# APTS 2013/14: Spatial and Longitudinal Data Analysis

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# Lecture topics

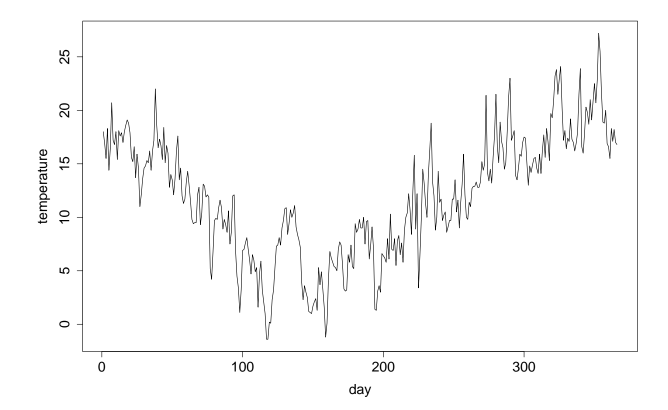
- Introduction: motivating examples
- Review of preliminary material
- Longitudinal data: linear Gaussian models; conditional and marginal models; why longitudinal and time series data are not the same thing.
- Continuous spatial variation: stationary Gaussian processes; variogram estimation; likelihood-based estimation; spatial prediction.
- Discrete spatial variation: joint versus conditional specification; Markov random field models.

- Spatial point patterns: exploratory analysis; Cox processes and the link to continuous spatial variation; pairwise interaction processes and the link to discrete spatial variation.
- Spatio-temporal modelling: spatial time series; spatiotemporal point processes; case-studies

### 1. Motivating examples

Example 1.1 Bailrigg temperature records

Daily maximum temperatures, 1.09.1995 to 31.08.1996

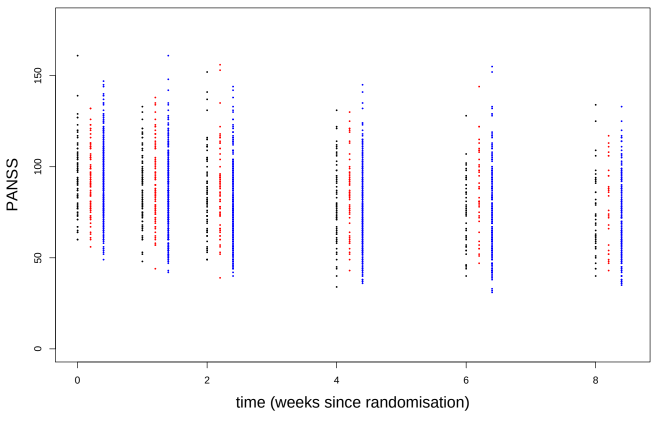


1.2 Schizophrenia clinical trial (PANSS)

- randomised clinical trial of drug therapies
- three treatments:
  - haloperidol (standard)
  - placebo
  - risperidone (novel)
- dropout due to "inadequate response to treatment"

Treatment	Number of non-dropouts at week					
	0	1	<b>2</b>	4	6	8
haloperidol	85	83	<b>74</b>	64	46	41
placebo	88	86	70	<b>56</b>	<b>40</b>	<b>29</b>
risperidone	<b>345</b>	<b>340</b>	307	<b>276</b>	<b>229</b>	199
total	<b>518</b>	509	451	396	<b>315</b>	<b>269</b>

#### Example 1.2: Schizophrenia trial data

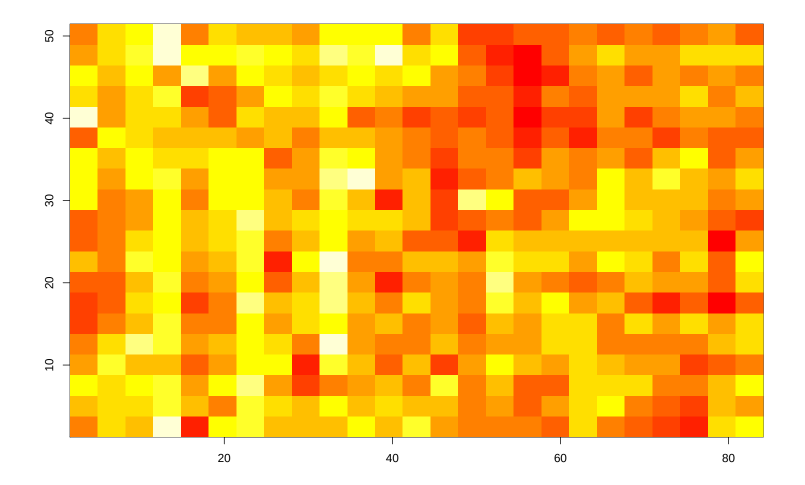


Diggle, Farewell and Henderson (2007)

Example 1.3 Wheat uniformity trial

- trial conducted at Rothamsted in summer of 1910
- wheat yield recorded in each of 500 rectangular plots (3.3m by 2.59m)
- same variety of wheat planted in all plots

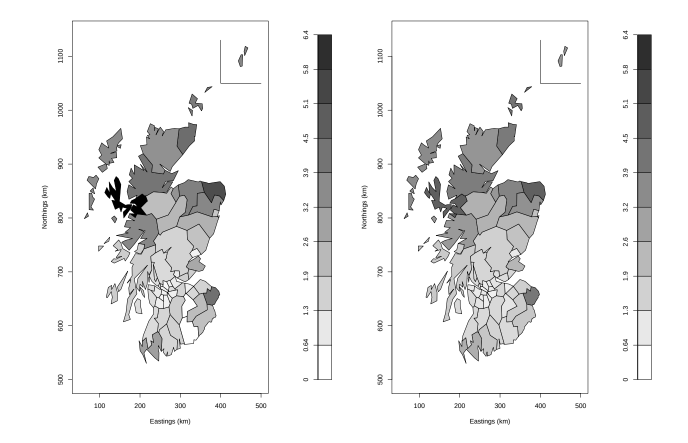
#### Mercer and Hall wheat yields



Mercer and Hall (1911)

#### **1.4 Cancer atlases**

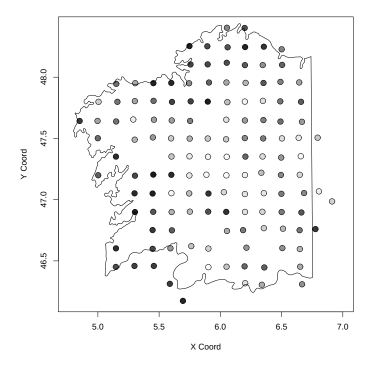
Raw and spatially smoothed relative risk estimates for lip cancer in 56 Scottish counties



#### Wakefield (2007)

**1.5 Galicia biomonitoring study** 

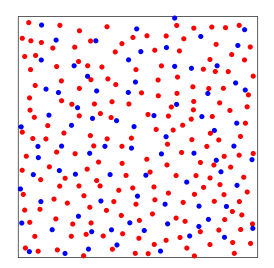
Lead concentrations measured in samples of moss, map shows locations and log-concentrations

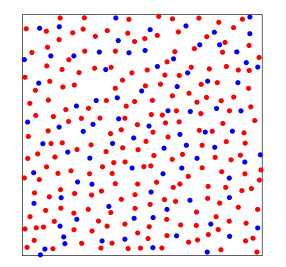


#### Diggle, Menezes and Su (2010)

**1.6 Retinal mosaics** 

Locations of two types of light-responsive cells in macaque retina (2 animals)





Eglen and Wong (2008)

# 2. Review of preliminary material

Time series

- trend and residual;
- autocorrelation;
- prediction;
- analysis of Bailrigg temperature data

# Analysis of Bailrigg temperature data

```
data<-read.table("maxtemp.data",header=F)
temperature<-data[,4]
n<-length(temperature)
day<-1:n
plot(day,temperature,type="l",cex.lab=1.5,cex.axis=1.5)
#
# plot shows strong seasonal variation,
# try simple harmonic regression
#</pre>
```

```
c1<-cos(2*pi*day/n)</pre>
s1<-sin(2*pi*day/n)</pre>
fit1<-lm(temperature~c1+s1)</pre>
lines(day,fit1$fitted.values,col="red")
#
# add first harmonic of annual frequency to check for
# non-sinusoidal pattern
#
c2<-cos(4*pi*day/n)
s2<-sin(4*pi*day/n)</pre>
fit2<-lm(temperature~c1+s1+c2+s2)
lines(day,fit2$fitted.values,col="blue")
#
# two fits look similar, but conventional F test says otherwise
#
summary(fit2)
RSS1<-sum(fit1$resid^2); RSS2<-sum(fit2$resid^2)
```

```
1-pf(F,2,361)
```

F<-((RSS1-RSS2)/2)/(RSS2/361)

```
#
  conventional residual plots
#
#
    residuals vs fitted values
#
#
plot(fit2$fitted.values,fit2$resid)
#
    residuals in time-order as scatterplot
#
#
plot(1:366,fit2$resid)
#
    and as line-graph
#
#
plot(1:366,fit2$resid,type="l")
```

```
#
 examine autocorrelation properties of residuals
#
#
residuals<-fit2$resid
par(mfrow=c(2,2),pty="s")
for (k in 1:4) {
   plot(residuals[1:(n-k)],residuals[(k+1):n],
         pch=19,cex=0.5,xlab=" ",ylab=" ",main=k)
   }
par(mfrow=c(1,1))
acf(residuals)
#
# exponentially decaying correlation looks reasonable
#
cor(residuals[1:(n-1)],residuals[2:n])
Xmat<-cbind(rep(1,n),c1,s1,c2,s2)</pre>
rho<-0.01*(60:80)
profile<-AR1.profile(temperature,Xmat,rho)</pre>
```

```
#
# examine results
#
plot(rho,profile$logl,type="l",ylab="L(rho)")
Lmax<-max(profile$logl)
crit.val<-0.5*qchisq(0.95,1)
lines(c(rho[1],rho[length(rho)]),rep(Lmax-crit.val,2),lty=2)
profile
#</pre>
```

- # Exercise: how would you now re-assess the significance of
  # the second harmonic term?
- # the second harmonic term?

```
#
# profile log-likelihood function follows
#
AR1.profile<-function(y,X,rho) {</pre>
   m<-length(rho)</pre>
   logl<-rep(0,m)</pre>
   n<- length(y)</pre>
   hold < -outer(1:n, 1:n, "-")
   for (i in 1:m) {
      Rmat<-rho[i] ^abs(hold)</pre>
      ev<-eigen(Rmat)</pre>
      logdet<-sum(log(ev$values))</pre>
      Rinv<-ev$vectors%*%diag(1/ev$values)%*%t(ev$vectors)</pre>
      betahat <- solve(t(X)%*%Rinv%*%X)%*%t(X)%*%Rinv%*%y
      residual <- y-X%*%betahat
      logl[i]<- - logdet - n*log(c(residual)%*%Rinv%*%c(residual))</pre>
      }
   max.index<-order(logl)[m]</pre>
   Rmat<-rho[max.index]^abs(hold)</pre>
      ev<-eigen(Rmat)</pre>
      logdet<-sum(log(ev$values))</pre>
      Rinv<-ev$vectors%*%diag(1/ev$values)%*%t(ev$vectors)</pre>
      betahat <- solve(t(X)%*%Rinv%*%X)%*%t(X)%*%Rinv%*%y
      residual <- y-X%*%betahat
      sigmahat<-sqrt(c(residual)%*%Rinv%*%c(residual)/n)</pre>
   list(logl=logl,rhohat=rho[max.index],sigmahat=sigmahat,betahat=betahat)
   }
```

Longitudinal data

- replicated time series;
- focus of interest often on mean values;
- modelling and inference can and should exploit replication

**Discrete spatial variation** 

- space is not like time;
- models for discrete spatial variation are tied to number of spatial units

**Real-valued continuous spatial variation** 

- direct specification of covariance structure;
- variogram as an exploratory and/or diagnostic tool

**Spatial point processes** 

- the Poisson process;
- crude classification of processes/patterns as regular, completely random or aggregated

# 3. Longitudinal data

- linear Gaussian models;
- conditional and marginal models;
- missing values

## Correlation and why it matters

- different measurements on the same subject are typically correlated
- and this must be recognised in the inferential process.

#### Estimating the mean of a time series

$$Y_1, Y_2, ..., Y_t, ..., Y_n$$
  $Y_t \sim N(\mu, \sigma^2)$ 

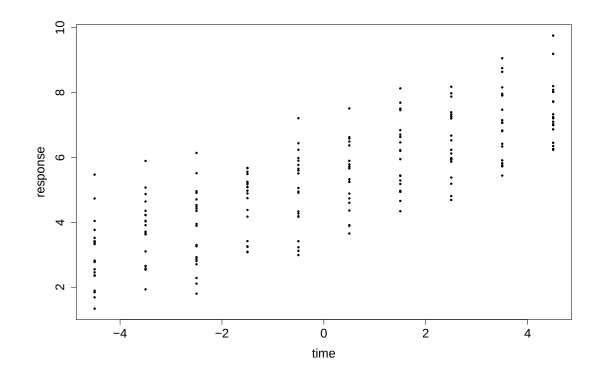
Classical result:  $ar{Y} \pm 2\sqrt{\sigma^2/n}$ 

But if  $Y_t$  is a time series:

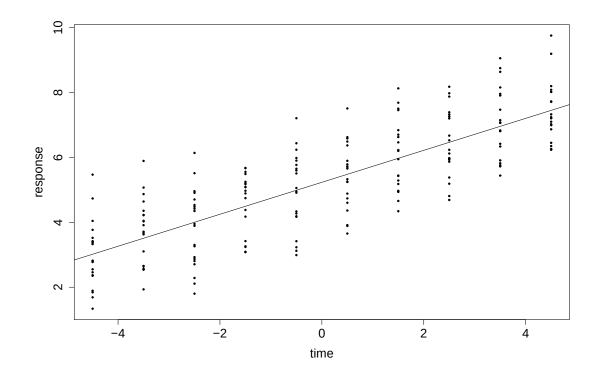
- $\mathrm{E}[\bar{Y}] = \mu$
- $\operatorname{Var}\{\bar{Y}\} = (\sigma^2/n) \times \{1 + n^{-1} \sum_{u \neq t} \operatorname{Corr}(Y_t, Y_u)\}$

Exercise: is the sample variance unbiased for  $\sigma^2 = Var(Y_t)$ ?

