## Exercises: multiple testing, FDR

1. Consider the following setting in the context of "multiple hypothesis tests". Let  $i=1,\ldots,n$  index individuals and  $j=1,\ldots,m$  index genes (or pixels in an image if you prefer). Assume we have measurements on each individual at each gene for "treatments" k=0,1. Let  $Y_{ijk}$  denote the measurement on individual i, gene j, treatment k, and let  $D_{ij}$  denote the difference between the measurements in the two treatments:  $D_{ij} := Y_{ij0} - Y_{ij1}$ . We will assume that D is sufficient for all our inferences, and so you can forget about Y now and work only with D: I just wanted you to understand where D might come from in principle.

We will assume a model for D:

$$D_{ij}|\beta,\sigma \sim N(\beta_j,\sigma_j^2)$$
 (1)

#m tests

where  $\beta = (\beta_1, \dots, \beta_m)$  is a vector of "treatment effect"s, where  $\beta_j$  is the effect at gene j, and  $\sigma = (\sigma_1, \dots, \sigma_m)$  is a vector of standard deviation parameters. For each gene j we wish to test the null  $H_j$ :  $\beta_j = 0$  (that is, that there is no treatment effect). For simplicity you can assume that  $\sigma_j = 1$  is known for all j. You can also assume that n = 10 and m = 1,000.

Assume that the <u>true effects</u>  $\beta_i$  are independent, and <u>identically distributed</u>, with iid

$$\frac{\text{i i d}}{\beta_j} \sim \frac{\pi_0 \delta_0 + (1 - \pi_0) N(0, \sigma_b^2)}{\pi_0 \delta_0 + (1 - \pi_0) N(0, \sigma_b^2)}.$$
(2)

where  $\delta_0$  denotes a point mass on zero. That is,  $\beta_j = 0$  with probability  $\pi_0$ , and  $\beta_j \sim N(0, \sigma_b^2)$  with probability  $1 - \pi_0$ .

- i) Write an R function to simulate data D under this model, for user-specified  $\pi_0$  and  $\sigma_b$ . The function should take  $\pi_0$  and  $\sigma_b$  as input, and return a list, with elements D (a matrix) and  $\beta$  (a vector).
- ii) Write an R function to compute a p value  $p_i$  for each column of the data matrix D, testing  $H_0: \beta_j = 0$ . This function should take as input the data matrix D and output a vector of p values. You can use any reasonable two-sided test, but state which test you use. Apply your R function to data simulated under a)  $\pi_0 = 1$ , b)  $\pi_0 = 0.5, \sigma_b = 3$ ; c)  $\pi_0 = 0, \sigma_b = 3$ . Provide histograms of the p values in each case and comment on their distributions.

- iii) Write an R function to apply the Benjamini-Hochberg rule to control FDR at a user-specified level  $\alpha$ . This function should input a vector of p values, and a level  $\alpha$ , and output a vector of binary (0/1) indicators,  $\gamma = (\gamma_1, \ldots, \gamma_m)$  say, where  $\gamma_i = 1$  indicates that the rule would reject  $H_i: \beta_i = 0$ .
- iv) Write an R function to compute the empirical False Discovery Rate (i.e. the number of false discoveries divided by the number of discoveries) for any given value for the vector  $\beta$  of true values of  $\beta$ , and the vector  $\gamma$  of reject decisions. That is, the function should return V/R in the notation of the notes. Remember to deal correctly with the special case of no discoveries, R = 0.
- v) Perform a simulation study to estimate the actual FDR (E(V/R)) achieved by the BH rule in the three cases a), b) and c) above. In each case perform the test procedure for different levels  $\alpha$ , and plot the estimated E(V/R) as a function of  $\alpha$  (say for  $\alpha = (0.05, 0.1, \dots, 0.5)$ ). Comment on the results. [NOTE: to estimate the actual FDR you have to estimate E(V/R) where the expectation is over datasets D. To do this you will want to do a simulation study where you simulate a large number of datasets D, not just one dataset!]
- vi) Repeat the simulation study, but this time estimate the pFDR instead of the FDR, and plot this as a function of  $\alpha$ .
- 2. The **qvalue** package in R implements Storey's approach to estimating FDR. To install this package use

```
source("http://bioconductor.org/biocLite.R")
biocLite("qvalue")
library("qvalue")
```

