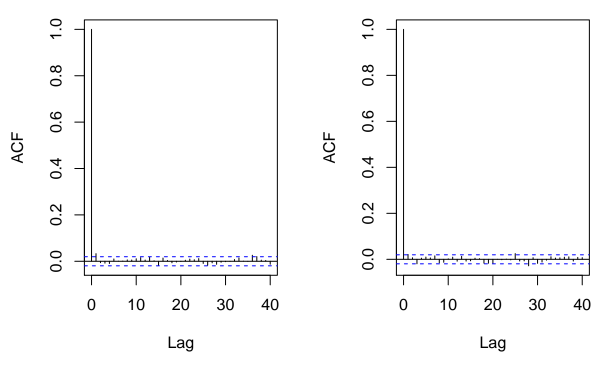
We plot the ACFs with:

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```
par(mfrow = c(1, 2))
acf(metro.hast.samp)
acf(accept.reject.samp)
```

Series metro.hast.samp

Series accept.reject.samp



Both samplers seem to have very little autocorrelation. If an uncorrelated sample is extremely important, to be safe, taking every other sample point would probably be sufficient.

We implement the Gelman-Rubin statistic with:

```
# samps: should be an L x J matrix, where L is the length of the samples and J
# is the number of samples (independent chains).
gelman.rubin <- function(samps) {
    L <- nrow(samps)
    J <- ncol(samps)
    x.bar <- apply(samps, 2, mean)
    B <- var(x.bar) * L
    W <- mean(apply(samps, 2, var))

    ((L - 1)/L * W + B/L)/W
}</pre>
```

Next, we compute Gelman-Rubin statistics on the computed independence chains.

```
chains <- 1000
samps <- matrix(nrow = m, ncol = chains)
for (i in 1:chains) {
    samps[, i] <- metro.hast.rgamma1(m, alpha, burn = 1000)
}
gelman.rubin.stat <- gelman.rubin(samps)
print(gelman.rubin.stat)</pre>
```

We see that the Gelman-Rubin statistic is given by

$$R = 1.0000067.$$

We adopt the rule of thumb that if $\sqrt{R} < 1.1$, the burn-in and chain length are sufficient. We compute \sqrt{R} to be

$$\sqrt{R} = 1.0000033,$$

and thus are satisfied with our burn-in choice and chain length.

Part (c)

Estimate $E(Y^2)$ using the generated chain. Compare with the estimate you get with acceptance-rejection sampling (Exam 1).

Theoretically,

$$E(Y^2) = \frac{\Gamma(2+\alpha)}{\Gamma(\alpha)} = \frac{\Gamma(4.5)}{\Gamma(2.5)} = 8.75.$$

We estimate $\mathrm{E}(Y^2)$ using the acceptance-rejection and Metropolis-Hastings by taking the square of each element in the samples they generated and then taking the mean:

```
tab <- matrix(nrow = 2, ncol = 1)
rownames(tab) <- c("acceptance-rejection", "metropolis-hastings")
colnames(tab) <- c("mean")
tab[1] <- c(mean(accept.reject.samp^2))
tab[2] <- c(mean(metro.hast.samp^2))</pre>
knitr::kable(data.frame(tab))
```

	mean
acceptance-rejection	8.436799
metropolis-hastings	8.749739

Both are quite close to the true value of 8.75.

Problem 2 (Problem 7.1)

Rework the textbook example. Consider the mixture normal $\delta N(7,0.5^2) + (1-\delta)N(10,0.5^2)$.

Part (a)

Simulate 200 realizations from the mixture distribution with $\delta = 0.7$. Draw a histogram of these data.

We implement the density and sampler for the mixture distribution with:

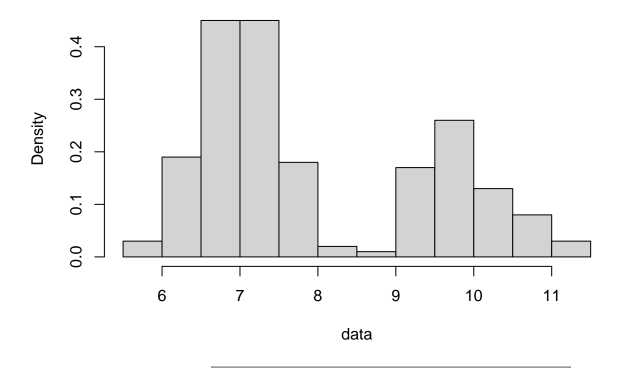
```
dmix <- function(x, delta) {
    delta * dnorm(x, 7, 0.5) + (1 - delta) * dnorm(x, 10, 0.5)
}</pre>
```