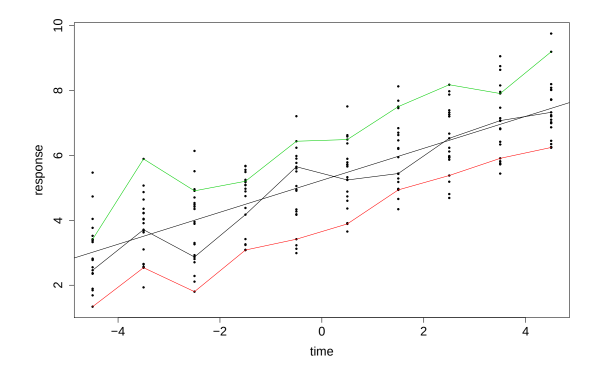
$$Y_{it}=lpha+eta(t-ar{t})+Z_{it} \qquad i=1,...,m \quad t=1,...,n$$



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$$Y_{it}=lpha+eta(t-ar{t})+Z_{it} \qquad i=1,...,m \quad t=1,...,n$$

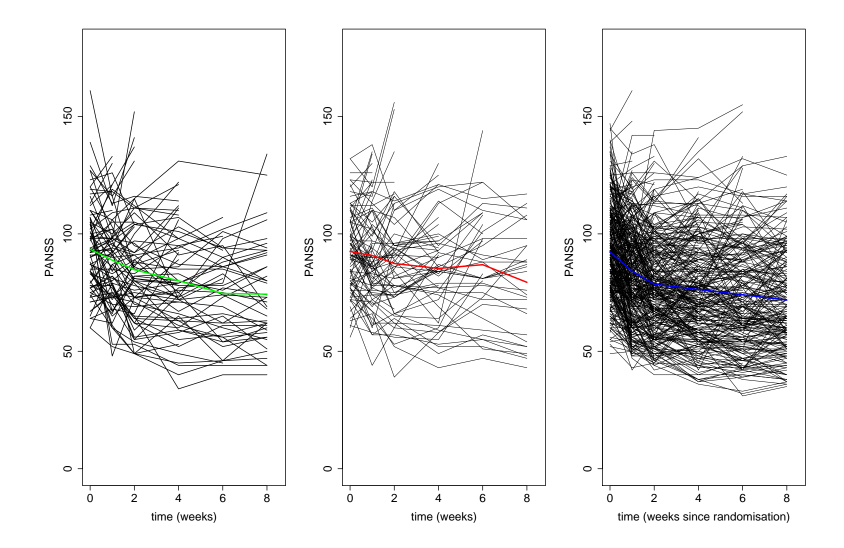


$$Y_{it}=lpha+eta(t-ar{t})+Z_{it} \qquad i=1,...,m \quad t=1,...,n$$

Parameter estimates and standard errors:

	ignoring correlation		recognising correlation		
	estimate	standard error	estimate	standard error	
$\alpha$	5.234	0.074	5.234	0.202	
$oldsymbol{eta}$	0.493	0.026	0.493	0.011	

## A spaghetti plot of the PANSS data



# Exploring covariance structure: balanced data

$$(Y_{ij}, t_j): j = 1, ..., n; i = 1, ..., m$$

- fit saturated treatments-by-times model to mean response
- compute sample covariance matrix of residuals

PANSS data:

SDY.t0Y.t1Y.t2Y.t4Y.t6Y.t8Y.t020.0191.0000.5330.3660.4480.2850.229Y.t120.1840.5331.0000.6930.5890.6580.535Y.t222.1200.3660.6931.0000.6700.5670.678Y.t420.9960.4480.5890.6701.0000.7180.648Y.t624.7460.2850.6580.5670.7181.0000.792Y.t823.6660.2290.5350.6780.6480.7921.000

- modest increase in variability over time
- correlation decays with increasing time-separation

# Exploring covariance structure: unbalanced data

$$(Y_{ij},t_{ij}): j=1,...,n_i; i=1,...,m$$

The variogram of a stochastic process Y(t) is

$$V(u) = \frac{1}{2} \operatorname{Var} \{ Y(t) - Y(t-u) \}$$

- well-defined for stationary and some non-stationary processes
- for stationary processes,

$$V(u) = \sigma^2 \{1 - \rho(u)\}$$

• V(u) easier to estimate than  $\rho(u)$  when data are unbalanced

#### Estimating the variogram

Data:  $(Y_{ij}, t_{ij}) : i = 1, ..., m; j = 1, ..., n_i$ 

 $r_{ij}$  = residual from preliminary model for mean response

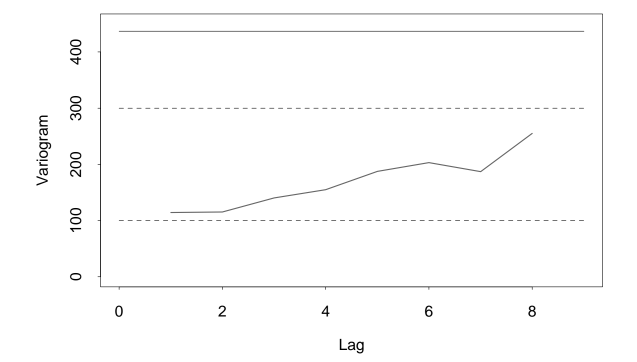
• Define

$$v_{ijk\ell}=rac{1}{2}(r_{ij}-r_{k\ell})^2$$

• Estimate

 $\hat{V}(u) = ext{average of all } v_{iji\ell} ext{ such that } |t_{ij} - t_{i\ell}| \simeq u$  $\hat{\sigma}^2 = ext{average of all } v_{ijk\ell} ext{ such that } i \neq k.$  **Example:** sample variogram of the PANSS data

Solid lines are estimates from data, horizontal lines are eye-ball estimates (explanation later)



## Where does the correlation come from?

- differences between subjects
- variation over time within subjects
- measurement error

#### General linear model, correlated residuals

i =subjects j = measurements within subjects

$$egin{array}{rcl} E(Y_{ij})&=&x_{ij1}eta_1+...+x_{ijp}eta_p\ &Y_i&=&X_ieta+\epsilon_i\ &Y&=&Xeta+\epsilon \end{array}$$

- measurements from different subjects independent
- measurements from same subject typically correlated.

# Parametric models for covariance structure

Three sources of random variation in a typical set of longitudinal data:

- Random effects (variation between subjects)
  - characteristics of individual subjects
  - for example, intrinsically high or low responders
  - influence extends to all measurements on the subject in question.

# Parametric models for covariance structure

Three sources of random variation in a typical set of longitudinal data:

- Random effects
- Serial correlation (variation over time within subjects)
  - measurements taken close together in time typically more strongly correlated than those taken further apart in time
  - on a sufficiently small time-scale, this kind of structure is almost inevitable

# Parametric models for covariance structure

Three sources of random variation in a typical set of longitudinal data:

- Random effects
- Serial correlation
- Measurement error
  - when measurements involve delicate determinations, duplicate measurements at same time on same subject may show substantial variation

Diggle, Heagerty, Liang and Zeger (2002, Chapter 5)

#### Some simple models

• Compound symmetry

$$egin{aligned} Y_{ij} - \mu_{ij} &= U_i + Z_{ij} \ U_i &\sim \mathrm{N}(0, 
u^2) \ Z_{ij} &\sim \mathrm{N}(0, au^2) \end{aligned}$$

Implies that  $\operatorname{Corr}(Y_{ij}, Y_{ik}) = \nu^2/(\nu^2 + \tau^2)$ , for all  $j \neq k$ 

• Random intercept and slope

$$Y_{ij}-\mu_{ij}=U_i+W_it_{ij}+Z_{ij}$$
 $(U_i,W_i)\sim \mathrm{BVN}(0,\Sigma)$  $Z_{ij}\sim \mathrm{N}(0, au^2)$ 

Often fits short sequences well, but extrapolation dubious, for example  $Var(Y_{ij})$  quadratic in  $t_{ij}$ 

#### • Autoregressive

$$egin{aligned} Y_{ij} - \mu_{ij} &= lpha(Y_{i,j-1} - \mu_{i,j-1}) + Z_{ij} \ Y_{i1} - \mu_{i1} &\sim \mathrm{N}\{0, au^2/(1-lpha^2)\} \ Z_{ij} &\sim \mathrm{N}(0, au^2), \quad j = 2, 3, ... \end{aligned}$$

# Not a natural choice for underlying continuous-time processes

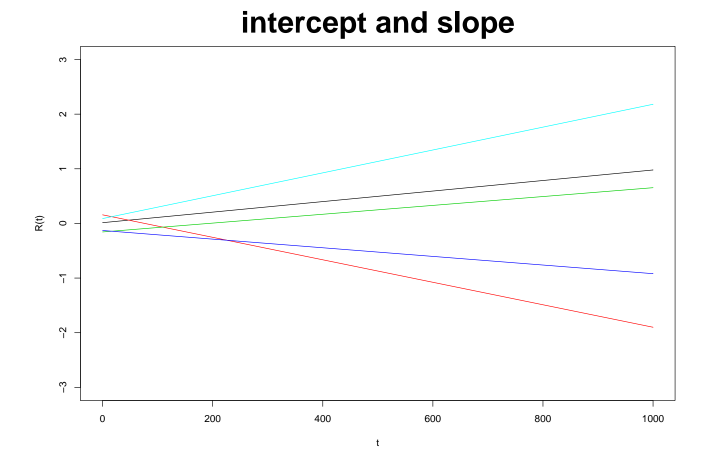
• Stationary Gaussian process

$$Y_{ij}-\mu_{ij}=W_i(t_{ij})$$

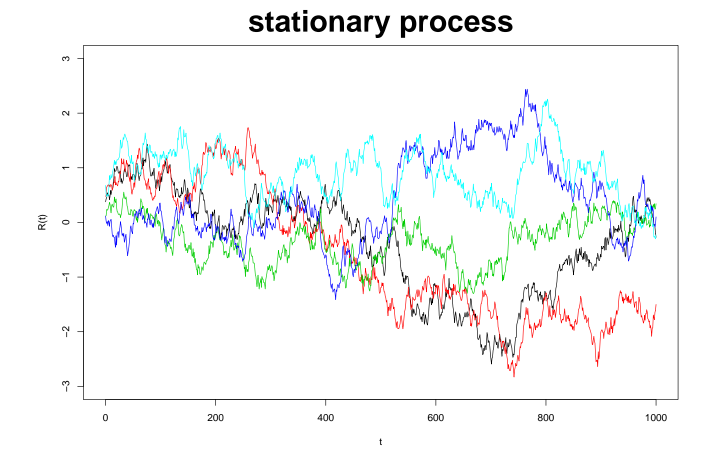
 $W_i(t)$  a continuous-time Gaussian process $\mathrm{E}[W(t)]=0\quad\mathrm{Var}\{W(t)\}=\sigma^2 \ \mathrm{Corr}\{W(t),W(t-u)\}=
ho(u)$ 

 $\rho(u) = \exp(-u/\phi)$  gives continuous-time version of the autoregressive model

# **Time-varying random effects**



# **Time-varying random effects: continued**



#### • A general model

$$Y_{ij}-\mu_{ij}=d_{ij}^{\prime}U_i+W_i(t_{ij})+Z_{ij}$$

 $U_i \sim \mathrm{MVN}(0, \Sigma) \ (\mathrm{random \ effects})$ 

 $d_{ij} =$ vector of explanatory variables for random effects

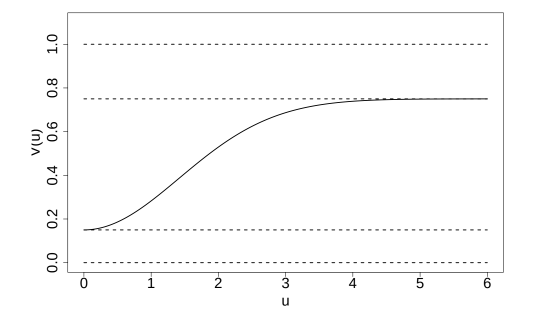
 $W_i(t) =$ continuous-time Gaussian process (serial correlation)

 $Z_{ij} \sim {
m N}(0, au^2) \ ({
m measurement\ errors})$ 

Even when all three components of variation are needed in principle, one or two may dominate in practice The variogram of the general model (stationary case)

