Finite mixture models

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Categorical distribution

Let $Z \sim Cat(H,p)$ represent a categorical distribution with

- $P(Z=h)=p_h$ for $h=1,\ldots,H$ and

Example: discrete choice model

Suppose we have a set of H categories and we label these $1,\ldots,H$. Independently, consumers choose one of the H categories with the same probability. Then a reasonable model is $Z_i \overset{ind}{\sim} Cat(H,p)$.

Multinomial distribution

If we count the number of times the consumer chose each category, i.e.

$$Y_h = \sum_{i=1}^n I(Z_i = h),$$

then the result is the multinomial distribution, i.e. $Y \sim Mult(n, p)$. The multinomial distribution has probability mass function

$$p(y; n, p) = \frac{n!}{y_1! \cdots y_H!} \prod_{h=1}^{H} p_h^{y_h}$$

which has

- $E[Y_i] = np_i$,
- $V[Y_i] = np_i(1 p_i)$, and
- $Cov[Y_i, Y_j] = -np_ip_j$ for $(i \neq j)$.

A special case is H=2 which is the binomial distribution.

Dirichlet distribution

The Dirichlet distribution (named after Peter Gustav Lejeune Dirichlet), i.e. $P \sim Dir(a)$, is a probability distribution for a probability vector of length H. The probability density function for the Dirichlet distribution is

$$p(P;a) = \frac{\Gamma(a_1 + \dots + a_H)}{\Gamma(a_1) \cdots \Gamma(a_H)} \prod_{h=1}^H p_h^{a_h - 1}$$

where $p_h \geq 0$, $\sum_{h=1}^{H} p_h = 1$, and $a_h > 0$.

Letting $a_0 = \sum_{h=1}^{H} a_h$, then some moments are

- $E[p_h] = \frac{a_h}{a_0}$,
- $V[p_h] = \frac{a_h(a_0 a_h)}{a_0^2(a_0 + 1)}$,
- $Cov(p_h, p_k) = -\frac{a_h a_k}{a_0^2(a_0+1)}$, and
- $\operatorname{mode}(p_h) = \frac{a_h 1}{a_0 H}$ for $a_h > 1$.

A special case is H=2 which is the beta distribution.

Conjugate prior for multinomial distribution

The Dirichlet distribution is the natural conjugate prior for the multinomial distribution. If

$$Y \sim Mult(n,\pi)$$
 and $\pi \sim Dir(a)$

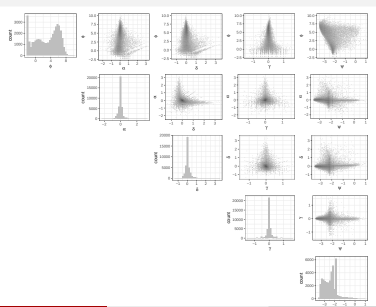
then

$$\pi|y \sim Dir(a+y).$$

Some possible default priors are

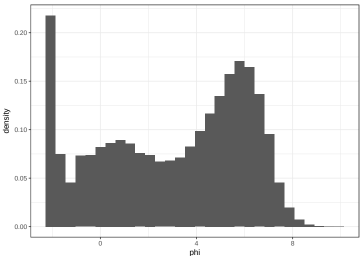
- a=1 which is the uniform density over π ,
- a = 1/2 is Jeffreys prior for the multinomial, and
- a=0, an improper prior that is uniform on $\log(\pi_h)$. The resulting posterior is proper if $y_h > 0$ for all h.

Complicated distributions



Finite mixtures

Let's focus on modeling the univariate distribution for $\boldsymbol{\phi}$



Finite mixture

A model for the marginal distribution for $Y_i = \phi_i$ is

$$Y_i \stackrel{ind}{\sim} \sum_{h=1}^{H} \pi_h N(\mu_h, \sigma_h^2)$$

where $\sum_{h=1}^{H} \pi_h = 1$.

Alternatively, we can introduce a latent variable $\zeta_i=h$ if observation i came from group h. Then

$$Y_i|\zeta_i = z \stackrel{ind}{\sim} N(\mu_z, \sigma_z^2)$$

 $\zeta_i \stackrel{ind}{\sim} Cat(H, \pi)$

where $\zeta \sim Cat(H,\pi)$ is a categorical random variable with $P(\zeta = h) = \pi_h$ for $h = 1, \dots, H$ and $\pi = (\pi_1, \dots, \pi_H)$.

A possible prior

Let's assume

$$\pi \sim Dir(a)
\mu_h | \sigma_h^2 \stackrel{ind}{\sim} N(m_h, v_h^2 \sigma_h^2)
\sigma_h^2 \stackrel{ind}{\sim} IG(c_h, d_h)$$

and π is independent of $\mu=(\mu_1,\ldots,\mu_H)$ and $\sigma^2=(\sigma_1^2,\ldots,\sigma_H^2)$.