```
rmix <- function(n, delta) {
    xs <- vector(length = n)
    for (i in 1:n) {
        xs[i] <- ifelse(runif(1) < delta, rnorm(1, 7, 0.5), rnorm(1, 10, 0.5))
    }
    xs
}</pre>
```

We generate a sample and plot its histogram with:

```
n <- 200
delta <- 0.7
data <- rmix(n, delta)
hist(data, freq = F)</pre>
```

Histogram of data



Part (b)

Now assume δ is unknown. Implement independence chain MCMC procedure to simulate from the posterior distribution of δ , using your data from part (a).

```
lmix <- Vectorize(function(delta, xs) {
    if (delta < 0 || delta > 1) {
        return(0)
    }
    p <- 1
    for (x in xs) {
        p <- p * dmix(x, delta)
    }
    p</pre>
```

```
}, "delta")

logmix <- Vectorize(function(delta, xs) {
    if (delta < 0 || delta > 1) {
        return(-Inf)
    }
    logp <- 0
    for (x in xs) {
        logp <- logp + log(dmix(x, delta))
    }
    logp
}, "delta")</pre>
```

A sample $\{x_t\}$ drawn from the mixture normal with density dmix is observed with likelihood $\text{lmix}(\delta|\vec{x})$ with respect to δ with prior distribution $p(\delta)$. Thus, the posterior distribution is given by

$$p(\delta|\vec{x}) \propto p(\delta) \operatorname{lmix}(\delta|\vec{x}).$$

In the independence chain MCMC, we may use the prior as the proposal density, $f(\delta) = p(\delta|\vec{x})$ and g = p, and thus

$$R = \frac{f(\delta^*)g(\delta^{(t)})}{f(\delta^{(t)})g(\delta^*)} = \frac{p(\delta^*|\vec{x})p(\delta^{(t)})}{p(\delta^{(t)}|\vec{x})p(\delta^*)}$$

which may be rewritten as

$$R = \frac{p(\delta^*) \operatorname{lmix}(\delta^* | \vec{x}) p(\delta^{(t)})}{p(\delta^{(t)}) \operatorname{lmix}(\delta^{(t)} | \vec{x}) p(\delta^*)} = \frac{\operatorname{lmix}(\delta^* | \vec{x})}{\operatorname{lmix}(\delta^{(t)} | \vec{x})}.$$

Numerical imprecision

Suppose we have a data type T that models real numbers. Since computers are physical, T can only represent a finite set of numbers.

In the likelihood function,

$$lmix: \mathbb{R} \times 2^{\mathbb{R}} \mapsto \mathbb{R},$$

if we model \mathbb{R} with T, i.e.,

lmix:
$$T \times 2^T \mapsto T$$
,

then if the true value of the likelihood function applied to a sufficiently large sample is some value $p \in (0, \epsilon)$ where ϵ is the smallest representable positive number of type T, the best we can do is round p to 0 or ϵ . As a consequence, the likelihood function evaluates to 0 on any sufficiently large sample size.

Suppose $\epsilon = 2^{-K}$. If we use the log-likelihood instead,

$$\text{logmix} \colon T \times 2^T \mapsto T$$

then, for instance, $\log_2 \epsilon = -K$ where -K is very likely to be at least approximately representatable by T, and much smaller values as well. We cannot map many of these log-likelihoods back to a likelihood, but as long as we only need to work with log-likelihoods, this is not a problem.

With the above in mind, we replace the likelihood function with the log-likelihood function to significantly increase the space of samples we can work with.