 $Y_{ij}-\mu_{ij}=U_i+W_i(t_{ij})+Z_{ij}$ 

 $V(u) = \tau^{2} + \sigma^{2} \{ 1 - \rho(u) \} \quad \text{Var}(Y_{ij}) = \nu^{2} + \sigma^{2} + \tau^{2}$ 



## Fitting the model: non-technical summary

- Ad hoc methods won't do
- Likelihood-based inference is the statistical gold standard
- But be sure you know what you are estimating when there are missing values

## Maximum likelihood estimation ( $V_0$ known)

Log-likelihood for observed data y is

$$L(\beta, \sigma^{2}, V_{0}) = -0.5\{nm \log \sigma^{2} + m \log |V_{0}| + \sigma^{-2}(y - X\beta)'(I \otimes V_{0})^{-1}(y - X\beta)\}, (1)$$

 $I\otimes V_0$  denotes block-diagonal matrix with non-zero blocks  $V_0$ Given  $V_0$ , estimator for  $\beta$  is

$$\hat{\beta}(V_0) = (X'(I \otimes V_0)^{-1} X)^{-1} X'(I \otimes V_0)^{-1} y, \qquad (2)$$

Explicit estimator for  $\sigma^2$  also available as

$$\hat{\sigma}^2(V_0) = RSS(V_0)/(nm) \tag{3}$$

 $RSS(V_0) = \{y - X \hat{eta}(V_0)\}' (I \otimes V_0)^{-1} \{y - X \hat{eta}(V_0)\}.$ 

#### Maximum likelihood estimation, $V_0$ unknown

Substitute (2) and (3) into (1) to give reduced log-likelihood

$$\mathcal{L}(V_0) = -0.5m[n\log\{RSS(V_0)\} + \log|V_0|].$$
(4)

Numerical maximization of (4) then gives  $\hat{V}_0$ , hence  $\hat{\beta} = \hat{\beta}(\hat{V}_0)$ and  $\hat{\sigma}^2 = \hat{\sigma}^2(\hat{V}_0)$ .

- Dimensionality of optimisation is  $\frac{1}{2}n(n+1) 1$
- Each evaluation of  $\mathcal{L}(V_0)$  requires inverse and determinant of an n by n matrix.

# A random effects model for CD4 cell counts

```
data<-read.table("CD4.data",header=T)
data[1:3,]
time<-data$time
CD4<-data$CD4
plot(time,CD4,pch=19,cex=0.25)
id<-data$id
uid<-unique(id)
for (i in 1:10) {
   take<-(id==uid[i])
   lines(time[take],CD4[take],col=i,lwd=2)
   }</pre>
```

```
# Simple linear model assuming uncorrelated residuals
#
fit1<-lm(CD4~time)
summary(fit1)
#
# random intercept and slope model
#
library(nlme)
?lme
fit2<-lme(CD4~time,random=~1|id)
summary(fit2)</pre>
```

```
# make fitted value constant before sero-conversion
#
timeplus<-time*(time>0)
fit3<-lme(CD4~timeplus,random=~1|id)</pre>
summary(fit3)
tfit<-0.1*(0:50)
Xfit<-cbind(rep(1,51),tfit)</pre>
fit<-c(Xfit%*%fit3$coef$fixed)</pre>
Vmat<-fit3$varFix
Vfit<-diag(Xfit%*%Vmat%*%t(Xfit))</pre>
upper<-fit+2*sqrt(Vfit)</pre>
lower<-fit-2*sqrt(Vfit)</pre>
#
# plot fit with 95% point-wise confidence intervals
#
plot(time,CD4,pch=19,cex=0.25)
lines(c(-3,tfit),c(upper[1],upper),col="red")
lines(c(-3,tfit),c(lower[1],lower),col="red")
```

# Missing values and dropouts

Issues concerning missing values in longitudinal data can be addressed at two different levels:

- technical: can the statistical method I am using cope with missing values?
- conceptual: *why* are the data missing? Does the fact that an observation is missing convey partial information about the value that would have been observed?

These same questions also arise with cross-sectional data, but the correlation structure of longitudinal data can sometimes be exploited to good effect, by modelling how the probability of dropout for each person depends on their previously observed measurements

### **Rubin's classification**

- MCAR (completely at random): P(missing) depends neither on observed nor unobserved measurements
- MAR (at random): P(missing) depends on observed measurements, but not on unobserved measurements
- MNAR (not at random): conditional on observed measurements, P(missing) depends on unobserved measurements.

Rubin (1976)

# Dropout

Once a subject goes missing, they never return

**Example : Longitudinal clinical trial** 

- completely at random: patient leaves the the study because they move house
- at random : patient leaves the study on their doctor's advice, based on observed measurement history
- not at random : patient misses their appointment because they are feeling unwell.

Little (1995)

# **Conventional wisdom**

- any sensible method of analysis valid if dropout is MCAR
- likelihood-based analysis valid if dropout is MAR

**But:** under MAR, target of likelihood-based inference is model for hypothetical dropout-free population **Proof:** Partition Y for each subject into observed and missing components,  $Y = (Y_o, Y_m)$  and let M denote binary vector of missingness indicators. Likelihood for observed data is

$$egin{array}{rcl} L = & g(y_o,m) = & \int f(y_o,y_m,m) dy_m \ &= & \int f(y_o) f(y_m|y_o) p(m|y_o,y_m) dy_m \end{array}$$

If  $p(m|y_o, y_m) = p(m|y_o)$ , take outside integral to give

 $L = p(m|y_o)f(y_o)$ 

and log-likelihood contribution

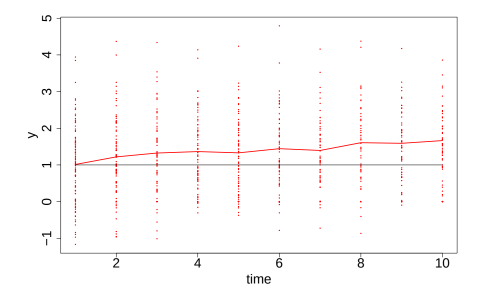
$$\log L = \log p(m|y_o; \theta) + \log f(y_o|\theta)$$

- OK to ignore first term for likelihood inference about  $\theta$
- and no loss of efficiency if  $\theta = (\theta_1, \theta_2)$  such that  $\theta_1$  and  $\theta_2$  parameterise  $p(\cdot)$  and  $f(\cdot)$ , respectively.

But is inference about  $f(\cdot)$  what you want?

#### Example

- Model is  $Y_{ij} = \mu + U_i + Z_{ij}$  (random intercept)
- Dropout is MAR:  $logit(p_{ij}) = -1 2 \times Y_{i,j-1}$



• Observed means increase over time, but population mean  $\mu$  is constant

# PJD's take on ignorability

For correlated data, dropout mechanism can be ignored only if dropouts are completely random

In all other cases, need to:

- think carefully what are the relevant practical questions,
- fit an appropriate model for both measurement process and dropout process
- use the model to answer the relevant questions.

Diggle, Farewell and Henderson (2007)

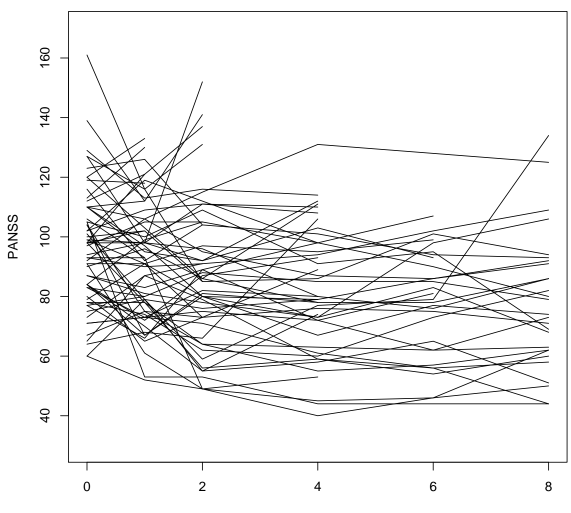
# Schizophrenia trial data

- Data from placebo-controlled RCT of drug treatments for schizophrenia:
  - Placebo; Haloperidol (standard); Risperidone (novel)
- Y = sequence of weekly PANSS measurements
- F = dropout time
- Total m = 516 subjects, but high dropout rates:

week	-1	0	1	2	4	6	8
missing	0	3	9	70	122	<b>205</b>	<b>251</b>
proportion	0.00	0.01	0.02	0.14	0.24	0.40	0.49

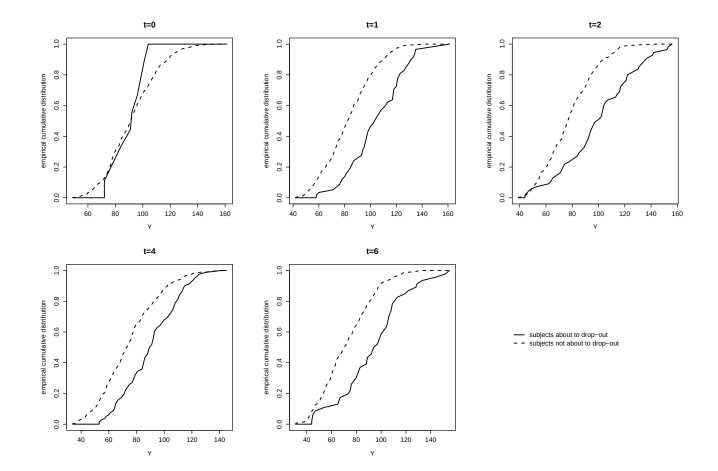
• Dropout rate also treatment-dependent (P > H > R)

#### Schizophrenia data PANSS responses from haloperidol arm

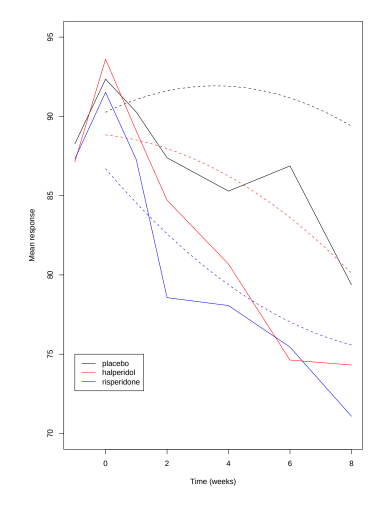


time (weeks)

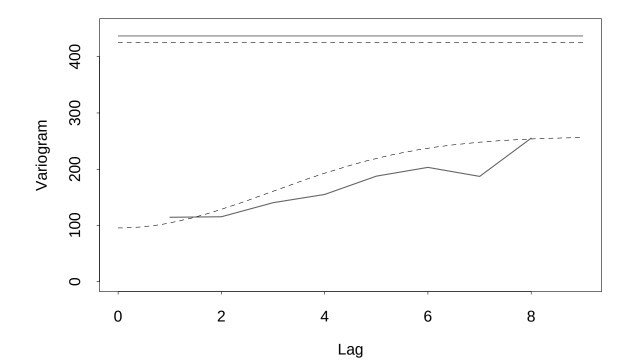
#### Dropout is not completely at random



#### Schizophrenia trial data Mean response (random effects model)



#### Schizophrenia trial data Empirical and fitted variograms



# Schizophrenia trial data: summary

- dropout is not MCAR
- MAR model apparently fits well, but:
  - hard to distinguish empirically between different MAR models;
  - and we haven't formally investigated evidence for informative dropout
- Fitted means relate to hypothetical, dropout-free population

**Exercise:** think about how you might embed the MAR model within an informative dropout model

# Generalized linear models for longitudinal data

- random effects models
- transition models
- marginal models

Diggle, Heagerty, Liang and Zeger (2002, Chapter 7)

#### Random effects GLM

Responses  $Y_1, \ldots, Y_n$  on an individual subject conditionally independent, given unobserved vector of random effects U

 $U \sim g(u)$  represents properties of individual subjects that vary randomly between subjects

- $E(Y_j|U) = \mu_j : h(\mu_i) = \mathbf{x}'_j \beta + U' \alpha$
- $\operatorname{Var}(Y_j|U) = \phi v(\mu_j)$
- $(Y_1, \ldots, Y_n)$  are mutually independent conditional on U.

Likelihood inference requires evaluation of

$$f(y) = \int \prod_{j=1}^n f(y_j|U)g(U)dU$$

## **Transition GLM**

Each  $Y_j$  modelled conditionally on preceding  $Y_1, Y_2, \ldots, Y_{j-1}$ .

- $E(Y_j|\text{history}) = \mu_j$
- $h(\mu_j) = \mathbf{x}'_j \beta + \sum_{k=1}^{j-1} Y'_{j-k} \alpha_k$
- $\operatorname{Var}(Y_j|\operatorname{history}) = \phi v(\mu_j)$

Construct likelihood as product of conditional distributions, usually assuming restricted form of dependence.

Example:  $f_k(y_j|y_1, ..., y_{j-1}) = f_k(y_j|y_{j-1})$ 

Need to condition on  $y_1$  as model does not directly specify marginal distribution  $f_1(y_1)$ .

# Marginal GLM

Let  $h(\cdot)$  be a link function which operates component-wise,

- $E(y) = \mu : h(\mu) = X\beta$
- $\operatorname{Var}(y_i) = \phi v(\mu_i)$
- $\operatorname{Corr}(y) = R(\alpha)$ .

