# Analysing Spatial Data in R: Worked examples: (Bayesian) disease mapping II

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31 August 2007

# Bayesian Disease mapping

- Bayesian Estimation in Disease Mapping has been one of the leading topics in spatial statistics in the last 20 years
- Bayesian Hierarchical Models can be used to model complex data structures
- ► The Bayesian approach offers an easy approach to the estimation of complex models via Markov Chain Monte Carlo
- Spatial analysis of routinoulsy collected health data is standard practise nowadays
- Spatio-temporal models can be used
- ▶ Waller & Gotway (2004) and Banerjee et al. (2003) account for a comprehensive summary on spatial models

# Bayesian Inference

- ▶ Bayesian Inference is based on estimating the probability density of the parameters  $\theta$  in the model *after* observing the data, i.e., their posterior distributions:  $p(\theta|y)$
- $p(\theta|y)$  is usually difficult to derive:

$$p(\theta|y) = \frac{p(y|\theta)p(\theta)}{\int_{\theta} p(y|\theta)p(\theta)} \propto p(y|\theta)p(\theta)$$

- ▶  $p(y|\theta)$  is the likelihood of the model, which reflects the relationship between the data and the parameters
- $\triangleright$   $p(\theta)$  is the *prior* distribution of the parameters, which reflects the initial information on the parameters
- ▶ Usually,  $p(\theta|y)$  is computed by simulation using *Markov Chain Monte Carlo* techniques
- ► WinBUGS in a generic software to fit a wide range of models. It uses the Gibbs sampler for that.

## Benefits of Bayesian Inference

- Suitable framework to deal with a large number of problems
- Priors can be used to account for initial information (for example, spatial dependence)
- ▶ If no prior information is available, vague (or non-informative) priors can be used so that the posterior distribution *will only depend* on the data and the model.
- Multilevel models can be used: Bayesian Hierarchical Models
- Complex effects, such as spatial and/or temporal dependence, can be modeled easily
- When the posterior distribution is not in a closed form, different simulation techniques can be used to approximate them.
- Missing values are treated similarly as the parameters in the model

# Markov Chain Monte Carlo/Gibbs sampler

MCMC aims at simulating a series of values for the parameters in the model, so that, in the end, these values will be draws from the posterior distribution.

- Assign initial values to every parameter in the model (and missing values)
- At every step, Gibbs sampler simulates from the full conditional distribution:

$$p(\theta_i|\theta_{-i},y)$$

- ▶ After a *burn in* period, the simulated values are draws from the posterior  $p(\theta|y)$
- Convergence of the simulated values should be assessed

### **WinBUGS**

- BUGS stands for Bayesian inference Using Gibbs Sampler
- Developed at the MRC and Imperial College London
- ▶ Provides a generic language to Bayesian Hierarchical models
- Models can be specified graphically as well
- Several utilities to assess the convergence of the chain and display results
- GeoBUGS is an extension to deal with spatial models and maps
- PkBUGS is another extension to deal with Pharmacokinetics models
- ► A developer interface has been included so that the user can extend the range of functions available
- OpenBUGS is the open source version of WinBUGS

## Calling WinBUGS from R

- ▶ Packages R2WinBUGS and BRUGS can call WinBUGS and OpenBUGS from R
- R2WinBUGS calls WinBUGS using the scripting language and then reads the output log file
- BRUGS is an interface to the actual OpenBUGS (NOT WinBUGS) routines
- R2WinBUGS can run on several platforms (Windows, Linux/Unix, Mac)
- Other alternatives to call WinBUGS externally in different ways are available at http://www.mrc-bsu.cam.ac.uk/ bugs/winbugs/remote14.shtml

## Leukemia Cancer Data revisited

#### We need...

- Model specification (using the BUGS language)
- Mortality Data (in a list)
- Spatial data describing the neighbourhood structure, in a specific format
- Initial values of the parameters
- Optionally, we may want to export the map information to be used within WinBUGS

# Bayesian Spatial Modelling

$$O_{i} \sim Poisson(\mu_{i})$$

$$\mu_{i} = \theta_{i}E_{i}$$

$$\log(\theta_{i}) = \alpha + \beta X_{i} + u_{i} + v_{i}$$

$$u_{i} \sim Normal(0, \sigma_{u}^{2})$$

$$v_{i}|v_{-i} \sim Normal(\sum_{j\sim i} v_{j}/n_{i}, \sigma_{v}^{2}/n_{i})$$

$$f(\alpha) \propto 1$$

$$f(\beta) \propto 1$$

$$\sigma_{u}^{2} \sim Gamma^{-1}(.001, .001)$$

$$\sigma_{v}^{2} \sim Gamma^{-1}(.001, .001)$$

$$O_{i}$$

$$O_{i}$$

$$O_{i}$$

$$O_{i}$$

$$O_{i}$$

$$O_{i}$$

$$O_{i}$$

J. Besag, J. York, A. Mollie (1991). Bayesian image restoration, with two applications in spatial statistics (with discussion). *Annals of the Institute of Statistical Mathematics* **43(1)**, 1-59

