## M.PHIL. IN STATISTICAL SCIENCE

Wednesday 4 June 2008 1.30 to 3.30

Script paper

## APPLIED BAYESIAN STATISTICS

Attempt no more than **THREE** questions.

There are FOUR questions in total.

The questions carry equal weight.

When asked to provide code, you need only give the approximate form for the statements: precise syntax is not expected.

**STATIONERY REQUIREMENTS** SPECIAL REQUIREMENTS Cover sheet None Treasury tag

> You may not start to read the questions printed on the subsequent pages until instructed to do so by the Invigilator.

1 Much of biomedical science is concerned with finding true relationships between 'exposures' (say genes) and 'outcomes' (say diseases). Ioannidis (2005) published a much-cited paper *Why most published research findings are false*, in which he supposed that relationships being studied were either 'true' (denoted T) or 'false' (denoted F): a false relationship corresponds to a null hypothesis  $H_0$ , while a true relationship corresponds to an alternative hypothesis  $H_1$ . He assumed, in any particular field, a ratio R between the proportion of true relationships and the proportion of false relationships. Furthermore, he assumed that each study was reported as being either 'positive' (denoted P), or 'negative' (denoted N), and that all studies were designed with Type 1 error  $\alpha$  (the probability that a false relationship is detected in the study - a 'false positive result') and Type II error  $\beta$  (the probability that a relationship that truly does exist is not detected in the study - a 'false negative result').

(a) What is the probability that a relationship is false, given that a study reported it as positive?

(b) Show that R has to be at least  $\alpha/(1-\beta)$ , in order for a reported positive result to be more likely true than false.

Suppose a study is designed to have a test statistic Y with a N(0,1) distribution under the null hypothesis that there is no relationship, and the alternative hypothesis is that Y has a N(2.49,1) [NB  $\Phi(1.65) \approx 0.95, \Phi(-0.84) \approx 0.2$ , where  $\Phi$  is the standard normal cumulative distribution function].

(c) If a positive result is declared if Y > 1.65, show that  $\alpha = 0.05, \beta = 0.20$ .

(d) If we observe y, show that the Bayes factor for the alternative hypothesis against the null hypotheses is  $e^{2.49(y-1.245)}$ .

(e) In a succeeding paper, Goodman and Greenland criticised Ioannidis' analysis, saying that it is not sensible to summarise results by just whether they are 'significant' or not. In the light of the preceding analysis, do you think this is a reasonable criticism?

Suppose that n independent studies, each designed with Type I error  $\alpha$  and Type II error  $\beta$ , are performed to test the same relationship.

(f) What is the chance that, if the relationship does not exist, at least one will show a positive result?

(g) If all we know is that there is at least one positive result among the n studies, what are the odds that this is a true finding?

(h) What happens to these odds as n increases? Why does Ioannidis claim that 'The hotter a scientific field (with more scientific teams involved), the less likely the research findings are to be true'?

(i) Goodman and Greenland also say it is unrealistic to assume that all we know is that there is at least one positive result among *n* studies. If we read a published account that claims a positive relationship based on the *full* results of many studies carried out in a field, should that increase or decrease our confidence in the positive finding, relative to what we would feel after reading a single study? **2** Suppose  $y_i$  is the numbers of cases of MRSA (a bacterial infection) in a single year, in the *i*th of *n* hospitals, each drawn respectively from a Poisson distribution with mean  $\theta_i$ . The  $\theta_i$ 's are assumed drawn from an exponential distribution  $p(\theta_i|\psi) = \psi e^{-\psi \theta_i}; \theta_i > 0$ .

(a) Draw a directed graph for this model.

(b) When using Gibbs sampling for this model, prove that the full conditional distribution for  $\theta_i$  depends only on  $y_i$  and  $\psi$ , and derive its form. [The Gamma distribution has the form  $p(x) = b^a x^{a-1} e^{-bx} / \Gamma(a)$ .]

(c) What does  $y_i$  contribute to the likelihood for  $\psi$ ? Show that contribution of  $y_i$  to the likelihood for  $\phi = 1/(1+\psi)$  is  $(1-\phi)\phi^{y_i}$ .