Not a fully specified probability model

May require constraints on variance function  $v(\cdot)$  and correlation matrix  $R(\cdot)$  for valid specification

Inference for  $\beta$  uses generalized estimating equations

Liang and Zeger (1986)

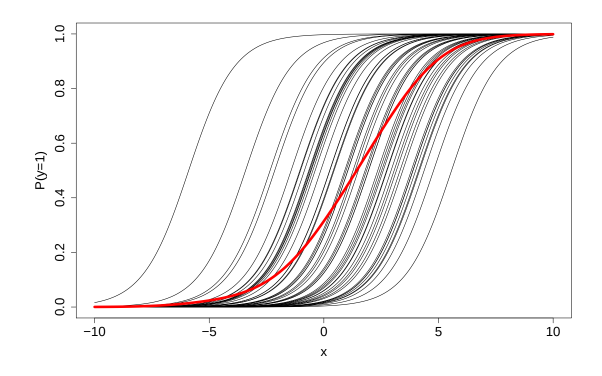
#### What are we estimating?

- in marginal modelling,  $\beta$  measures population-averaged effects of explanatory variables on mean response
- in transition or random effects modelling,  $\beta$  measures effects of explanatory variables on mean response of an individual subject, conditional on
  - subject's measurement history (transition model)
  - subject's own random characteristics  $U_i$  (random effects model)

Example: Simulation of a logistic regression model, probability of positive response from subject i at time t is  $p_i(t)$ ,

 $logit{p_i(t)}: \alpha + \beta x(t) + \gamma U_i,$ 

x(t) is a continuous covariate and  $U_i$  is a random effect



Example: Effect of mother's smoking on probability of intra-uterine growth retardation (IUGR).

Consider a binary response Y = 1/0 to indicate whether a baby experiences IUGR, and a covariate x to measure the mother's amount of smoking.

Two relevant questions:

- 1. public health: by how much might population incidence of IUGR be reduced by a reduction in smoking?
- 2. clinical/biomedical: by how much is a baby's risk of IUGR reduced by a reduction in their mother's smoking?

Question 1 is addressed by a marginal model, question 2 by a random effects model

## **R** software

The following is almost certainly an incomplete list.

• Marginal models

Function gee within package of same name

• Random effects models

Function glmmPQL within MASS package or lmer within lme4 (but note evaluation of likelihood uses approximate methods that may perform badly if random effects are highdimensional). Package glmmBUGS is a Bayesian alternative.

• Transition models

Standard glm function, after computing values of required functions of lagged responses to be used as explanatory variables.

# 4. Continuous spatial variation

- stationary Gaussian processes;
- variogram estimation;
- likelihood-based estimation;
- spatial prediction.

What is this thing called geostatistics?

## biostatistics = bio-statistics

## geostatistics $\neq$ geo-statistics

The core geostatistical problem: given a set of measured values  $Y_i$  at locations  $x_i \in A$  of some spatial phenomenon  $S(\cdot)$ , what can you say about the complete surface  $\{S(x) : x \in A\}$ ?

Krige, 1951; Matérn, 1960; Mathéron, 1963; Watson, 1972; Ripley, 1981

### **Recall from LDA lectures**

- Stationary Gaussian process  $Y_{ij} \mu_{ij} = W_i(t_{ij})$  $W_i(t)$  a continuous-time Gaussian process  $E[W(t)] = 0, Var\{W(t)\} = \sigma^2,$  $Corr\{W(t), W(t-u)\} = \rho(u)$
- Variogram of a stochastic process Y(t) is

$$V(u)=rac{1}{2}\mathrm{Var}\{Y(t)-Y(t-u)\}$$

For stationary processes,

$$V(u)=\sigma^2\{1-
ho(u)\}$$

For geostatistics, simply substitute a spatial process S(x) for the temporal process W(t), and off you go

# **Model-based Geostatistics**

- the application of general principles of statistical modelling and inference to geostatistical problems
- Example: kriging as minimum mean square error prediction under Gaussian modelling assumptions Diggle, Moyeed and Tawn, 1998; Diggle and Ribeiro, 2007
- An important practical difference between LDA and geostatistics is its typical scientific focus:
  - LDA: estimating (mean response profiles) from multiple realisations
  - geostatistics: predicting (a spatially continuous phenomenon) from a single, incomplete realisation

# Computation with geoR

```
library(geoR)
lead<-read.table("lead2000.data",header=T)
lead<-as.geodata(lead)
summary(lead)
plot(lead)
?points.geodata
points(lead,cex.min=1,cex.max=4)
points(lead,cex.min=0.5,cex.max=2)
points(lead,cex.min=0.5,cex.max=2,pt.div="quint")
loglead<-lead
loglead$data<-log(loglead$data)
points(loglead,cex.min=0.5,cex.max=2,pt.div="quint")
```

## Notation

- $Y = \{Y_i : i = 1, ..., n\}$  is the measurement data
- $\{x_i: i = 1, ..., n\}$  is the sampling design
- A is the region of interest
- $S^* = \{S(x) : x \in A\}$  is the signal process

• 
$$S = \{S(x_i) : i = 1, ..., n\}$$

- $T = \mathcal{F}(S^*)$  is the target for prediction
- $[S^*, Y] = [S^*][Y|S^*]$  is the geostatistical model

Typically,  $[S^*, Y]$  can be further factorised and simplified as

$$[S^{\ast},Y] = [S][S^{\ast}|S][Y|S^{\ast},S] = [S][S^{\ast}|S][Y|S]$$

Exercise: why is this helpful?

## Gaussian model-based geostatistics

Model specification:

- Stationary Gaussian process  $S(x): x \in \mathbb{R}^2$ 
  - $\cdot \operatorname{E}[S(x)] = \mu$
  - $\cdot \operatorname{Cov}\{S(x),S(x')\} = \sigma^2 \rho(\|x-x'\|)$
- Mutually independent  $Y_i | S(\cdot) \sim \mathcal{N}(S(x), \tau^2)$

### Minimum mean square error prediction

$$[S^*,Y]=[Y][S^*|Y] \qquad \quad T=\mathcal{F}(S^*)$$

- $\hat{T} = t(Y)$  is a point predictor
- $MSE(\hat{T}) = E[(\hat{T} T)^2]$

Theorem:  $MSE(\hat{T})$  takes its minimum value when  $\hat{T} = E(T|Y)$ . Proof uses result that for any predictor  $\tilde{T}$ ,

$$\mathbf{E}[(T-\tilde{T})^2] = \mathbf{E}_{Y}[\operatorname{Var}_{T}(T|Y)] + \mathbf{E}_{Y}\{[\mathbf{E}_{T}(T|Y) - \tilde{T}]^2\}$$

Immediate corollary is that

$$\operatorname{E}[(T - \hat{T})^2] = \operatorname{E}_Y[\operatorname{Var}(T|Y)] \approx \operatorname{Var}(T|Y)$$

Simple and ordinary kriging

**Recall Gaussian model:** 

- Stationary Gaussian process  $S(x): x \in \mathbb{R}^2$ 
  - $\cdot \operatorname{E}[S(x)] = \mu$
  - $\cdot \operatorname{Cov}\{S(x),S(x')\} = \sigma^2 \rho(\|x-x'\|)$
- Mutually independent  $Y_i | S(\cdot) \sim \mathcal{N}(S(x), \tau^2)$

Gaussian model implies

$$Y \sim \mathrm{MVN}(\mu 1, \sigma^2 V)$$

$$V = R + (\tau^2 / \sigma^2) I$$
  $R_{ij} = \rho(||x_i - x_j||)$ 

Suppose target for prediction is T = S(x) for arbitrary x, write  $r = (r_1, ..., r_n)$  where

$$r_i = \rho(\|x - x_i\|)$$

Standard results on multivariate Normal then give [T|Y] as multivariate Gaussian with mean and variance

$$\hat{T} = \mu + r' V^{-1} (Y - \mu 1)$$
(5)

$$\operatorname{Var}(T|Y) = \sigma^2 (1 - r'V^{-1}r).$$
 (6)

Simple kriging:  $\hat{\mu} = \bar{Y}$  Ordinary kriging:  $\hat{\mu} = (1'V^{-1}1)^{-1}1'V^{-1}Y$ 

### The Matérn family of correlation functions

$$ho(u)=2^{\kappa-1}(u/\phi)^\kappa K_\kappa(u/\phi)$$

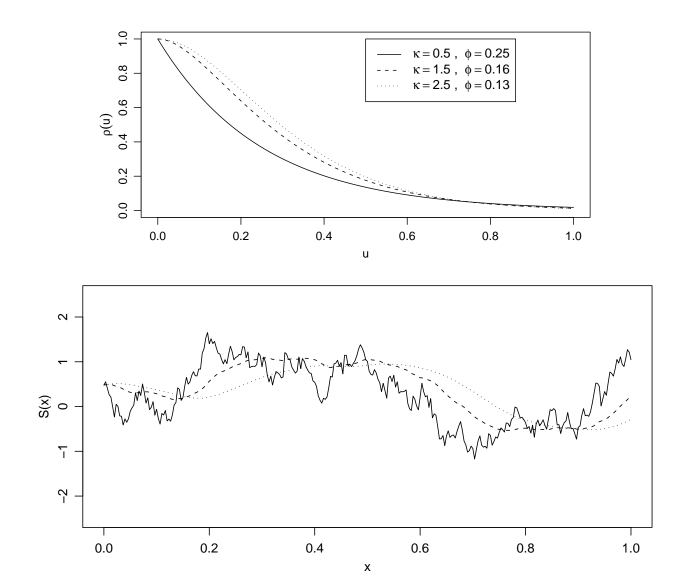
- parameters  $\kappa > 0$  and  $\phi > 0$
- $K_{\kappa}(\cdot)$  : modified Bessel function of order  $\kappa$

• 
$$\kappa = 0.5$$
 gives  $ho(u) = \exp\{-u/\phi\}$ 

• 
$$\kappa \to \infty$$
 gives  $\rho(u) = \exp\{-(u/\phi)^2\}$ 

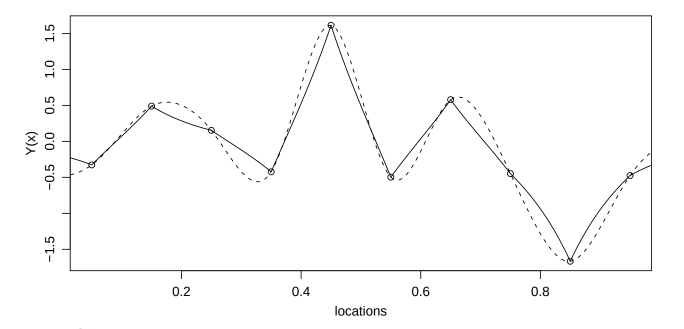
- $\kappa$  and  $\phi$  are not orthogonal:
  - more nearly orthogonal re-parametrisation to  $\alpha = 2\phi \sqrt{\kappa}$
  - estimation of  $\kappa$  is difficult
  - but helpful interpretation: S(x) is k times meansquare differentiable if  $\kappa > k$

## Matérn correlation functions: varying $\kappa$



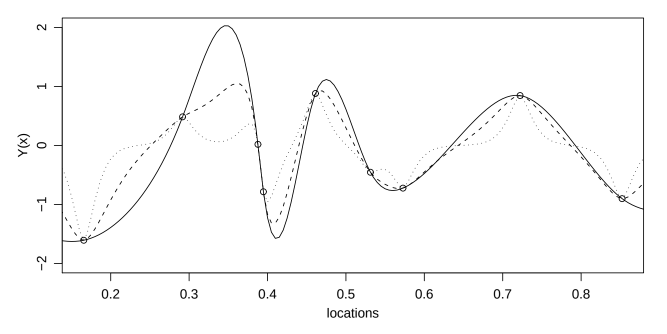
## Simple kriging: three examples

1. Varying  $\kappa$  (smoothness of S(x))



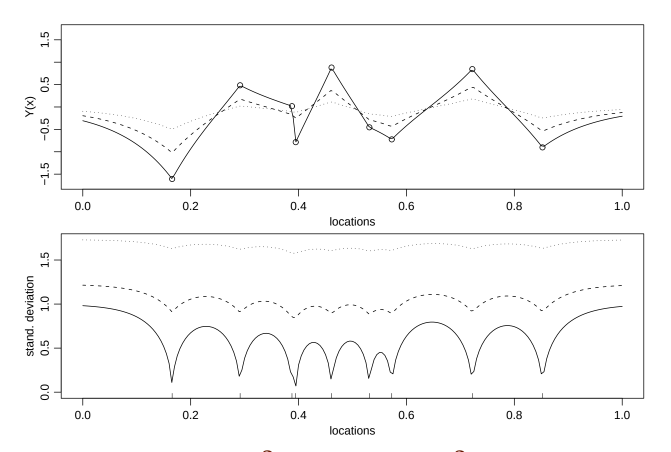
Assuming  $\tau^2 = 0$  (no measurement error), kriging predictors interpolate the data

#### 2. Varying $\phi$ (range of spatial correlation)



For  $\tau^2 = 0$  and fixed  $\kappa$ , all kriging predictors have the same analytic smoothness, but the smaller the value of  $\phi$ , the more quickly the predictions revert to the assumed constant mean,  $\mu$ , at non-data locations

#### 3. Varying $\tau^2/\sigma^2$ (noise-to-signal ratio)



The smaller the value of  $\tau^2$ , for fixed  $/\sigma^2$ , the more closely the predictions approach the data at the data-locations, and the smaller are the prediction variances.