Commonly, we have $m_h=m$, $v_h=v$, $c_h=c$, and $d_h=d$. If the data have been standardized (scaled and centered), a reasonable default prior is

- m = 0,
- v = 1,
- c = 2,
- d = 4, (BDA3 pg 535) and
- a is 1/H (BDA3 pg 536).

MCMC

The steps of a Gibbs sampler with stationary distribution

$$p(\pi, \mu, \sigma^2, \zeta|y) \propto p(y|\zeta, \mu, \sigma^2)p(\zeta|\pi)p(\mu|\sigma^2)p(\sigma^2)p(\pi)$$

has steps

1. For $i=1,\ldots,n$, sample ζ_i from its full conditional (as the ζ are conditionally independent across i):

$$P(\zeta_i = h | \ldots) \propto \pi_h N(y_i; \mu_h, \sigma_h^2)$$

- 2. Jointly sample π and μ, σ^2 as they are conditionally independent.
 - a. Sample $\pi \sim Dir(a+Z)$ where $n=(Z_1,\ldots,Z_H)$ and $Z_h=\sum_{i=1}^n \mathrm{I}(\zeta_i=h)$.
 - b. For $h=1,\ldots,H$, sample μ_h,σ_h^2 from their full conditional (as these are conditionally independent across h):

$$\sigma_h^2 \overset{ind}{\sim} IG(c_h', d_h') \quad \mu_h | \sigma_h^2 \overset{ind}{\sim} N(m_h', v_h'^2 \sigma_h^2)$$

where

$$\begin{array}{ll} v_h'^2 &= (1/v_h^2 + Z_h)^{-1} \\ m_h' &= v_h'^2 (m_h/v_h^2 + Z_h \overline{y}_h) \\ c_h' &= c_d + Z_h/2 \\ d_h' &= d_h + \frac{1}{2} \left(\sum_{i:\zeta_i = h} (y_i - \overline{y}_h)^2 + \left(\frac{Z_h}{1 + Z_h/v_h^2} \right) (\overline{y}_h - m_h)^2 \right) \\ \overline{y}_h &= \frac{1}{Z_h} \sum_{i:\zeta_i = h} y_i \end{array}$$

```
library("rjags")
jags_model = "
model {
 for (i in 1:n) {
   y[i] ~ dnorm(mu[zeta[i]], tau[zeta[i]])
   zeta[i] ~ dcat(pi[])
 for (i in 1:H) {
   mu[i] ~ dnorm(0,1e-5)
   tau[i] ~ dgamma(1,1)
    sigma[i] <- 1/sqrt(tau[i])
 pi ~ ddirich(a)
```

```
tmp = hat[sample(nrow(hat), 1000),]
dat = list(n=nrow(tmp), H=3, y=tmp$phi, a=rep(1,3))
```

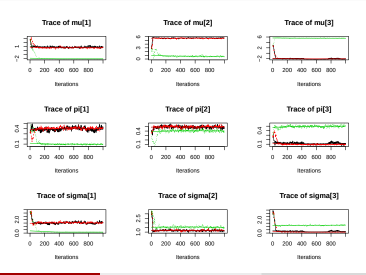
```
jm = jags.model(textConnection(jags_model), data = dat, n.chains = 3)
r = coda.samples(jm, c('mu', 'sigma', 'pi'), 1e3)
```

Convergence diagnostics

```
gelman.diag(r)
## Error in chol.default(W): the leading minor of order 6 is not positive definite
gelman.diag(r, multivariate=FALSE)
## Potential scale reduction factors:
##
           Point est. Upper C.I.
## mu[1]
                21.76
                         55.54
## mu[2]
               41.55
                      90.41
## mu[3]
                91.54
                      222,09
## pi[1]
               11.79
                       29.52
## pi[2]
               3.46
                         6.36
                      40.16
## pi[3]
              18.63
## sigma[1]
              13.70
                        35.60
## sigma[2]
               4.26
                         8.87
## sigma[3]
               13.93
                          37.09
```

Convergence diagnostics (2)

plot(r, density=FALSE)



Prior distributions

The parameters of the model are unidentified due to label-switching, i.e.

$$Y_i \stackrel{ind}{\sim} \sum_{h=1}^{H} \pi_h N(\mu_h, \sigma_h^2) \stackrel{d}{=} \sum_{h'=1}^{H} \pi_{h'} N(\mu_{h'}, \sigma_{h'}^2)$$

for some permutation h'.

