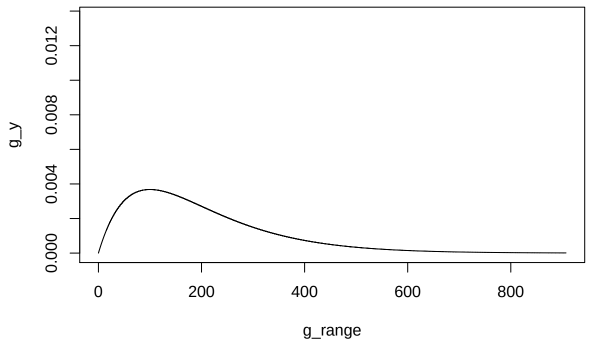
Version 2: 2 clusters; Fix cluster label switching u,v; unif; fix w [0,1]; only force ordering of means in 1 sample; change V prior; 5 chains

Simulate data

```
I <- 50
K <- 2
S <- 10

# choose diffuse priors for gamma
a_gamma <- 2
b_gamma <- 100

avrg <- a_gamma * b_gamma
std.dv <- sqrt(a_gamma*b_gamma^2)
g_range = seq(0, avrg + 5*std.dv, 0.01)
g_y = dgamma(g_range, a_gamma, rate = 1/b_gamma)
plot(g_range, g_y, type = "l", ylim=c(0, max(g_y) + 0.01))</pre>
```



```
set.seed(123)

a <- matrix(NA, nrow=K, ncol=S)
b <- matrix(NA, nrow=K, ncol=S)
for (s in 1:S) {
   a[, s] <- rgamma(K, a_gamma, rate = 1/b_gamma)
   b[, s] <- rgamma(K, a_gamma, rate = 1/b_gamma)
}</pre>
```

```
pi <- as.vector(rdirichlet(1, rep(1, K)))</pre>
z <- sample(1:K, size = I, replace = T, prob = pi)
w <- matrix(NA, nrow=I, ncol=S)
for (s in 1:S) {
  w[, s] <- rbeta(I, a[,s][z], b[,s][z])
tcn <- matrix(2, nrow=I, ncol=S)</pre>
m <- matrix(rep(sample(1:2, size = I, replace = T), S), nrow=I, ncol=S)
calcTheta <- function(m, tcn, w) {</pre>
  (m * w) / (tcn * w + 2*(1-w))
theta <- calcTheta(m, tcn, w)
n <- replicate(S, rpois(I, 100))</pre>
y <- matrix(NA, nrow=I, ncol=S)
for (i in 1:I) {
  for (s in 1:S) {
    y[i, s] <- rbinom(1, n[i, s], theta[i,s])
}
```

JAGS

```
jags.file <- file.path(models.dir, "v2_uv_unif_fix3.jags")</pre>
test.data <- list("I" = I, "S" = S, "K" = K,
                   "y" = y, "n" = n,
                   "m" = m, "tcn" = tcn)
jags.m <- jags.model(jags.file, test.data,</pre>
                       n.chains = numChains)
## Compiling model graph
##
      Resolving undeclared variables
##
      Allocating nodes
## Graph information:
##
      Observed stochastic nodes: 500
##
      Unobserved stochastic nodes: 591
##
      Total graph size: 8400
##
## Initializing model
params <- c("z", "w", "a", "b", "U", "V")
update(jags.m, n.iter=1000)
samps <- coda.samples(jags.m, params, n.iter=1000)</pre>
s <- summary(samps)</pre>
#effectiveSize(window(samps, start=5001))
pdf(file.path(trace.dir, paste0(runName, "_trace.pdf")))
plot(samps)
dev.off()
```

```
## pdf
##
mcmc_vals <- s$statistics</pre>
mcmc_w <- mcmc_vals[substr(rownames(mcmc_vals), 1, 1) == "w", "Mean"]</pre>
plot(as.vector(w), mcmc_w, type = "p")
      0.8
mcmc_w
      9.0
      0.4
      0.2
                0.2
                                   0.4
                                                      0.6
                                                                         8.0
                                                                                            1.0
                                              as.vector(w)
mcmc_z <- as.vector(mcmc_vals[substr(rownames(mcmc_vals), 1, 1) == "z", "Mean"])</pre>
\#mcmc_z \leftarrow round(mcmc_z, 0)
plot(z, mcmc_z, type = "p")
      1.8
      1.6
mcmc_z
      4.
      1.2
      1.0
                                                                                           0
             1.0
                            1.2
                                            1.4
                                                           1.6
                                                                          1.8
                                                                                          2.0
```

Z

```
mcmc_U <- mcmc_vals[substr(rownames(mcmc_vals), 1, 1) == "U", "Mean"]</pre>
mcmc_U <- matrix(mcmc_U, nrow=K)</pre>
mcmc_V <- mcmc_vals[substr(rownames(mcmc_vals), 1, 1) == "V", "Mean"]</pre>
mcmc_V <- matrix(mcmc_V, nrow=K)</pre>
{\tt mcmc} U
                        [,2]
                                   [,3]
                                             [,4]
                                                        [,5]
                                                                  [,6]
##
              [,1]
                                                                             [,7]
## [1,] 0.6422038 0.9407988 0.5646119 0.7001590 0.4680595 0.2468498 0.3582272
## [2,] 0.8403647 0.9177366 0.6096788 0.2682265 0.6737750 0.2386912 0.2802439
                        [.9]
             [,8]
                                 Γ.107
## [1,] 0.7960865 0.7414597 0.4121112
## [2,] 0.5449510 0.2687351 0.5818920
p \leftarrow seq(0, 1, length = 100)
for (s in 1:S) {
  for (k in 1:K) {
    if (k == 1) {
      # plot mcmc mean U, V
      plot(p, dbeta(p, mcmc_U[k,s] * mcmc_V[k,s], (1-mcmc_U[k,s])*mcmc_V[k,s]),
           main = paste0("S", s),
           ylab = "density", xlab = "w", type = "l", col = k,
           ylim = c(0, 12)
      # plot truth
      lines(p, dbeta(p, a[k,s], b[k,s]), type = "l", col = k, lty=2)
      # add legend
      legend(x = "topleft",
             legend = paste0(c("mcmc k", "true k"), rep(1:K, each=2)),
             col = rep(1:K, each=2),
             lty = rep(1:2, K),
             cex=0.8)
    } else {
      # plot mcmc mean U, V
      lines(p, dbeta(p, mcmc_U[k,s] * mcmc_V[k,s], (1-mcmc_U[k,s])*mcmc_V[k,s]),
            type = "l", col = k)
      # plot truth
      lines(p, dbeta(p, a[k,s], b[k,s]), type = "1", col = k, lty=2)
    }
  }
}
```