## **Predicting non-linear functionals**

- minimum mean square error prediction is not invariant under non-linear transformation
- the complete answer to a prediction problem is the predictive distribution, [T|Y]
- Recommended strategy:
  - draw repeated samples from  $[S^*|Y]$ (conditional simulation)
  - calculate corresponding values of  $T = t(S^*)$ from each sample (examples to follow)

### The variogram re-visited

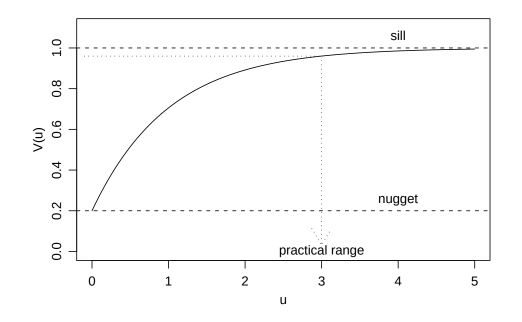
• the variogram of a process Y(x) is the function

$$V(x,x') = \frac{1}{2} \operatorname{Var} \{Y(x) - Y(x')\}$$

• for the spatial Gaussian model, with u = ||x - x'||,

$$V(u) = \tau^2 + \sigma^2 \{1 - \rho(u)\}$$

• provides a summary of the basic structural parameters of the spatial Gaussian process



- the nugget variance:  $\tau^2$
- the sill:  $\sigma^2 = \operatorname{Var}\{S(x)\}$
- the practical range: directly related to  $\phi$ ; specifically, the value of u such that

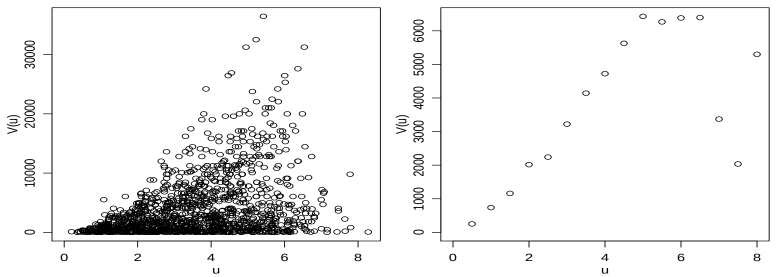
$$ho(u)=
ho_0(u/\phi)=0.05$$

## **Empirical variograms**

$$u_{ij} = \|x_i - x_j\|$$
  $v_{ij} = 0.5[y(x_i) - y(x_j)]^2$ 

- the variogram cloud is a scatterplot of the points  $(u_{ij}, v_{ij})$
- the empirical variogram smooths the variogram cloud by averaging within bins:  $u h/2 \le u_{ij} < u + h/2$
- for a process with non-constant mean (covariates), use residuals  $r(x_i) = y(x_i) \hat{\mu}(x_i)$  to compute  $v_{ij}$

### Limitations of $\hat{V}(u)$



- 1.  $v_{ij} \sim V(u_{ij}) \chi_1^2$
- 2. the  $v_{ij}$  are correlated

#### **Consequences:**

- variogram cloud is unstable, pointwise and in overall shape
- binning addresses point 1, but not point 2

Parameter estimation using the variogram

What not to do and how to do it

• weighted least squares criterion:

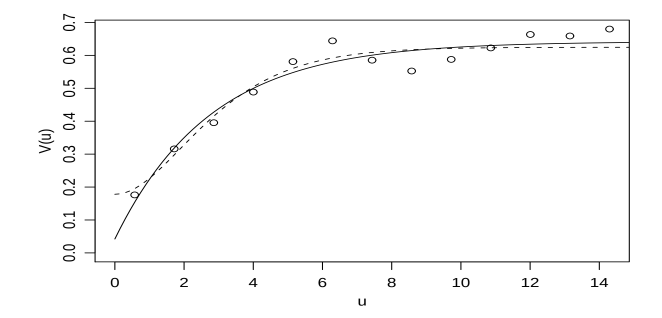
$$W( heta) = \sum_k n_k \{ar{V}_k - V(u_k; heta)\}^2$$

where  $\theta$  denotes vector of covariance parameters and  $\bar{V}_k$ is average of  $n_k$  variogram ordinates  $v_{ij}$ .

- need to choose upper limit for u (arbitrary?)
- variations include:
  - fitting models to the variogram cloud
  - other estimators for the empirical variogram
  - different proposals for weights

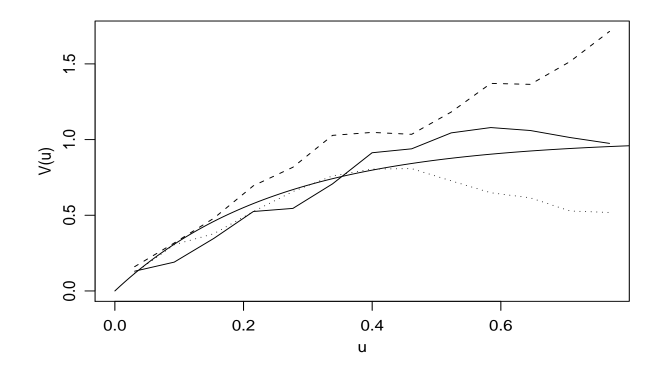
**Comments on variogram fitting** 

1. Can give equally good fits for different extrapolations at origin.

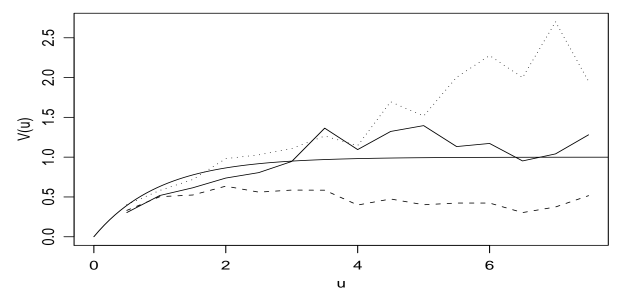


2. Correlation between variogram points induces smoothness.

Empirical variograms for three simulations from the same model.



- 3. Fit is sensitive to specification of the mean. Illustration with linear trend surface:
  - solid smooth line: theoretical variogram;
  - dotted line: from data;
  - solid line: from true residuals;
  - dashed line : from estimated residuals.



Note: no analogue of saturated model used in LDA to counteract bias

### Parameter estimation: maximum likelihood

$$Y \sim \mathrm{MVN}(\mu 1, \sigma^2 R + \tau^2 I)$$

R is the  $n \times n$  matrix with  $(i, j)^{th}$  element  $\rho(u_{ij})$  where  $u_{ij} = ||x_i - x_j||$ , Euclidean distance between  $x_i$  and  $x_j$ . Or more generally:

$$\mu(x_i) = \sum_{j=1}^k f_k(x_i)eta_k$$

where  $d_k(x_i)$  is a vector of covariates at location  $x_i$ , hence

 $Y \sim \mathrm{MVN}(Deta, \sigma^2 R + au^2 I)$ 

Gaussian log-likelihood function:

$$egin{aligned} L(eta, au,\sigma,\phi,\kappa) \propto & -0.5\{\log|(\sigma^2R+ au^2I)|+\ & (y-Deta)'(\sigma^2R+ au^2I)^{-1}(y-Deta)\}. \end{aligned}$$

• write  $\nu^2 = \tau^2/\sigma^2$ , hence  $\sigma^2 V = \sigma^2 (R + \nu^2 I)$ 

• log-likelihood function is maximised for

$$\hat{\beta}(V) = (D'V^{-1}D)^{-1}D'V^{-1}y$$
$$\hat{\sigma}^2 = n^{-1}(y - D\hat{\beta})'V^{-1}(y - D\hat{\beta})$$

• substitute  $(\hat{\beta}, \sigma^2)$  to give reduced maximisation problem

$$L^*(
u^2, \phi, \kappa) \propto -0.5\{n \log |\hat{\sigma^2}| + \log |(R + 
u^2 I)|\}$$

• usually just consider  $\kappa$  in a discrete set, for example  $\{0.5, 1.5, 2.5\}$  corresponding to continuous, differentiable and twice differentiable

# Comments on maximum likelihood

- likelihood-based methods preferable to variogram-based methods
- restricted maximum likelihood is widely recommended but in PJD's experience is sensitive to mis-specification of the mean model.
- in spatial models, distinction between  $\mu(x)$  and S(x) is not sharp.
- composite likelihood treats contributions from pairs  $(Y_i, Y_j)$ as if independent
- examining profile likelihoods is advisable, to check for poorly identified parameters
- for large data-sets it can be useful to partition the studyregion and compare fitted models in different sub-regions

A word on asymptotics

Two different asymptotic regimes are:

- increasing domain
- infill

Inferential implications are:

- increasing domain  $\Rightarrow$  consistent parameter estimation
- infill  $\Rightarrow$  consistent prediction

Stein, 1999

# Computation with geoR

```
loglead2<-loglead
loglead2$coords<-loglead$coords/100000
mlfit<-likfit(loglead2,ini.cov.pars=c(0.25,1),
    cov.model="matern",kappa=0.5)
region<-matrix(c(4.5,46.0,7.0,46.0,7.0,48.5,4.5,48.5),4,2,T)
grid<-pred_grid(region,by=0.1)
KC<-krige.control(obj.model=mlfit)
OC<-output.control(n.predictive=100)
set.seed(24367)
predictions<-krige.conv(geodata=loglead2,locations=grid,
    borders=region,krige=KC,output=0C)
```

```
image(predictions)
points(loglead2,add=T)
coast<-read.table("galicia_coastline.txt",header=T)
lines(coast[,1],coast[,2])
par(mfrow=c(1,2))
hist(loglead2$data,main="data")
predict.max<-NULL
for (sim in 1:100) {
    predict.max<-c(predict.max,max(predictions$simulations[,sim]))
    }
hist(predict.max,main="predicted maximum")</pre>
```

### **Trans-Gaussian models**

- assume Gaussian model holds after point-wise transformation
- Box-Cox family is widely used

$$Y_i^* = h_\lambda(Y_i) = \left\{egin{array}{cc} (Y_i^\lambda - 1)/\lambda & ext{if } \lambda 
eq 0 \ \log(Y_i) & ext{if } \lambda = 0 \end{array}
ight.$$

• bias-correction? only if point prediction is required?

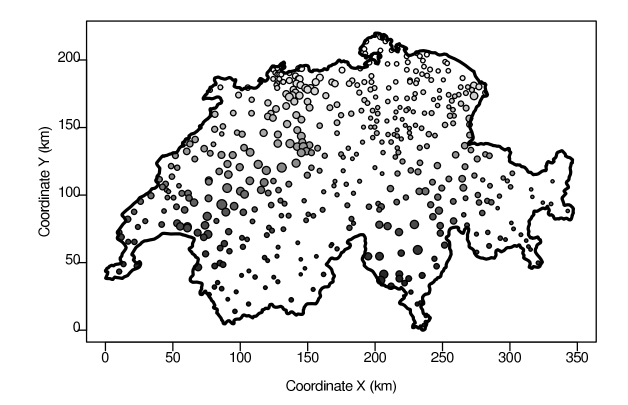
**Example:** log-Gaussian kriging

- $T(x) = \exp\{S(x)\}$   $\hat{T}(x) = \exp\{\hat{S}(x) + v(x)/2\}$
- $S_1, ..., S_m$  are a sample from [S|Y]

•  $T_i = \exp(S_i) \Rightarrow T_1, ..., T_m$  are a sample from [T|Y]

**Exercise:** is  $T(x) = \exp\{S(x)\}$  really the correct target?

# Swiss rainfall data



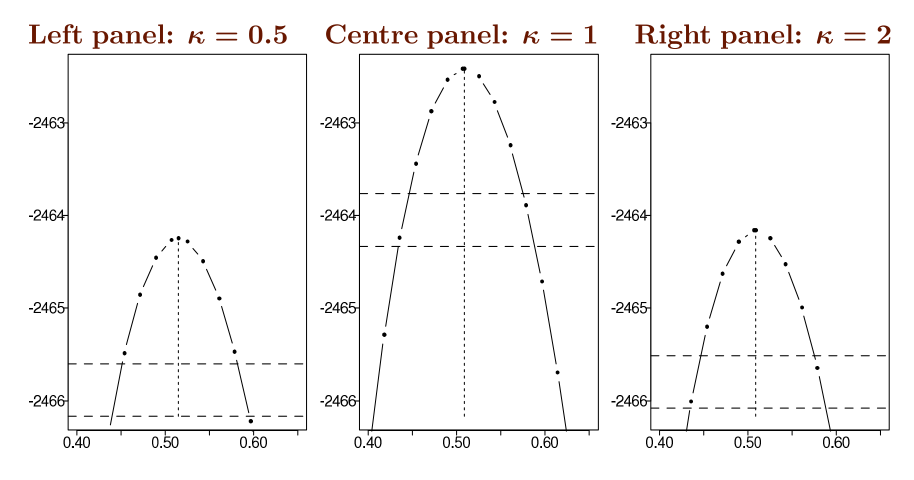
## Swiss rainfall: trans-Gaussian model

$$Y_i^* = h_\lambda(Y_i) = \left\{egin{array}{cc} (Y_i^\lambda - 1)/\lambda & ext{if } \lambda 
eq 0 \ \log(Y_i) & ext{if } \lambda = 0 \end{array}
ight.$$