 $\mu_1 < \mu_2 < \cdots < \mu_H$.

One way to resolve this issue is to enforce identifiability in the prior. For example, in one-dimension, we can order the component means:

To ensure the posterior is proper

- ullet Maintain proper prior for π
- Ensure proper prior for ratios of variances (perhaps by ensuring prior is proper for variances themselves)

Two conditionally conjugate prior options

Option 1:

$$Dir(\pi; a)I(\mu_1 < \dots < \mu_H) \prod_{h=1}^H N(\mu_h; m_h, v_h^2) IG(\sigma_h^2; c_h, d_h)$$

Option 2:

$$Dir(\pi; a)I(\mu_1 < \dots < \mu_H) \prod_{h=0}^{1} N(\mu_h; m_h, v_h^2 \sigma_h^2) IG(\sigma_h^2; c_h, d_h)$$

```
library("rjags")
jags_model = "
model {
    for (i in 1:n) {
        y[i] ~ dnorm(mu[zeta[i]], tau[zeta[i]])
        zeta[i] ~ dcat(pi[])
    }

    for (i in 1:H) {
        mu0[i] ~ dnorm(0,1e=5)
        tau[i] ~ dgamma(1,1)
        sigma[i] <- 1/sqrt(tau[i])
    }

    mu[1:H] <- sort(mu0)
    pi ~ ddirich(a)
}"</pre>
```

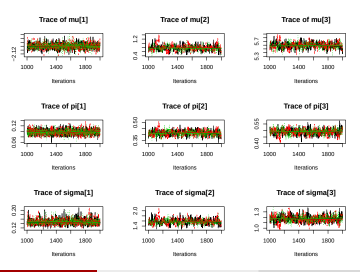
```
jm = jags.model(textConnection(jags_model), data = dat, n.chains = 3)
r = coda.samples(jm, c('mu', 'sigma', 'pi'), 1e3)
```

Convergence diagnostics

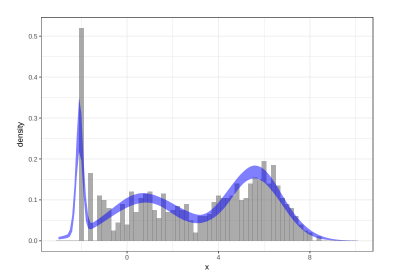
```
gelman.diag(r)
## Error in chol.default(W): the leading minor of order 6 is not positive definite
gelman.diag(r, multivariate=FALSE)
## Potential scale reduction factors:
##
           Point est. Upper C.I.
## mu[1]
                 1.00
                            1.00
## mu[2]
                 1.01
                           1.04
## mu[3]
                 1.00
                        1.01
                 1.00
                           1.00
## pi[1]
## pi[2]
                 1.01
                           1.03
## pi[3]
                 1.01
                           1.02
## sigma[1]
                1.00
                           1.00
## sigma[2]
               1.01
                           1.03
## sigma[3]
                 1.01
                            1.02
```

Convergence diagnostics (2)

plot(r, density=FALSE)



Posterior on data density



Group membership

Group membership can be obtained using the ζ_i , e.g.

$$P(\text{gene }i\text{ in cluster }h) = P(\zeta_i = h|y) \approx \sum_{m=1}^M \mathrm{I}\left(\zeta_i^{(m)} = h\right).$$

```
## parameter p1 p2 p3
## 1 zeta[10] 0.000 0.002666667 0.99733333
## 2 zeta[10] 0.000 0.00233333 0.90966667
## 3 zeta[100] 0.000 0.206000000 0.79400000
## 4 zeta[100] 0.000 0.95966667 0.04033333
## 5 zeta[101] 0.000 0.000 0.000000000 0.000000000
## 6 zeta[102] 0.929 0.071000000 0.00000000
```

Clustering

Genes can then be clustered by assigning them to a group based on their posterior probabilities of group membership, i.e. for gene i, we assign the group according to

$$\operatorname{argmax}_{h} P(\zeta_{i} = h|y).$$

Unfortunately clustering is extremely sensitive to the parametric model chosen, e.g. normal in this example, and the cluster could change dramatically with a different choice, e.g. t.

Choosing H

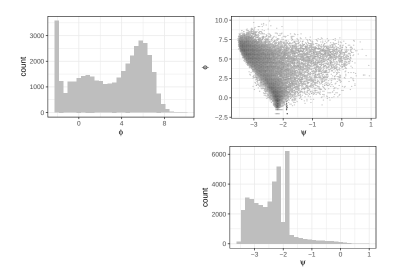
When using finite mixture models one of the key choices is to choose H, the number of clusters.