```
delta.estimator.ic <- function(n, data, delta0 = runif(1), burn = 0) {
   m <- n + burn
   deltas <- vector(length = m)
   deltas[1] <- delta0</pre>
```

```
for (i in 2:m) {
    delta <- runif(1) # draw candidate from prior
    delta.old <- deltas[i - 1]
    log.R <- logmix(delta, data) - logmix(delta.old, data)
    if (log(runif(1)) <= log.R) {
        deltas[i] <- delta
    } else {
        deltas[i] <- delta.old
    }
}
deltas[(burn + 1):m]</pre>
```

Part (c)

Implement a random walk chain with $\delta^* = \delta^{(t)} + \epsilon_t$ with $\epsilon \sim \text{UNIF}(-1, 1)$.

We observe that $\epsilon_t \sim \text{UNIF}(-1,1)$ for $t=0,\ldots,t$ are the only random components. Thus, the conditional distribution of ϵ_{t+1} given ϵ_t is

 $\epsilon_{t+1} \sim f(\delta^* - \delta^{(t)})$

where f is the density of UNIF(-1,1).

```
delta.estimator.rw <- function(n, data, delta0 = runif(1), burn = 0) {</pre>
    m <- n + burn
    deltas <- vector(length = m)</pre>
    deltas[1] <- delta0</pre>
    for (i in 2:m) {
         delta.old <- deltas[i - 1]
         delta <- delta.old + runif(1, -1, 1)
         log.R <- logmix(delta, data) - logmix(delta.old, data)</pre>
         if (log(runif(1)) <= log.R) {</pre>
             deltas[i] <- delta
         } else {
             deltas[i] <- delta.old</pre>
         }
    }
    deltas[(burn + 1):m]
}
```

Part (d)

Reparameterize the problem letting $U = \log(\delta/(1-\delta))$ and $U^* = u(t) + \epsilon_t$. Implement a random walk chain with U as in Equation (7.8) page 208.

```
logit <- function(delta) {
    log(delta/(1 - delta))
}
logit.inv <- function(u) {</pre>
```

```
\exp(u)/(1 + \exp(u))
}
logit.inv.J <- function(u) {</pre>
    \exp(u)/(1 + \exp(u))^2
}
delta.estimator.u.rw <- function(n, data, delta0 = runif(1), burn = 0) {</pre>
    m <- n + burn
    u <- vector(length = m)</pre>
    u[1] <- logit(delta0)</pre>
    for (i in 2:m) {
         u.old <- u[i - 1]
         u.star <- u.old + runif(1, -1, 1)
         R <- lmix(logit.inv(u.star), data) * logit.inv.J(u.star)/(lmix(logit.inv(u.old),</pre>
             data) * logit.inv.J(u.old))
         if (runif(1) <= R) {</pre>
             u[i] <- u.star
         } else {
             u[i] \leftarrow u.old
         }
    }
    logit.inv(u[(burn + 1):m])
}
```

Part (e)

Compare the estimates and convergence behavior of three algorithms.

We do not do a burn-in, since we are interested in seeing how quickly the three methods converge. We only plot chains of length 1000.

We generate the data sets with:

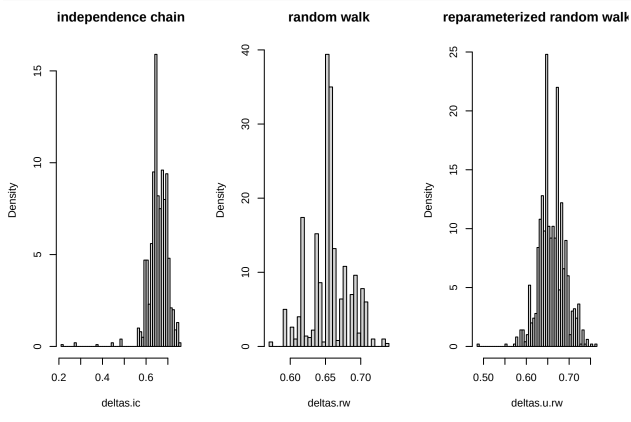
```
chain <- 1000
burn <- 0
deltas.ic <- delta.estimator.ic(chain, data, burn = burn)
deltas.rw <- delta.estimator.rw(chain, data, burn = burn)
deltas.u.rw <- delta.estimator.u.rw(chain, data, burn = burn)
tab <- matrix(nrow = 3, ncol = 1)
rownames(tab) <- c("independence chain", "random walk", "reparameterized random walk")
colnames(tab) <- c("mu")
tab[1, ] <- mean(deltas.ic)
tab[2, ] <- mean(deltas.rw)
tab[3, ] <- mean(deltas.u.rw)
knitr::kable(data.frame(tab))</pre>
```

	mu
independence chain	0.6554597
random walk	0.6557555
reparameterized random walk	0.6596772

As the table of estimations shows, all three methods provide a good estimate of δ . Next, we consider their convergence and mixing behavior.

We plot the histograms with:

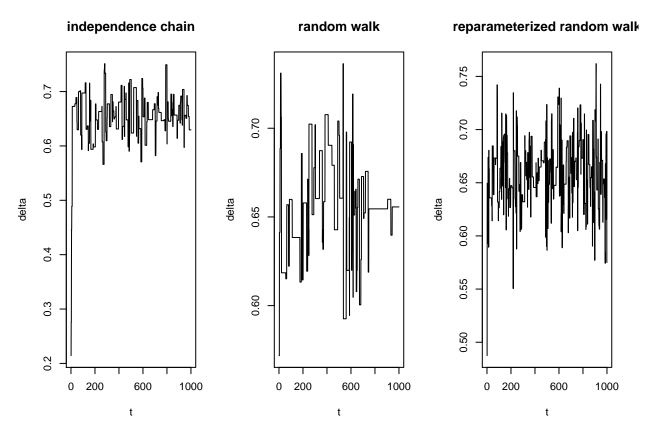
```
par(mfrow = c(1, 3))
hist(deltas.ic, freq = F, breaks = 50, main = "independence chain")
hist(deltas.rw, freq = F, breaks = 50, main = "random walk")
hist(deltas.u.rw, freq = F, breaks = 50, main = "reparameterized random walk")
```



The reparameterized random walk metropolis has a histogram that is most compatible with normality, i.e., characteristic bell curve with a mode at $\delta = 0.7$. That said, all three histograms arguably satisfy normality with approximately the same mean at $\delta = 0.7$.

We plot the sample paths with:

```
par(mfrow = c(1, 3))
plot(deltas.ic, pch = ".", type = "l", xlab = "t", ylab = "delta", main = "independence chain")
plot(deltas.rw, pch = ".", type = "l", xlab = "t", ylab = "delta", main = "random walk")
plot(deltas.u.rw, pch = ".", type = "l", xlab = "t", ylab = "delta", main = "reparameterized random walk")
```



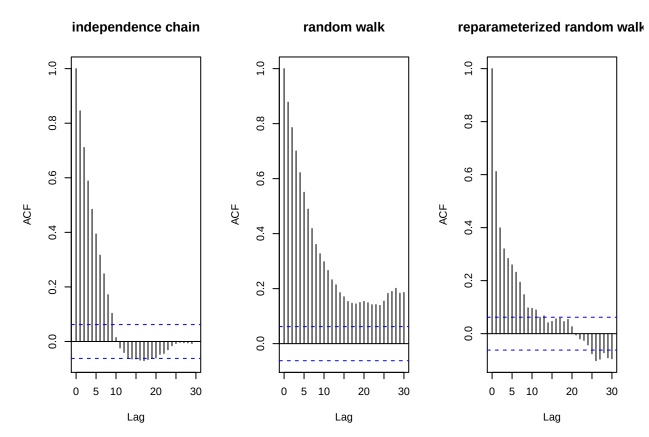
We see that the random walk demonstrates relatively poor mixing. It has a high rejection rate (stays at the same level for long periods of time), causing it to explore the support of the likelihood slowly.

The sample path of the indepedence chain also can be said to demonstrate poor mixing.

The reparameterized random walk exihibits good mixing, vigorously jiggling around the true value.

We plot the ACFs with:

```
par(mfrow = c(1, 3))
acf(deltas.ic, main = "independence chain")
acf(deltas.rw, main = "random walk")
acf(deltas.u.rw, main = "reparameterized random walk")
```



They all decay quickly, but in order of increasing autocorrelation: the reparameterized random walk, the independence chain, and the random walk.

In particular, the reparameterized random walk shows autocorrelation that decays quite rapidly with respect to lag time.

Problem 3

Consider an i.i.d. sample X_1, \dots, X_n from $N(\mu, \sigma^2)$. Consider the Bayesian analysis to estimate μ and $\tau = (\sigma^2)^{-1}$. We put prior $\mu \sim N(m, p^{-1})$ and $\tau \sim \text{GAM}(a, b)$.

Part (a)

Write out the posterior distribution of $(\mu, \tau) | \vec{x}$. You may ignore the normalizing constant.

Note that the posterior distribution is given by

$$\pi(\mu, \tau | \vec{x}) = f(\vec{x} | \tau, \mu) f(\tau) f(\mu) / Z$$

where $Z = f(\vec{x})$ is the normalizing constant. We may rewrite this as

$$\pi(\mu, \tau | \vec{x}) \propto L(\tau, \mu | \vec{x}) f(\tau) f(\mu).$$

The likelihood function of (μ, τ) conditioned on the data \vec{x} is given by the product of the normal density on the sample,

$$L(\mu,\tau|\vec{x}) = \prod_{i=1}^{n} \frac{1}{\sqrt{2\pi\sigma^{2}}} \exp\left\{-\frac{1}{2\sigma^{2}}(x_{i}-\mu)^{2}\right\}$$

which may be simplified to

$$L(\mu,\tau|\vec{x}) = \left(2\pi\sigma^2\right)^{-n/2} \exp\left\{-\frac{1}{2\sigma^2}\sum_{i=1}^n (x_i-\mu)^2\right\}.$$

Observing $\tau = 1/\sigma^2$, we obtain the result

$$L(\mu,\tau|\vec{x}) \propto \tau^{n/2} \exp\left\{-\frac{\tau}{2} \sum_{i=1}^n (x_i - \mu)^2\right\}.$$

The prior for μ is $f(\mu) \propto \exp\left(-\frac{p}{2}(\mu - m)^2\right)$ and the prior for τ is $f(\tau) \propto \tau^{a-1} \exp(-b\tau)$. Putting it all together, we obtain the result

$$\pi(\mu,\tau|\vec{x}) \propto \tau^{n/2+a-1} \exp\left\{-\frac{\tau}{2} \sum_{i=1}^n (x_i - \mu)^2 - \frac{p}{2} (\mu - m)^2 - b\tau\right\}.$$

Part (b)

Show the posterior conditional distribution of $\mu|(\tau, \vec{x})$ is

$$N\left(\frac{n\tau\bar{x}+pm}{n\tau+p},\frac{1}{n\tau+p}\right)$$

and the posterior conditional distribution of $\tau|(\mu, \vec{x})$ is

$${\rm GAM}(a+n/2,b+n/2[s^2+(u-\bar{x})^2])$$
 .

Distribution of $\mu|(\tau, \vec{x})$

The conditional distribution of $\mu|(\tau, \vec{x})$ is given by

$$\pi(\mu|\tau, \vec{x}) = \frac{\pi(\mu, \tau|\vec{x})}{\pi(\tau|\vec{x})}$$

where

$$\pi(\tau|\vec{x}) = \int_{-\infty}^{\infty} \pi(\mu, \tau|\vec{x}) d\mu.$$

Note that $\pi(\tau|\vec{x})$ is not a function of μ , and thus

$$\pi(\mu|\tau,\vec{x}) \propto \pi(\mu,\tau|\vec{x}).$$

We discard any factor that is not a function of μ ,

$$\pi(\mu|\tau,\vec{x}) \propto \exp\left\{-\frac{\tau}{2}\sum_{i=1}^n(x_i-\mu)^2\right\} \exp\left(-\frac{p}{2}(\mu-m)^2\right).$$

Now, we wish to show that this is the kernel of the given normal distribution. We do this by discarding any factors that are not a function of μ and rewriting the result that fits the pattern of a normal kernel

$$\pi(\mu|\tau,\vec{x}) = \exp\left\{-\frac{1}{2k_1}\left(\mu - k_2\right)^2\right\}$$

with a mean k_2 and variance k_1 .