For log-likelihood, write  $h_{\lambda} = h_{\lambda}(Y_1), ..., h_{\lambda}(Y_n),$ 

$$egin{aligned} \ell(eta, heta,\lambda) &=& -rac{1}{2}\{\log|\sigma^2 V|+(h_\lambda-Deta)'\{\sigma^2 V\}^{-1}(h_\lambda-Deta)\}\ &+(\lambda-1)\sum_{i=1}^n\log(Y_i) \end{aligned}$$

# Swiss rainfall: profile log-likelihoods for $\lambda$

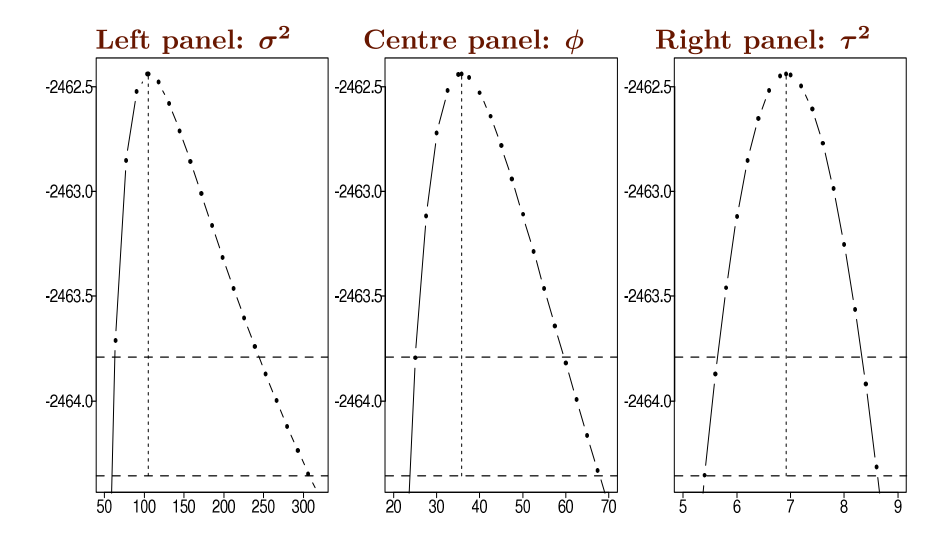


# Swiss rainfall: MLE's ( $\lambda = 0.5$ )

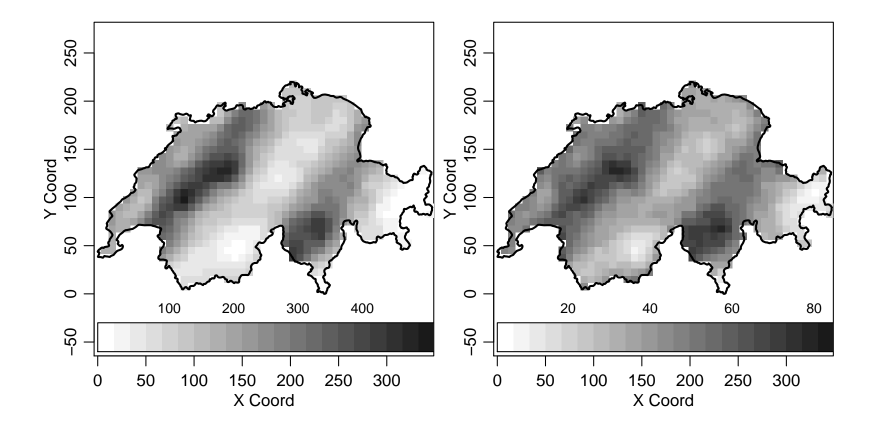
$\kappa$	$\hat{\mu}$	$\hat{\sigma}^2$	$\hat{\phi}$	$\hat{ au}^2$	$\log \hat{L}$
0.5	18.36	118.82	87.97	2.48	-2464.315
1	<b>20.13</b>	105.06	35.79	6.92	-2462.438
<b>2</b>	<b>21.36</b>	88.58	17.73	8.72	-2464.185

Likelihood criterion favours  $\kappa=1$ 

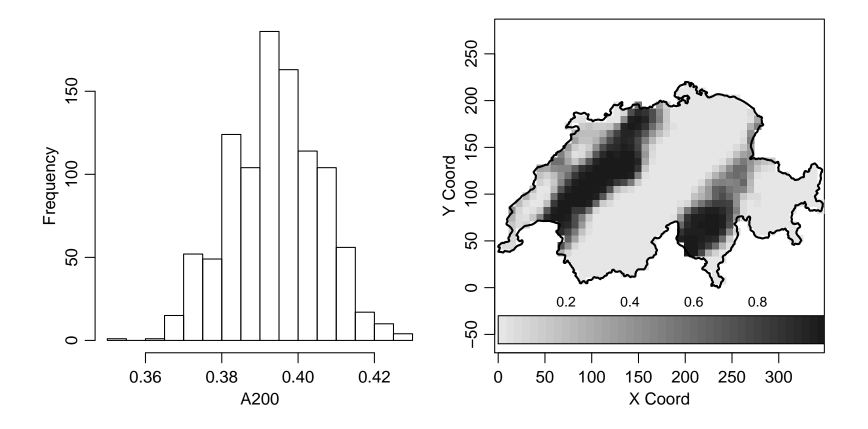
# Swiss rainfall: profile log-likelihoods $(\lambda = 0.5, \kappa = 1)$



# Swiss rainfall: plug-in predictions and prediction variances



## Swiss rainfall: non-linear prediction



Left-panel: plug-in prediction for proportion of total area with rain exceeding  $200 \ (= 20 \text{mm})$ 

Right-panel: plug-in prediction for P(rain > 250|Y)

# **Bayesian inference: basics**

Model specification

 $[Y,S,\theta] = [\theta][S|\theta][Y|S,\theta]$ 

**Parameter estimation** 

• integration gives

$$[Y, heta]=\int [Y,S, heta]dS$$

• Bayes' Theorem gives posterior distribution

 $[\theta|Y] = [Y|\theta][\theta]/[Y]$ 

• where  $[Y] = \int [Y|\theta][\theta] d\theta$ 

#### Prediction: $S \to S^*$

• expand model specification to

 $[Y,S^*, heta]=[ heta][S| heta][Y|S, heta][S^*|S, heta]$ 

• plug-in predictive distribution is

 $[S^*|Y, \hat{ heta}]$ 

• Bayesian predictive distribution is

$$[S^*|Y] = \int [S^*|Y, heta][ heta|Y]d heta$$

• for any target  $T = t(S^*)$ , required predictive distribution [T|Y] follows

# Notes

- likelihood function is central to both classical and Bayesian inference
- Bayesian prediction is a weighted average of plug-in predictions, with different plug-in values of  $\theta$  weighted according to their conditional probabilities given the observed data.
- Bayesian prediction is usually more conservative than plug-in prediction

# **Bayesian computation**

- 1. Evaluating the integral that defines  $[S^*|Y]$  is often difficult
- 2. Markov Chain Monte Carlo methods are widely used
- 3. but for geostatistical problems, reliable implementation of MCMC is not straightforward (no natural Markovian structure)
- 4. INLA is a serious competitor to MCMC (Rue, Martino and Chopin, 2009), but in current implementations only delivers marginal predictive distributions

 $\{[S(x_k)|Y]: k = 1, ..., N\}$  NOT  $[\{S(x_k: k = 1, ..., N\}|Y]$ 

5. for the Gaussian model, direct simulation is available

# Gaussian models: known $(\sigma^2, \phi)$

#### $Y \sim \mathcal{N}(D\beta, \sigma^2 R(\phi))$

- choose conjugate prior  $\beta \sim \mathrm{N}\left(m_{m eta} \; ; \; \sigma^2 V_{m eta} 
  ight)$
- posterior for  $\beta$  is  $\left[\beta|Y, \sigma^2, \phi\right] \sim N\left(\hat{\beta}, \sigma^2 V_{\hat{\beta}}\right)$

$$\hat{\beta} = (V_{\beta}^{-1} + D'R^{-1}D)^{-1}(V_{\beta}^{-1}m_{\beta} + D'R^{-1}y)$$

$$V_{\hat{\beta}} = \sigma^{2} (V_{\beta}^{-1} + D'R^{-1}D)^{-1})$$

• predictive distribution for  $S^*$  is

$$p(S^*|Y,\sigma^2,\phi) ~=~ \int p(S^*|Y,eta,\sigma^2,\phi) \, p(eta|Y,\sigma^2,\phi) \, deta.$$

# Notes

- mean and variance of predictive distribution can be written explicitly (but not given here)
- predictive mean compromises between prior and weighted average of Y
- predictive variance (not shown) has three components:
  - a priori variance,
  - minus information in data
  - plus uncertainty in  $\beta$
- limiting case  $V_{\beta} \to \infty$  corresponds to ordinary kriging.

# Gaussian models: unknown $(\sigma^2, \phi)$

Convenient choice of prior is:

 $[eta|\sigma^2,\phi] \sim \mathrm{N}\left(m_b,\sigma^2 V_b
ight) \quad [\sigma^2|\phi] \sim \chi^2_{ScI}\left(n_\sigma,S^2_\sigma
ight) \quad [\phi] \sim \mathrm{arbitrary}$ 

- results in explicit expression for  $[\beta, \sigma^2 | Y, \phi]$  and computable expression for  $[\phi | Y]$  whose form depends on choice of prior for  $\phi$
- in practice, use arbitrary discrete prior for  $\phi$  and combine posteriors conditional on  $\phi$  by weighted averaging

#### Algorithm 1:

- 1. choose lower and upper bounds for  $\phi$ , assign a discrete uniform prior for  $\phi$  over the chosen range
- 2. compute posterior  $[\phi|Y]$  on this discrete support set
- 3. sample  $\phi$  from posterior,  $[\phi|Y]$
- 4. attach sampled value of  $\phi$  to conditional posterior,  $[\beta, \sigma^2 | y, \phi]$ , and sample  $(\beta, \sigma^2)$  from this distribution
- 5. repeat steps (3) and (4) as many times as required, to generate a sample from the joint posterior,  $[\beta, \sigma^2, \phi|Y]$

Predictive distribution  $[S^*|Y, \phi]$  is tractable, hence write

$$p(S^*|Y) = \int p(S^*|Y,\phi) \; p(\phi|y) \, d\phi = \mathrm{E}_{\phi|Y}[p(S^*|Y,\phi)]$$

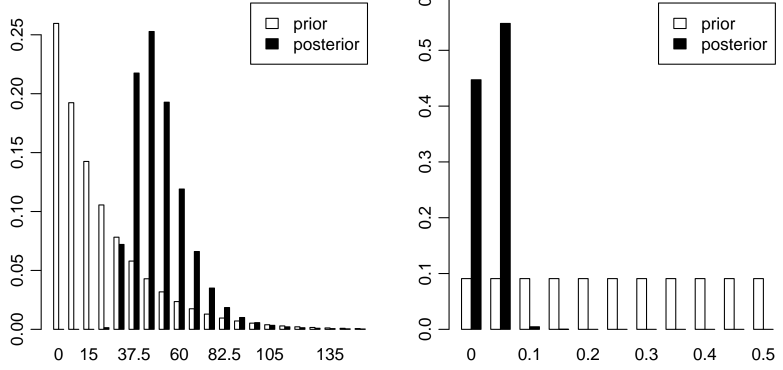
#### Algorithm 2:

- 1. discretise  $[\phi|Y]$ , as in Algorithm 1.
- 2. compute posterior  $[\phi|Y]$
- 3. sample  $\phi$  from posterior  $[\phi|Y]$
- 4. attach sampled value of  $\phi$  to  $[S^*|y, \phi]$  and sample from this to obtain realisations from  $[S^*|Y]$
- 5. repeat steps (3) and (4) as required

Note: Extends immediately to multivariate  $\phi$  (but may be computationally awkward)

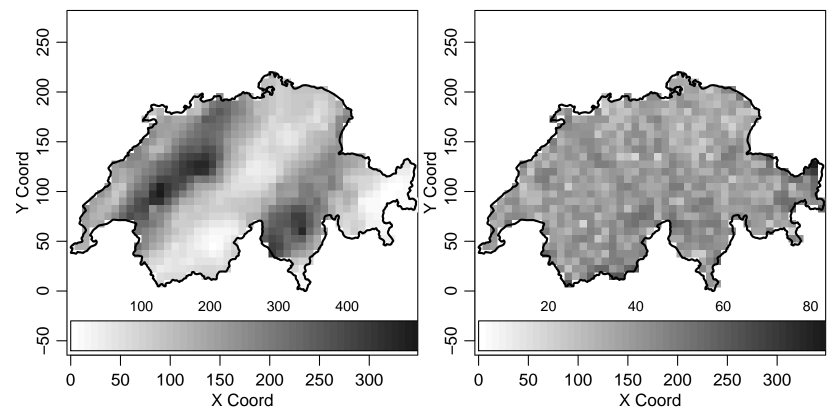
# Swiss rainfall





## Swiss rainfall

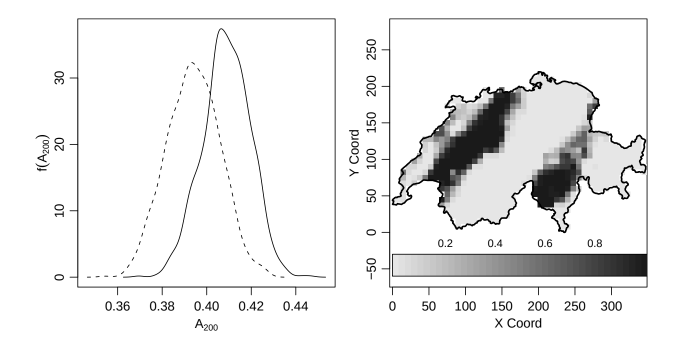
# Mean (left-panel) and variance (right-panel) of predictive distribution



