```
options(scipen=999) #for removing scientific notation on the plots
n = 50
s = 13
#alpha and beta in prior Beta distribution
alpha0 = 5
beta0 = 5
#alpha and beta in posterior Beta distribution
alpha = alpha0 + s
beta = beta0 + (n - s)
#true mean and standard deviation of the posterior Beta distribution
(formulas from wikipedia for Beta distribution)
true mean = alpha / (alpha + beta)
true sd = sqrt(alpha * beta / ((alpha + beta)^2 * (alpha+beta+1)))
draws = seg(5000, 500000, 50000) #different number of draws, i.e. sample
sizes to use
means = c(1:length(draws)) #vector to store the means of the different
sample sizes
sds = c(1:length(draws)) #vector to store the sd's of the different sample
sizes
#for each of the different number of draws, compute and store the estimated
mean and sd.
#storing the absolute value for easier comparison.
for (i in 1:length(draws)) {
 set.seed(12345)
 data = rbeta(draws[i], alpha, beta)
 mean = mean(data)
 sd = sd(data)
 means[i] = abs(true mean - mean)
 sds[i] = abs(true sd - sd)
}
#As we see in the plot, the difference between the true mean/sd and the
estimated mean/sd from the simulations
#go to 0 as we increase the number of draws, i.e. the posterior mean and sd
converges to the true values of 0.3 and 0.05867387.
max y = max (means, sds)
plot(draws, means, type="1", lwd = 2, col="blue", ylim=c(0, max y), main =
"Deviance from true mean and sd for different number of draws",
    xlab="Number of draws", ylab="Deviance between true mean/sd and
simulated mean/sd" )
points(draws, sds, type="l", lwd = 2, col="green")
legend(x = max(draws)*0.8, y = max_y*0.95, legend = c("mean", "sd"), col =
c("blue", "green"), lwd = 3)
#pbeta gives the distribution function for the Beta distribution,
#i.e. the accumulated probability mass up to some value (area under curve
less or equal to the value)
true prob = pbeta(0.3, alpha, beta)
true prob
nDraws = 10000
```

```
set.seed(12345)
data = rbeta(nDraws, alpha, beta)
#using our 10000 draws from the posterior,
#we can calculate the probability by taking number of samples < 0.3 divided
by total number of samples
sim prob = length(data[data<0.3]) / nDraws</pre>
sim prob
#The probability from simulation with 10000 draws, 0.5156, is very close to
the exact probability of 0.5150.
#Plot of density estimates of our data, a value of around 0.515 looks
reasonable.
plot(density(data))
abline (v=0.3, col="red")
nDraws = 10000
set.seed(12345)
data = rbeta(nDraws, alpha, beta)
#posterior distribution
hist(data, breaks=25)
plot(density(data), main = "Density estimation of posterior distribution")
\#simulating draws from the posterior distribution of the log-odds
\#(\text{logit maps the probability values from }(0,1) \text{ to }(-\text{inf, inf}))
log odds = log(data / (1-data))
#posterior distribution of the log-odds
hist(log odds, breaks=25)
plot(density(log odds), main = "Density estimation of posterior
distribution of log odds")
```

```
data = c(33, 24, 48, 32, 55, 74, 23, 76, 17)
n <- length(data)</pre>
mju = 3.5
tao2 = sum((log(data)-mju)^2)/n
NDraws = 10000
PostDraws = c(1:NDraws)
set.seed(123465)
#drawing 10000 sigma^2 from the inv-chi^2 distribution
PostDraws = ((n) *tao2)/rchisq(NDraws, n)
plot(density(PostDraws), xlim=c(0,2), main = expression(paste('Posterior
distribution of ', sigma^2,)))
#phi(z) is the cumulative distribution function (CDF) for the standard
normal distribution with mean 0 and variance 1.