(d) What is the conjugate prior for  $\phi$ ?

Suppose the logarithm of the hyper-parameter  $\psi$  is given an improper uniform prior

 $p(\log \psi) \propto 1$ .

- (e) What is the induced prior for  $\phi$ ?
- (f) Find the posterior distribution for  $\phi$  using this prior.
- (g) Show that the posterior expectation of  $\theta_i$  is a weighted average of  $y_i$  and  $\overline{y}$ .

(h) In real hospitals would you expect the assumptions about the prior distribution for the  $\theta_i$ 's to be correct?

(i) What might be a more reasonable model?

**3** The testosterone/epitestosterone (T/E) ratio is used as a biomarker for detecting athletes' abuse of testosterone, and a Bayesian model has been suggested for monitoring repeated measures of the logarithm of an athlete's T/E ratio.

An athlete *i* is assumed, if he/she is not taking additional testosterone, to have log(T/E) measurements drawn independently from a normal distribution with mean  $\theta_i$  and standard deviation  $\sigma_i$ . We assumed that the population of  $\theta_i$ 's are independently drawn from a N( $\mu, \tau^2$ ) distribution.

Suppose for the moment that we know values for  $\mu$  and  $\tau$ . We select an athlete *i* for whom we have no data, and assume we know  $\sigma_i$ .

(a) What is the predictive distribution for  $Y_{i1}$ , the first observation to be taken on athlete *i*?

(b) We want to set an upper limit  $u_{i1}$ , so that the predictive probability that an observation lies above that limit, if the athlete is not abusing testosterone, is 1/1000. Write down an explicit expression for  $u_{i1}$  in terms of  $\mu, \tau, \sigma_i$  and  $\Phi$ , where  $\Phi$  is the standard normal distribution function.

Suppose we now take a measurement  $y_{i1}$  on athlete *i*.

(c) What is the posterior distribution for  $\theta_i$ ?

(d) Denoting the posterior mean and variance of  $\theta_i$  by  $m_{i1}$  and  $v_{i1}$  respectively, what is the predictive distribution for  $Y_{i2}$ , a future second measurement?

(e) What monitoring limit  $u_{i2}$  would you set for  $Y_{i2}$ , such that there was a 1/1000 predicted probability that an 'innocent' athlete would exceed that limit?

(f) Explain in words how the monitoring limits will change as more data becomes available on athlete i.

In fact the parameters  $\mu, \tau$  and  $\sigma$ 's are unknown, but there is a historical database available of repeated measurements on athletes who were not abusing testosterone, denoted  $y_{j1}, ..., y_{jn_j}$ , for j in 1 to J. The following code has been suggested for analysing the database.

```
for (j in 1:J) {
   for (k in 1:n [j]) {
       Y [j,k] ~ dnorm (theta [j], invsigma2 [j])
   }
   theta [j]
                        ~ dnorm (mu, invtau2)
   log (invsigma2 [j]) <- - log.sigma2 [j]</pre>
   log.sigma2 [j]
                        ~ dnorm (phi, invpsi2)
                                                     # (1)
}
mu
     ~ dunif (-100, 100)
invtau2 <- 1 / (tau*tau) ; tau ~ dunif (0, 100)
                                                     # (2)
phi ~ dunif (-100, 100)
invpsi2 <- 1 / (psi*psi) ; psi ~ dunif (0, 100)</pre>
                                                     # (3)
```

(g) Explain why this code may be reasonable, with special reference to the numbered lines of code.

Question continued/...

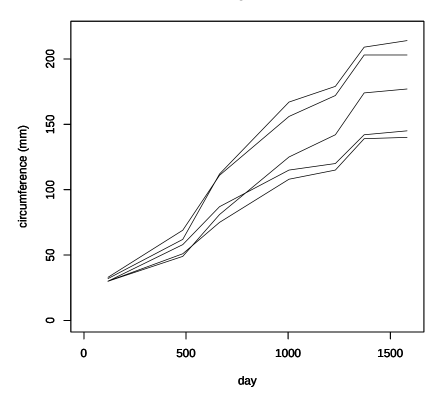
(h) Suppose you now have three measurements available on a new athlete, and you want to check whether the third observation is 'extreme' relative to the first two (*i.e.* outside a 99.9% prediction interval). How would you adapt the code to carry out this analysis?