- A Bayesian approach would place a prior on H, e.g. a Poisson or truncated Poisson, and then use reversible jump MCMC to estimate it.
- ullet A more pragmatic approach is to start with a small H and then determine whether there is some feature of the data that is not being adequately addressed, e.g. via posterior predictive pvalues.
- An empirical Bayes finds an MLE (or MAP) via

$$\hat{H} = \mathrm{argmax}_H p(y|H) = \int p(y|\pi,\mu,\sigma^2,H) p(\pi,\mu,\sigma^2|H) d\pi d\mu d\sigma^2$$

and then condition on \hat{H} in the analysis. Typically this MLE (or MAP) is found via the EM algorithm.

Multivariate density estimation



Finite mixture

A model for the joint distribution for $Y_i = (\phi_i, \psi)^{\top}$ is

$$Y_i \stackrel{ind}{\sim} \sum_{h=1}^{H} \pi_h N(\mu_h, \Sigma_h)$$

where $\sum_{h=1}^{H} \pi_h = 1$.

Alternatively, we can introduce a latent variable $\zeta_i=h$ if observation i came from group h. Then

$$Y_i|\zeta_i = z \stackrel{ind}{\sim} N(\mu_z, \Sigma_z)$$

 $\zeta_i \stackrel{ind}{\sim} Cat(H, \pi)$

where $\zeta \sim Cat(H,\pi)$ is a categorical random variable with $P(\zeta = h) = \pi_h$ for $h = 1, \dots, H$ and $\pi = (\pi_1, \dots, \pi_H)$.

A possible prior

Let's assume

$$\pi \sim Dir(a)$$

$$\mu_h | \Sigma_h \stackrel{ind}{\sim} N_p(m_h, v_h^2 \Sigma_h)$$

$$\Sigma_h \stackrel{ind}{\sim} IW(D_h, c_h)$$

where $c_h > p-1$ is the degrees of freedom and D is the scale matrix. The mean of this distribution is $D_h/(c_h-p-1)$ for $c_h > p+1$.

MCMC

The steps of a Gibbs sampler with stationary distribution

$$p(\pi,\mu,\Sigma,\zeta|y) \propto p(y|\zeta,\mu,\Sigma) p(\zeta|\pi) p(\mu|\Sigma) p(\Sigma) p(\pi)$$

has steps

1. For $i=1,\ldots,n$, sample ζ_i from its full conditional

$$P(\zeta_i = h|\ldots) \propto \pi_h N(y_i; \mu_h, \Sigma_h)$$

- 2. Jointly sample π and μ,σ^2 as they are conditionally independent.
 - a. Sample $\pi \sim Dir(a+Z)$ where $Z=(Z_1,\ldots,Z_H)$ and $Z_h=\sum_{i=1}^n \mathrm{I}(\zeta_i=h)$.
 - b. For $h=1,\ldots,H$, sample μ_h,Σ_h from their full conditional

$$\Sigma_h \stackrel{ind}{\sim} IW(D_h', c_h') \quad \mu_h | \Sigma_h \stackrel{ind}{\sim} N(m_h', v_h'^2 \Sigma_h)$$

where

$$\begin{array}{ll} v_h'^2 &= (1/v_h^2 + Z_h)^{-1} \\ m_h' &= v_h'^2 (m_h/v_h^2 + Z_h \overline{y}_h) \\ c_h' &= c_d + Z_h \\ D_h' &= D_h + \sum_{i:\zeta_i = h} (y_i - \overline{y}_h) (y_i - \overline{y}_h)^\top + \left(\frac{Z_h}{1 + Z_h/v_h^2}\right) (\overline{y}_h - \mu_h) (\overline{y}_h - \mu_h)^\top \\ \overline{y}_h &= \frac{1}{Z_h} \sum_{i:\zeta_i = h} y_i \end{array}$$

```
library("rjags")
joint_mixture_model = "
model {
  for (i in 1:n) {
    y[i,1:p] ~ dmnorm(mu[,zeta[i]], Tau[,,zeta[i]])
    zeta[i] ~ dcat(pi[])
  for (h in 1:H) {
    mu[1:p,h] ~ dmnorm(mu0,Tau[,,h])
    Tau[1:p.1:p.h] ~ dwish(D[.].c)
    Sigma[1:p,1:p,h] <- inverse(Tau[,,h])
  pi ~ ddirich(a)
tmp = hat[sample(nrow(hat), 1000),]
dat = list(n=nrow(tmp), y = tmp[,c('phi','psi')], p=2, H=10)
dat$a = rep(1/dat$H, dat$H)
dat$D = diag(1, dat$p)
dat$c = dat$p+1
dat\$mu0 = c(3,0)
```