#pnorm gives the distribution function for the Normal distribution with
default mean=1 and sd=1 (the CDF),
#i.e. the accumulated probability mass up to some value (area under curve
less or equal to the value)
G = 2*pnorm(sqrt(PostDraws/2)) - 1
plot(density(G), main = "Posterior distribution of the Gini coefficient G
for the current data set ")
######################################
#estimation of the the 95% equal tail credible interval using mean and sd
mean = mean(G)
sd = sd(G)
lowerBound = mean - 1.96*sd
upperBound = mean + 1.96*sd
lowerBound
upperBound
#more exact values using gnorm
lowerBound = qnorm(0.025, mean, sd)
upperBound = qnorm(0.975, mean, sd)
lowerBound
upperBound
plot(density(G), main = "Posterior distribution of the Gini coefficient G
for the current data set, \n with 95% equal tail credible interval. ")
abline(v=lowerBound, col="red")
abline(v=upperBound, col="red")
#extract x- and y-values from the density estimate and sort these on y-
value
\#density = density(G, n=10000)
density = density(G)
```

```
x = density$x
y = density$y
xy = cbind(x, y)
xy sorted = xy[order(xy[,2],decreasing=TRUE),]
#calculate what value 95% of the accumulated y-values corresponds to, i.e.
the accumulated value to stop at
#when deciding which (x,y)-pairs to look at
stop value = sum(xy sorted[,2])*0.95
#accumulated sum of the sorted y-values
#find out at which index we reach 95% mass (i.e. the stop value),
#then index 1 to stop index corresponds to 95% mass
acumulatedMass <- function(stop val) {</pre>
  for (i in 1:length(xy_sorted)) {
    if (sum(xy sorted[1:i,2]) > stop val) {
     return (i-1)
    }
  1
stop index = acumulatedMass(stop value)
#alternative to using above function:
#cumSum = cumsum(xy sorted[,2])
#stop index = min(which(cumSum > stop value)) - 1
#only keep the 95% of the mass corresponding to highest densities
xy sorted 95 = xy sorted[1:stop index,]
#find the indexes corresponding to lower and upper bound
lower index = which.min(xy sorted 95[,1])
upper index = which.max(xy sorted 95[,1])
lower index
upper index
#extract actual lower and upper bounds
HPDIlower = xy sorted 95[lower index,1]
HPDIupper = xy_sorted_95[upper_index,1]
HPDIlower
HPDIupper
plot(density(G), main = "Posterior distribution of the Gini coefficient G
for the current data set, \n with 95% Equal Tail Credible Interval, \n and
95% Highest Posterior Density Interval HPDI ")
legend(x = 0.6, y = 5, legend = c("Equal Tail Credible Interval", "HPDI"),
col = c("red", "blue"), lwd = 3)
abline(v=lowerBound, col="red")
abline(v=upperBound, col="red")
abline(v=HPDIlower, col="blue")
abline(v=HPDIupper, col="blue")
```

```
##################################
degrees = c(285, 296, 314, 20, 299, 296, 40, 303, 326, 308)
radians = c(1.83, 2.02, 2.33, -2.79, 2.07, 2.02, -2.44, 2.14, 2.54, 2.23)
mju = 2.51
lambda = 1
besselDegree = 0
#posterior proportional to likelihood * prior
#likelihood proportional to exp(k*cos(y-mju)) / IO(k)
#K~Exp(lambda=1) which gives the prior: lambda*exp(-lambda*k)
posterior = function(k, y, mju, degree, lambda) {
 #posterior proportional to:
 posterior = prod( exp(k*cos(y-mju))/besselI(k, degree) ) * lambda*exp(-
lambda*k)
 \#posterior = prod( exp(k*cos(y-mju))/besselI(k, degree) ) * dexp(k,
rate=lambda)
#grid of k-values to plot over
gridstep = 0.005
k = seq(0, 7, gridstep)
#calculate and store the value for each of the k-values
post = c(1:length(k))
for(i in 1:length(k)) {
 post[i] = posterior(k[i], radians, mju, besselDegree, lambda)
#normalizing the posterior distirbution of k so that it integrates to 1
post = post/(sum(post)*gridstep)
plot(k, post, type="1", main="Posterior distribution of k", col="blue")
#the approximate posterior mode of k is the largest value in the posterior
approx post K = k[which.max(post)]
approx post K
plot(k, post, type="l", main="Posterior distribution of k", col="blue")
abline(v=approx post K, col="red")
```