$$\pi(\mu|\tau,\vec{x}) \propto \exp\left\{-\frac{\tau}{2}\left[\sum x_i^2 - 2\mu\sum x_i + n\mu^2\right] - \frac{p}{2}\left[\mu^2 - 2\mu m + m^2\right]\right\} \tag{1}$$

$$\propto \exp\left\{-\frac{1}{2}\left[-2n\tau\bar{x}\mu + n\tau\mu^2 + p\mu^2 - 2pm\mu\right]\right\}$$
 (2)

$$\propto \exp\left\{-\frac{1}{2}\left[(n\tau+p)\mu^2 - (2pm+2n\tau\bar{x})\mu\right]\right\}$$
 (3)

$$\propto \exp\left\{-\frac{n\tau+p}{2}\left[\mu^2 - \frac{2(pm+n\tau\bar{x})}{n\tau+p}\mu\right]\right\}. \tag{4}$$

Completing the square, we obtain

$$\pi(\mu|\tau,\vec{x}) \propto \exp\left\{-\frac{n\tau+p}{2}\left[\mu - \frac{pm+n\tau\bar{x}}{n\tau+p}\right]^2 - \left[\frac{pm+n\tau\bar{x}}{n\tau+p}\right]^2\right\}$$
 (5)

$$\propto \exp\left\{-\frac{n\tau + p}{2} \left[\mu - \frac{pm + n\tau\bar{x}}{n\tau + p}\right]^2\right\}. \tag{6}$$

Thus, we see that this is the kernel of a normal density with mean

$$\frac{pm + n\tau \bar{x}}{n\tau + p}$$

and variance

$$\frac{1}{n\tau+p}.$$

Distribution of $\tau | (\mu, \vec{x})$

The conditional distribution of $\tau | (\mu, \vec{x})$ is given by

$$\pi(\tau|\mu, \vec{x}) = \frac{\pi(\mu, \tau|\vec{x})}{\pi(\mu|\vec{x})}$$

where

$$\pi(\mu|\vec{x}) = \int_{-\infty}^{\infty} \pi(\mu, \tau|\vec{x}) d\tau.$$

Note that $\pi(\mu|\vec{x})$ is not a function of τ , and thus

$$\pi(\tau|\mu, \vec{x}) \propto \pi(\mu, \tau|\vec{x}).$$

We discard any factor that is not a function of τ ,

$$\pi(\tau|\mu,\vec{x}) \propto \tau^{n/2+a-1} \exp\left\{-\frac{\tau}{2} \sum_{i=1}^n (x_i - \mu)^2 - b\tau\right\}.$$

We seek to match it to the kernel of the gamma density for $GAM(\alpha, \beta)$, which is given by

$$\tau^{\alpha-1} \exp(-\beta \tau)$$
.

So, we rewrite the above as

$$\pi(\tau|\mu,\vec{x}) \propto \tau^{n/2+a-1} \exp\left\{-\frac{\tau}{2} \sum (x_i - \mu)^2 - b\tau\right\} \tag{7}$$

$$\propto \tau^{n/2+a-1} \exp\left\{-\left(\frac{1}{2}\sum_{i}(x_i-\mu)^2+b\right)\tau\right\}. \tag{8}$$

Thus, we see that $\alpha = n/2 + a$ and $\beta = b + \frac{1}{2} \sum (x_i - \mu)^2$. The β is not in the form requested, so we continue, focusing strictly on β .

We may rewrite $\sum (x_i - \mu)^2$ as

$$\sum (x_i - \mu)^2 = \sum (x_i - \bar{x} + \bar{x} - \mu)^2 \tag{9}$$

$$= \overline{\sum} (x_i - \bar{x})^2 + \sum (\bar{x} - \mu)^2 - 2 \sum (x_i - \bar{x})(\bar{x} - \mu) \tag{10}$$

$$= \sum_{i=1}^{n} (x_i - \bar{x})^2 + n(\bar{x} - \mu)^2 - 2\left(\sum_{i=1}^{n} x_i - n\bar{x}\right)(\bar{x} - \mu). \tag{11}$$

Since $\sum x_i - n\bar{x} = 0$, we may drop the last term,

$$\sum (x_i - \mu)^2 = \sum (x_i - \bar{x})^2 + n(\bar{x} - \mu)^2.$$

We note that $s^2 = \frac{1}{n} \sum_i (x_i - \bar{x})^2$ and thus

$$\sum (x_i - \mu)^2 = n(s^2 + (\mu - \bar{x})^2),$$

which shows that $\beta = b + \frac{n}{2}(s^2 + (\mu - \bar{x})^2)$ and $\alpha = n/2 + a$ and is thus the kernel of

$${\rm GAM}\big(n/2 + a, b + n(s^2 + (\mu - \bar{x})^2)/2\big)\,.$$

Part (c)

First, generate some "observed" sample data using x = rnorm(200, mu = 5, sigma = 2). Hand-code Gibbs Sampler algorithm to sample (μ, τ) from the posterior using x. You make take prior parameters a = 0.0001; b = 0.0001; p = 0.0001; m = 0. Use the estimated posterior mean and compare your estimates with the true parameters $\mu = 5$ and $\tau = 0.25$.

We generate the sample with:

```
x \leftarrow rnorm(200, mean = 5, sd = 2)
```

We implement the Gibbs sampling with the function:

```
mu.tau.gibbs <- function(n, x, burn = 1000, theta0 = NULL, p = 1e-04, m = 0, a = 1e-04,
    b = 1e-04) {
    x.mu <- mean(x)
    x.s2 <- var(x)
    x.n <- length(x)

rmu <- function(tau) {
    mean <- (x.n * tau * x.mu + p * m)/(x.n * tau + p)
    var <- 1/(x.n * tau + p)
    rnorm(1, mean = mean, sd = sqrt(var))
}

rtau <- function(mu) {
    rgamma(1, shape = a + x.n/2, rate = b + x.n * (x.s2 + (mu - x.mu)^2)/2)
}

prior <- function() {
    c(rnorm(1, m, 1/p), rgamma(1, a, b))</pre>
```

```
}
    N \leftarrow n + burn
    thetas <- matrix(nrow = N, ncol = 2)
    if (is.null(theta0)) {
         thetas[1, ] <- prior()</pre>
    } else {
         thetas[1, ] <- theta0
    }
    for (i in 1:(N - 1)) {
         tau.new <- rtau(thetas[i, 1])</pre>
         mu.new <- rmu(tau.new)</pre>
         thetas[i + 1, ] <- c(mu.new, tau.new)
    }
    thetas <- thetas [(burn + 1):N, ]
    mu.est <- mean(thetas[, 1])</pre>
    tau.est <- mean(thetas[, 2])</pre>
    sigma.est <- sqrt(1/tau.est)</pre>
    list(theta.dist = thetas, mu.est = mu.est, tau.est = tau.est, sigma.est = sigma.est)
}
```

We use the Gibbs sampler to estimate (μ, τ) with:

```
# set up hyper-parameters
a <- 1e-04
b <- 1e-04
m <- 0
p <- 1e-04

res <- mu.tau.gibbs(1e+05, x, burn = 10000, p = p, a = a, m = m, b = b)
mu.est <- round(as.numeric(res$mu.est), digits = 4)
tau.est <- round(as.numeric(res$tau.est), digits = 4)
var.est <- round(1/tau.est, digits = 4)
c(mu.est, tau.est)</pre>
```

[1] 5.0265 0.2953

We estimate (μ, τ) to be (5.0265, 0.2953)'. Therefore, we estimate that the data is being sampled from the normal distribution

$$X_i \sim N(\mu = 5.0265, \sigma^2 = 3.3864). \tag{12}$$

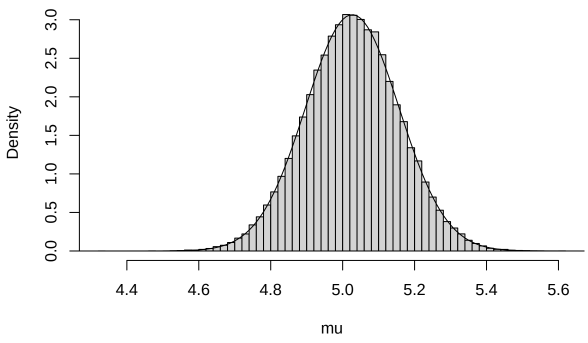
Additional analysis

Out of curiosity, we decided to plot the marginals of the sample superimposed with their respective conditional densities with:

```
x.mu <- mean(x)
x.n <- length(x)
x.s2 <- var(x)
mu.dist <- res$theta.dist[, 1]
tau.dist <- res$theta.dist[, 2]
hist(mu.dist, freq = F, breaks = 50, main = "marginal of mu", xlab = "mu")</pre>
```

```
lines(seq(4, 6, by = 0.01), dnorm(seq(4, 6, by = 0.01), mean = (x.n * tau.est * x.mu)/(x.n * tau.est + p), sd = sqrt(1/(x.n * tau.est + p)))
```

marginal of mu



```
hist(tau.dist, freq = F, breaks = 50, main = "marginal of tau", xlab = "tau")
lines(seq(0.15, 0.4, by = 0.01), dgamma(seq(0.15, 0.4, by = 0.01), shape = a + x.n/2,
rate = b + x.n * (x.s2 + (mu.est - x.mu)^2)/2))
```

marginal of tau