The assumption that the  $\theta$ 's are normally distributed is questionable, and the following modification to the prior distribution has been suggested:

```
theta [i] ~ dnorm (mu , invtau2 [i])
invtau2 [i] <- lambda [i] / (4*tau*tau)
lambda [i] ~ dchisqr (4)</pre>
```

- (i) What prior distribution will this imply for each  $(\theta_i \mu)/\tau$ ?
- (j) Why might this be an appropriate model?
- (k) How might you assess if it was a more appropriate model for the historical data?

4 The graph below shows the circumferences in mm of 5 orange trees, each measured on day x.



**Orange trees** 

An investigator uses the following WinBUGS code to fit an independent 'logistic growth curve' to each tree (Model A).

```
for (i in 1:5) {
   for (j in 1:7) {
      Y [i,j] ~ dnorm (m [i,j], invsigma2)
      m [i,j] <- phi [i,1] / ( 1 + phi [i,2]</pre>
                                   * exp ( phi [i,3]
                                   * ( x [j] - mean (x []) ) / sd (x []) ) )
   }
   log ( phi [i,1] ) <-
                           theta [i,1]
   log ( phi [i,2] ) <- - theta [i,2]
         phi [i,3]
                      <- - theta [i,3]
   for (p in 1:3) {
      theta [i,p] ~ dunif (-100,100)
                                          *
   }
                                        #
                                          *
}
                                          *
                                        #
invsigma2 ~ dgamma (1.0E-3, 1.0E-3)
sigma <- 1 / sqrt (invsigma2)</pre>
```

(a) Why is x transformed?

(b) Interpret the parameters phi[i,1], and 1/(1 + phi[i,2]). Why do you think this is called a logistic curve?

(c) Do you think the prior distributions are reasonable?

The investigator decides to replace the starred lines of code in Model A by the following (Model B).

```
theta [i,1:3] ~ dmnorm (mu [], Omega [,])
}
for (p in 1:3) { mu[p] ~ dunif (-100,100) }
Omega [1:3, 1:3] ~ dwish (R [,], 3)
R [1,1] <- 1; R [1,2] <- 0; R [1,3] <- 0;
R [2,1] <- 0; R [2,2] <- 1; R [2,3] <- 0;
R [3,1] <- 0; R [3,2] <- 0; R [3,3] <- 1</pre>
```

Some of the output is shown below:

		theta [,1]		theta [,2]		theta	theta [,3]	
		mean	sd	mean	sd	mean	sd	
theta	[1,]	5.063	0.090	0.753	0.357	1.443	0.281	
theta	[2,]	5.392	0.059	0.679	0.252	1.601	0.223	
theta	[3,]	5.057	0.103	0.557	0.354	1.412	0.280	
theta	[4,]	5.427	0.053	0.673	0.234	1.715	0.227	
theta	[5,]	5.302	0.098	0.291	0.289	1.498	0.241	

(d) Comment on the appropriateness of the new code

(e) Why might it be reasonable to assume that all trees have the same values for theta[i,2] and theta[i,3]?

(f) How might you change the code to reflect this assumption (keeping theta[i,1] as a random effect)? Call this Model C.

After 50,000 iterations the following output for DICs were obtained for all three fitted models:

<pre>Dbar = post.mean of -2logL;</pre>									
		Dbar	рD	DIC					
Model	Α	248.1	12.0	260.1					
Model	В	244.2	13.9	258.1					
Model	С	247.3	8.2	255.6					

(g) Comment on the how the values of Dbar (the mean deviance), pD (the effective number of parameters) and DIC depend on the models.

(h) What might be considered unsatisfactory about Dbar and pD for Model A? Why might this have happened?

(i) In the final model, say very briefly how you would go about testing the normality assumption for the residual error. Would you also be testing the adequacy of the normal assumption for the random effects?

## END OF PAPER