# Swiss rainfall: posterior means and 95% credible intervals

parameter	estimate	95% interval
$\beta$	144.35	[53.08,  224.28]
$\sigma^2$	13662.15	[8713.18, 27116.35]
$oldsymbol{\phi}$	<b>49.97</b>	[30,  82.5]
$ u^2$	0.03	[0, 0.05]

### Swiss rainfall: non-linear prediction



Left-panel: Bayesian (solid) and plug-in (dashed) prediction for proportion of total area with rainfall exceeding 200 (= 20mm) Right-panel: Bayesian predictive map of P(rainfall > 250|Y)

# Computation with geoR

```
MC<-model.control()
?model.control
PC<-prior.control(beta.prior="flat",sigmasq.prior="sc.inv.chisq",
    sigmasq=0.2,df.sigmasq=4,phi.discrete=0.1*(1:5),
    tausq.rel.prior="uniform",tausq.rel.discrete=0.1*(0:3))
OC<-output.control(n.posterior=100,n.predictive=100,
    simulations.predictive=T,signal=T,moments=F)
set.seed(24367)
results.bayes<-krige.bayes(geodata=loglead2,locations=grid,
    borders=region,model=MC,prior=PC,output=OC)</pre>
```

```
names(results.bayes)
posterior.bayes<-results.bayes$posterior
names(posterior.bayes)
posterior.sample<-posterior.bayes$sample
par.names<-names(posterior.sample)
par(mfrow=c(2,2))
for (i in 1:4) {
    hist(posterior.sample[,i],xlab=par.names[i],main=" ")
    }
par(mfrow=c(1,1))
plot(posterior.sample[,2],posterior.sample[,3],
    xlab=par.names[2],ylab=par.names[3])</pre>
```

```
par(mfrow=c(1,1),pty="s")
predictions.bayes <- results.bayes $predictive
image(unique(grid[,1]),unique(grid[,2]),
   matrix(predictions.bayes$mean.simulations,26,26))
points(loglead2,add=T); lines(coast[,1],coast[,2])
par(mfrow=c(1,2))
predict.max<-NULL</pre>
for (sim in 1:100) {
   predict.max<-c(predict.max,max(predictions$simulations[,sim]))</pre>
   }
hist(predict.max,xlab="predictive distribution of maximum",
   main="plug-in",breaks=0.1*(16:28))
predict.bayes.max<-NULL</pre>
for (sim in 1:100) {
   predict.bayes.max<-c(predict.bayes.max,</pre>
      max(predictions.bayes$simulations[,sim]))
   }
hist(predict.bayes.max,xlab="predictive distribution of maximum",
   main="Bayesian", breaks=0.1*(16:28))
```

# Generalized linear geostatistical model (GLGM)

- Latent spatial process $S(x) \sim \mathrm{SGP}\{0,\sigma^2,
  ho(u)\}$  $ho(u) = 
  ho_0(-|u|/\phi)$
- Linear predictor

$$\eta(x) = d(x)'\beta + S(x)$$

- Link function $\mathrm{E}[Y_i] = \mu_i = h\{\eta(x_i)\}$
- Conditional distribution for  $Y_i: i=1,...,n$  $Y_i|S(\cdot)\sim f(y;\eta)$  mutually independent

# GLGM

- usually just a single realisation is available, in contrast with GLMM for longitudinal data analysis
- GLGM approach is most appealing when there is a natural sampling mechanism, for example Poisson model for counts or logistic-linear models for proportions
- transformed Gaussian models may be more useful for non-Gaussian continuous responses
- theoretical variograms can be derived but are less natural as summary statistics than in Gaussian case
- but empirical variograms of GLM residuals can still be useful for exploratory analysis

# The Loa loa prediction problem

Ground-truth survey data

- random sample of subjects in each of a number of villages
- blood-samples test positive/negative for Loa loa

Environmental data (satellite images)

- measured on regular grid to cover region of interest
- elevation, green-ness of vegetation

Objectives

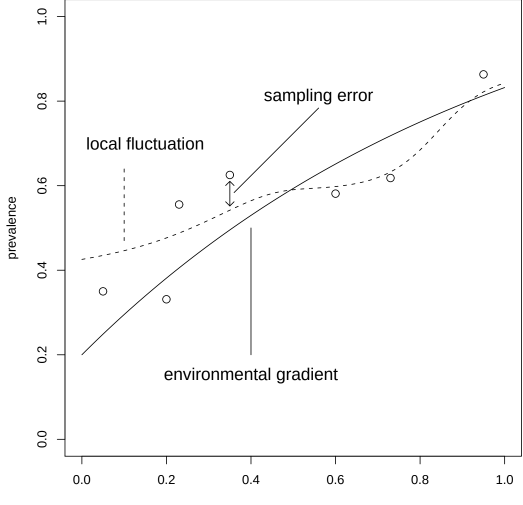
- predict local prevalence throughout study-region (Cameroon)
- compute local exceedance probabilities,

P(prevalence > 0.2|data)

# Loa loa: a generalised linear model

- Latent spatial process $S(x) \sim \mathrm{SGP}\{0, \sigma^2, \rho(u))\}$  $ho(u) = \exp(-|u|/\phi)$
- Linear predictor
  - $egin{aligned} d(x) &= ext{environmental variables at location } x \ \eta(x) &= d(x)'eta + S(x) \ p(x) &= \log[\eta(x)/\{1-\eta(x)\}] \end{aligned}$
- Error distribution $Y_i | S(\cdot) \sim \mathrm{Bin}\{n_i, p(x_i)\}$

# Schematic representation of Loa loa model

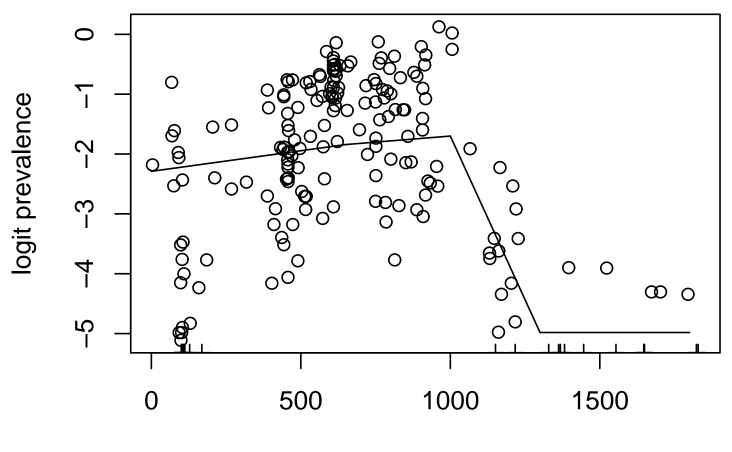


location

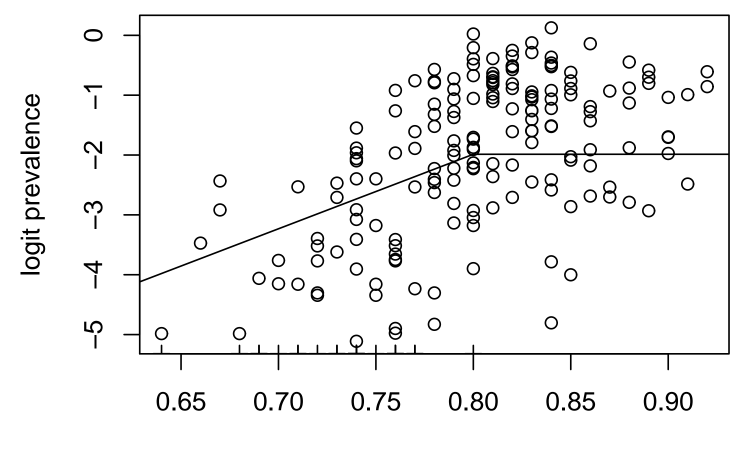
# The modelling strategy

- use relationship between environmental variables and groundtruth prevalence to construct preliminary predictions via logistic regression
- use local deviations from regression model to estimate smooth residual spatial variation
- Bayesian paradigm for quantification of uncertainty in resulting model-based predictions

logit prevalence vs elevation



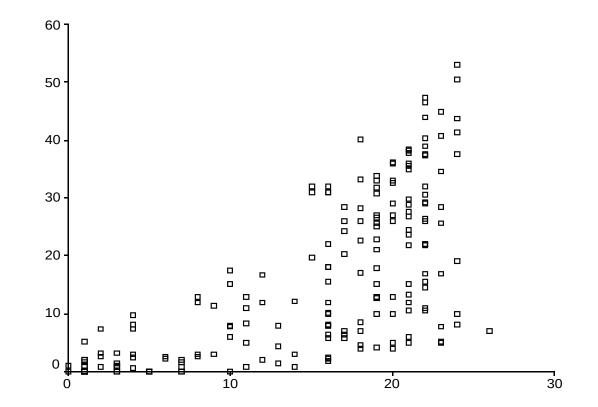
elevation



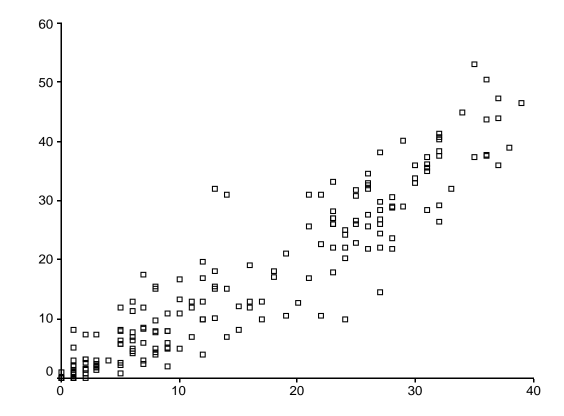
Max Greeness

# Comparing non-spatial and spatial predictions in Cameroon

#### **Non-spatial**



### **Spatial**



# **Probabilistic prediction in Cameroon**

Observed prevalence of loa loa (IRD-TDR)

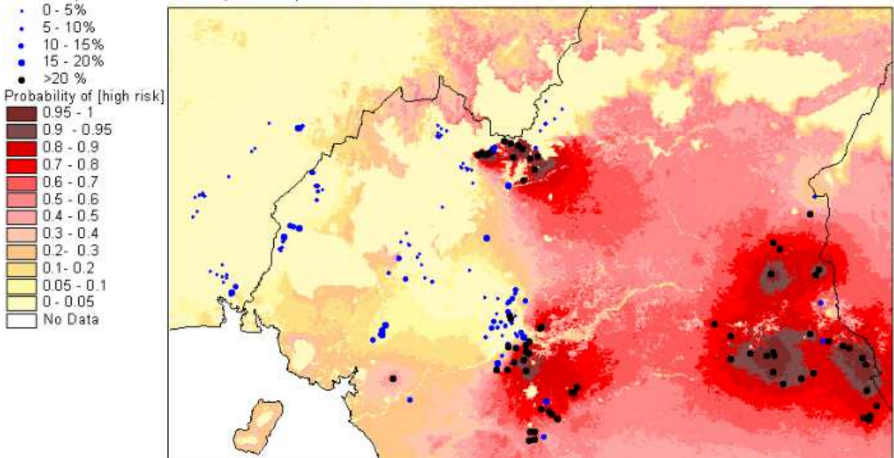


Figure 6: PCM for [high risk] in Cameroon based on ERMr with ground truth data.

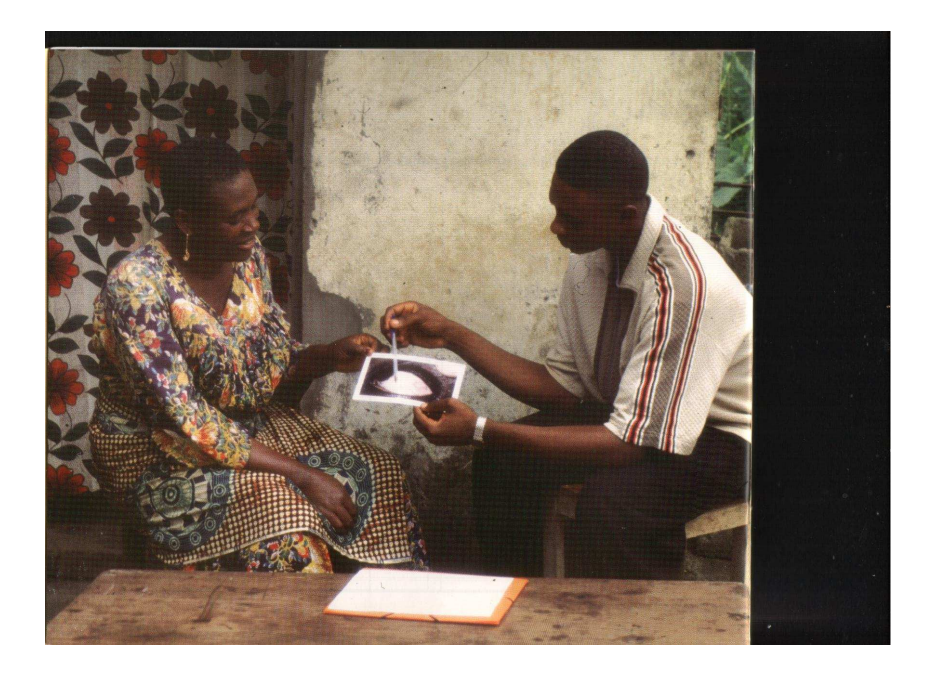
# Next Steps

How can we improve the precision of our predictive inferences?