# Model specification using the BUGS language

```
model{
        for(i in 1:N)
                O[i] ~ dpois(mu[i])
                mu[i]<-theta[i] * E[i]
            log(theta[i]) <- alpha + beta[1]*PCTAGE65P[i]+
                beta[2]*PCTOWNHOME[i]+beta[3]*AVGIDIST[i]+u[i] + v[i]
                u[i] ~ dnorm(0, precu)
                SMR[i]<- O[i] / E[i]
                prob[i] <- step(theta[i]-1)</pre>
        }
        v[1:N]~car.normal(adj[], weights[], num[], precv)
        alpha~dflat()
        for(i in 1:3) {beta[i] ~dflat()}
        precu~dgamma(0.001, 0.001)
        precv~dgamma(0.001, 0.001)
        sigmau<-1/precu
        sigmav<-1/precv
```

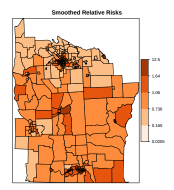
```
Preparing data...
    1.- Read maps
    > library(maptools)
    > nymap <- readShapePoly("NY8_utm18")</pre>
    2.- Create list of observed, expected
    > nymap$EXP <- nymap$POP8 * sum(nymap$Cases)/sum(nymap$POP8)</pre>
    3.- Create adjacency matrix
    > library(spdep)
    > nynb <- poly2nb(nymap)</pre>
    4.- Create weights
    > nyWBweights <- nb2WB(nynb)</pre>
    > d <- c(list(0 = nymap$Cases, E = nymap$EXP), N = 281,
          list(PCTAGE65P = nymap$PCTAGE65P, PCTOWNHOME = nymap$PCTOWNHOME,
               AVGIDIST = nymap$AVGIDIST))
    > inits1 <- list(alpha = 1, beta = c(0, 0, 0), u = rep(0, 1)
          281), v = rep(0, 281), precu = 1, precv = 1)
    > inits2 <- list(alpha = 10, beta = c(1, 1, 1), u = rep(1, 1)
          281), v = rep(1, 281), precu = 0.1, precv = 0.1)
```

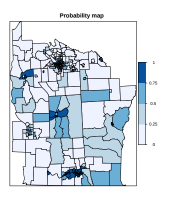
# Calling WinBUGS using R2WinBUGS

#### 5.- Call WinBUGS

```
> library(R2WinBUGS)
> mfile <- paste(getwd(), "/model.txt", sep = "", collapse = "")
> tdir <- paste(getwd(), "/NYoutput", sep = "", collapse = "")</pre>
> dir.create(tdir)
> res <- bugs(data = c(d, nyWBweights), inits = list(inits1,
      inits2), parameters.to.save = c("u", "v", "theta",
+
     "prob", "sigmau", "sigmav"), model.file = mfile,
+ working.directory = tdir, n.thin = 3, n.chains = 2,
+ n.iter = 6000, n.burnin = 3000)
6.- Add results to map object
> nymap$prob <- res$mean$prob</pre>
> nymap$theta <- res$mean$theta
> nymap$u <- res$mean$u
> nymap$v <- res$mean$v</pre>
> logfile <- paste(getwd(), "/NYoutput/log.txt", sep = "",</pre>
+ collapse = "")
> reslog <- bugs.log(file = logfile)</pre>
```

# Mapping the results





## Exporting the data to work directly with WinBUGS

1. Export the maps with spdep

```
> sp2WB(map = nymap, file = "NY_WB.txt")
```

- 2. Import map with WB first, and "reboot"
- 3. Use bugs.data (from R2WinBUGS) to create the files with data and initial values

```
> bugs.data(d)
```

- > file.rename("data.txt", "dataNY.txt")
- > bugs.data(nyWBweights)
- > file.rename("data.txt", "data-spatialNY.txt")
- > bugs.data(inits1)
- > file.rename("data.txt", "inits1NY.txt")
- > bugs.data(inits2)
- > file.rename("data.txt", "inits2NY.txt")

# Running WinBUGS directly

- 1. Open all needed files in WinBUGS
- 2. Check the model syntax
- 3. Load data (health and spatial)
- 4. Compile the model
- 5. Load inital values
- 6. Run the model (burn in period)
- 7. Monitor parameters of interest and DIC
- 8. Rerun the model
- 9. Assess convergence of the simulations
- 10. Show summary statistics of the parameters of the model
- 11. Display results on a map

#### Further references

- ▶ S. Banerjee, B.P. Carlin and A.E. Gelfand (2003). *Hierarchical Modeling and Analysis for Spatial Data*. Chapman & Hall.
- ▶ A.B. Lawson, W.J. Browne and C.L. Vidal Rodeiro (2003). Disease Mapping with WinBUGS and MLwiN. Wiley & Sons.
- ▶ OpenBUGS: http://mathstat.helsinki.fi/openbugs/
- R programming language: http://www.r-project.org
- D.J. Spiegelhalter, N.G. Best, B.P. Carlin and A. Van der Linde (2002). Bayesian Measures of Model Complexity and Fit (with Discussion), *Journal of the Royal Statistical Society*, Series B 64(4), 583-616.
- L.A. Waller and C.A. Gotway (2004). *Applied Spatial Statistics for Public Health Data*. Wiley & Sons.