The package takes a vector of p values, and outputs a list which includes an estimate of  $\pi_0$  (obtained using the p values near 1) and a vector of q values. Try, for example, for a vector of p values p,

```
res=qvalue(p)
res$pi0
res$qvalues
```

The q value for a particular observation is an estimate of the pFDR if you reject all things that are as or more significant than that observation. You can convert the vector of q values into a list of reject decisions at a given  $\alpha$  level (the  $\gamma$  vector above) using, say,

## compute.gamma=function(q,alpha){return(q<alpha)}</pre>

- i) Repeat the simulation study above, using qvalue instead of the BH procedure. Produce plots of the FDR vs the  $\alpha$  level for qvalue and compare them with those obtained for BH.
- ii) Perform a simulation study (e.g. by modifying the simulations you have already performed), to see how accurately qvalue is able to estimate the proportion of nulls  $\pi_0$ . Try varying  $\pi_0$  from 0 to 1 for at least 3 different values of  $\sigma_b$ , and in each case provide plots of the true  $\pi_0$  vs the estimated  $\pi_0$  from qvalue. Comment on the results.
- 3. Now consider implementing an Empirical Bayes approach to this problem. To do so, given data D we will need two steps:
  - A Estimate the hyper parameters  $\pi_0, \sigma_b$  in (2) by maximum likelihood. Call the estimates  $\hat{\pi}_0, \hat{\sigma}_b$ .
  - B Compute the posterior distribution  $p(\beta_j|D,\hat{\pi}_0,\hat{\sigma}_b)$  for each j.

This question takes you through these two steps.

- i) Define  $\bar{D}_j = (1/n) \sum_i D_{ij}$ . Show that the vector  $\bar{D} := (\bar{D}_1, \dots, \bar{D}_m)$  is sufficient for  $\beta$ . That is,  $p(D|\beta) \propto p(\bar{D}|\beta)$  where the constant of proportionality does not depend on  $\beta$ . [This means that, as far as inference for  $\beta$  is concerned, the likelihood  $p(D|\beta)$  is equivalent to the likelihood  $p(\bar{D}|\beta)$ , so from now on you can treat  $\bar{D}$  as your data instead of D.]
- ii) Derive an expression for the log-likelihood  $l(\pi_0, \sigma_b) := \log(p(\bar{D}|\pi_0, \sigma_b))$ . [Hint: note that the  $\bar{D}_i$  are independent given  $\pi_0, \sigma_b$ ]
- iii) Write an R function to compute the log-likelihood  $l(\pi_0, \sigma_b)$ , or alternatively  $l(\theta_1, \theta_2)$  where  $\theta_1 = \log(\pi_0/(1 \pi_0))$ ,  $\theta_2 = \log(\sigma_b)$ . [The motivation for this reparameterization is that  $\theta_1, \theta_2$  can take any value on the real line.] Try using the R function optimize

- (or another method if you prefer) to maximize the likelihood over  $\pi_0$ ,  $\sigma_b$  (or  $\theta_1$ ,  $\theta_2$ ). [You may or may not find that this works... it is a somewhat tricky numerical problem. The reparameterization may help. Alternatively if you know about the EM algorithm you can try that.]
- iv) Derive the posterior distribution  $\beta_j|D, \pi_0, \sigma_b$ . Hint: this posterior should be a mixture of a point mass at zero and a normal distribution. It may help to first derive  $p(\beta_i = 0|D, \pi_0, \sigma_b)$ , and then  $p(\beta_i|D, \pi_0, \sigma_b, \beta_i \neq 0)$ .
- v) Implement a method that computes  $p(\beta_i = 0|D, \hat{\pi}_0, \hat{\sigma}_b)$ . Implement another method that takes these probabilities and rejects those tests j for which this probability is  $< \alpha$ . Add this method to your simulation study and see how it performs. (If you are unable to get the optimization for  $\pi$ ,  $\sigma_b$  to work then you can "cheat" in this step and use the true value of  $\pi$ ,  $\sigma_b$ .)