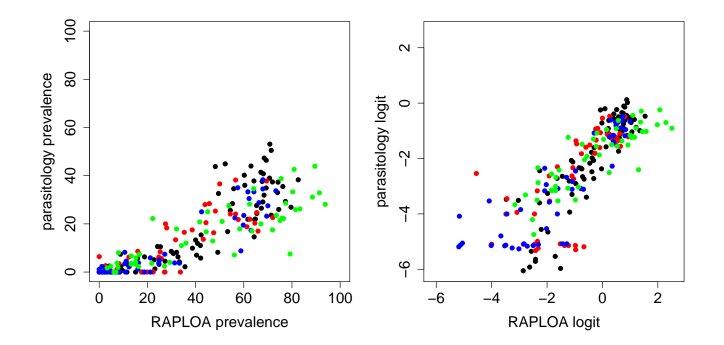


# RAPLOA

- A cheaper alternative to parasitological sampling:
  - have you ever experienced eye-worm?
  - did it look like this photograph?
  - did it go away within a week?
- RAPLOA data to be collected:
  - in sample of villages previously surveyed (to calibrate parasitology vs RAPLOA estimates)
  - in villages not previously surveyed (to reduce local uncertainty)
- Calibration model needed to reconcile parasitological and RAPLOA prevalence estimates



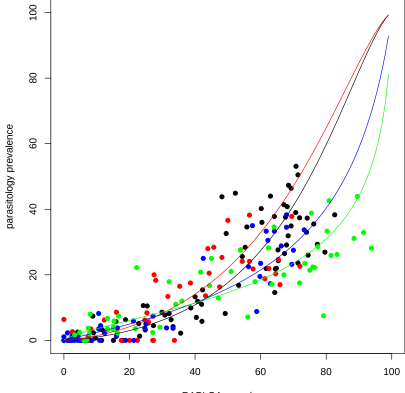
### **RAPLOA** calibration



Empirical logit transformation linearises relationship Colour-coding corresponds to four surveys in different regions

# **RAPLOA** calibration (ctd)

Fit linear functional relationship on logit scale and back-transform



**RAPLOA** prevalence

### **Bivariate geostatistical models**

$$egin{array}{rll} Y_{1i} &=& S_1(x_{1i})+Z_{1i}:i=1,...,n_1 \ Y_{2j} &=& S_2(x_{2j})+Z_{2j}:j=1,...,n_2 \end{array}$$

- OK to assume  $Z_{1i}, Z_{2j}$  independent?
- how to model correlation between  $S_1(x)$  and  $S_2(x')$ ?
- common sampling locations?
- symmetric or asymmetric association?

**Open question:** how to construct bivariate geostatistical models with spatially invariant calibration properties.

# 5. Discrete spatial variation

- Joint vs conditional specification
- Markov random field models

# **Conditional specification of joint distributions**

Theorem

$$rac{f(y)}{f(z)} = \prod_{i=1}^n rac{f_i(y_i|y_1,...,y_{i-1},z_{i+1},...,z_n)}{f_i(z_i|y_1,...,y_{i-1},z_{i+1},...,z_n)}$$

**Outline of proof** 

Case n = 3 sufficient to show the idea, as follows

$$egin{aligned} f(y_1,y_2,y_3) &= & f(y_3|y_1,y_2) imes f(y_2,y_1) \ &= & rac{f(y_3|y_1,y_2)}{f(z_3|y_1,y_2)} imes f(z_3|y_1,y_2) imes f(y_1,y_2) \ &= & rac{f(y_3|y_1,y_2)}{f(z_3|y_1,y_2)} imes f(y_1,y_2,z_3) \end{aligned}$$

Same argument gives

$$egin{array}{rll} f(y_1,y_2,z_3)&=&f(y_2|y_1,z_3) imes f(y_1,z_3)\ &=&rac{f(y_2|y_1,z_3)}{f(z_2|y_1,z_3)} imes f(y_1,z_2,z_3) \end{array}$$

and so on, to give required result.

Exercise 2.2.2 (from preliminary material) re-visited

$$Y_i = \alpha(Y_{i-1} + Y_{i+1}) + Z_t : Z_i \sim N(0, \tau^2)$$

Full conditional of  $Y_i$  depends on  $Y_{i-2}$ ,  $Y_{i-1}$ ,  $Y_{i+1}$  and  $Y_{i+2}$ .

• Re-write model in vector-matrix notation as

$$Y = AY + Z \Leftrightarrow Y = (I - A)^{-1}Z$$

where (using n = 5 for illustration)

$$A = \left[egin{array}{cccccccc} 0 & lpha & 0 & 0 & 0 \ lpha & 0 & lpha & 0 & 0 \ 0 & lpha & 0 & lpha & 0 \ 0 & lpha & 0 & lpha & 0 \ 0 & 0 & lpha & 0 & lpha \ 0 & 0 & lpha & 0 & lpha \end{array}
ight]$$

• Then,  $Y \sim \text{MVN}(0, \tau^2 (I - A)^{-2})$ 

- Standard result from graphical modelling is that non-zero elements in  $Var(Y)^{-1}$  identify conditional dependencies (eg Whittaker, 1990, Proposition 5.7.3)
- Straightforward matrix algebra gives

$$(I-A)^2 = egin{bmatrix} 1+lpha^2 & -2lpha & lpha^2 & 0 & 0 \ -2lpha & 1+2lpha^2 & -2lpha & lpha^2 & 0 \ lpha^2 & -2lpha & 1+2lpha^2 & -2lpha & lpha^2 \ 0 & lpha^2 & -2lpha & 1+2lpha^2 & -2lpha \ 0 & 0 & lpha^2 & -2lpha & 1+lpha^2 \end{bmatrix}$$

• Third row of  $(I - A)^2$  gives required result (no non-zero elements)

# Hammersley-Clifford

Previous result says joint distribution of Y is determined by full conditionals provided full conditionals are self-consistent

General result: for any  $A \subset \{1, 2, ..., n\}$ , write  $\mathcal{Y}_A = \{y_i : i \in A\}$ , then

$$f(y) = \exp\left\{\sum_{A \subset \{1,2,\dots,n\}} h(\mathcal{Y}_A)
ight\}$$
 (1)

**Definitions:** 

- 1) for any set of full conditionals  $f_i(y_i | \{y_j : j \neq i\})$ , index j is a neighbour of i if  $f_i(\cdot)$  depends on  $y_j$
- 2) a clique is a set of mutual neighbours.

Theorem (Hammersley-Clifford)

Expression (1) gives valid specification of f(y) if and only if:

- 1.  $h(\mathcal{Y}_A) = 0$  for all non-cliques A
- 2. f(y) integrable (so can scale to  $\int f(y) = 1$ )
- 3. if  $f(y_j) > 0$  for all  $j \in A$ , then  $f(\mathcal{Y}_A) > 0$

Besag, 1974

### Markov Random Field (MRF) models

- Random vector  $Y = (Y_1, ..., Y_n)$
- joint distribution [Y] fully specified by full conditionals,  $[Y_i|\{Y_j:j\neq i\}]:i=1,...,n$
- neighbourhood of i is  $\mathcal{N}(i) \subset \{1,2,...,n\}$
- MRF:  $[Y_i | \{Y_j : j \neq i\}] = [Y_i | Y_j : j \in \mathcal{N}(i)] : i = 1, ..., n$

# **Examples of MRF models**

1. Binary  $Y_i$ : auto-logistic model

$$p_i = \mathrm{P}(Y_i = 1 | \{Y_j : j \neq i\}) \quad \mathrm{logit} p_i = lpha + eta \sum_{j \in \mathcal{N}(i)} Y_j$$

Higher-order models defined naturally on regular lattices:

	4	3	4	
4	2	1	2	4
3	1	•	1	3
4	2	1	2	4
	4	3	4	

$$ext{logit} p_i = lpha + \sum_{k=1}^m eta_k \sum_{j \in \mathcal{N}_k(i)} Y_j$$

#### 2. Count $Y_i$ : auto-Poisson model

$$\mu_i = \mathrm{E}[Y_i|\{Y_j: j 
eq i\}] \quad \log \mu_i = lpha + eta \sum_{j \in \mathcal{N}(i)} Y_j$$

Restriction: the auto-Poisson model only defines a proper distribution when  $\beta \leq 0$ 

3. Hierarchical model with latent Gaussian MRF

A better way to model spatial count data:

• latent Gaussian MRF  $S = (S_1, ..., S_n)$ 

• conditionally independent  $Y_i | S \sim \text{Poiss}(\alpha + \beta S_i)$ 

Even better if  $\alpha$  is replaced by  $\alpha_i = d'_i \theta$  for vector of spatial explanatory variables  $d_i$ 

Besag, York and Mollié, 1991

# **Computational appeal of MRF models**

• Gaussian MRF

Mean  $\mu$ , precision matrix  $\Omega = {\operatorname{Var}(Y)}^{-1}$ , log-likelihood is

$$L=0.5n\log|\Omega|-0.5(Y-\mu)'\Omega(Y-\mu)$$

Markov structure implies that  $\Omega$  is sparse

• Gaussian or non-Gaussian MRF

Gibbs sampler for MCMC follows directly from model specification through full conditionals,

$$[Y_i | \{Y_j : j \neq i\}] : i = 1, ..., n$$

# Limitations of MRF models for spatial data

- MRF's are just multivariate probability distributions
  - parameterised in a way that has a spatial interpretation
  - but specific to a fixed set of locations  $x_1, ..., x_n$
- neighbourhood specification can be problematic
  - natural hierarchy of models on regular lattices
  - not so for irregular lattices
  - and arguably un-natural for spatially aggregated data,

$$Y_i = \int_{A_i} Y(x) dx$$

# 6. Spatial point processes

- exploratory analysis
- Cox processes and the link to continuous spatial variation
- pairwise interaction processes and the link to discrete spatial variation.

# Notation

- spatial point process: countable set of events  $x_i \in \mathbb{R}^2$
- $N(A) = \#(x_i \in A)$  for spatial region  $A \subset \mathbb{R}^2$
- stationary if properties invariant under translation
- isotropic if properties invariant under rotation
- orderly if no multiple coincident events

### The Poisson Process

1.  $N(A) \sim \text{Poiss}(\mu(A))$ , where

$$\mu(A) = \int_A \lambda(x) dx$$

2. given N(A) = n, events  $x_i \in A$  iid,  $pdf \propto \lambda(x)$ 

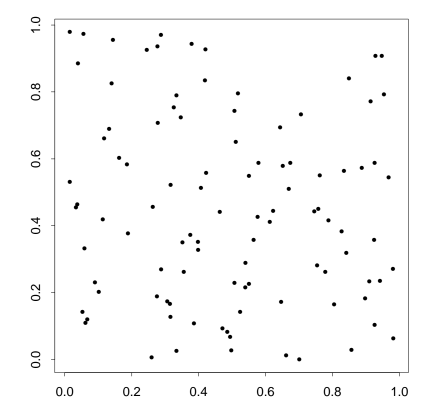
Complete spatial randomness:  $\lambda(x) = \lambda$ 

Properties

- 1. N(A) and N(B) independent when A and B disjoint
- 2.  $Var{N(A)}/E[N(A)] = 1$ , for all A
- 3. distance from an arbitrary point to the nearest event:

$$F(x) = 1 - \exp(-\pi\lambda x^2) : x > 0$$

# Partial realisation of a Poisson process



### **Point process intensities**

Def 6.1. The (first-order) intensity function of a spatial point process is

$$\lambda(x) = \lim_{|dx| o 0} \left\{ rac{E[N(dx)]}{|dx|} 
ight\}$$

Def 6.2. The second-order intensity function of a spatial point process is

$$\lambda_2(x,y) = \lim_{\substack{|dx| o 0 \ |dy| o 0}} \left\{ rac{E[N(dx)N(dy)]}{|dx||dy|} 
ight\}$$

Def 6.3. The covariance density of a spatial point process is

$$\gamma(x,y) = \lambda_2(x,y) - \lambda(x)\lambda(y).$$

What if process is stationary and isotropic?

(i)  $\lambda(x) \equiv \lambda = E[N(A)]/|A|$ ,

(constant, for all A).

(ii)  $\lambda_2(x, y) \equiv \lambda_2(||x - y||)$  (depends only on distance)

(iii)  $\gamma(u) = \lambda_2(u) - \lambda^2$ .

### The K-function

Def 6.4 The reduced second moment function of a stationary, isotropic spatial point process is

$$K(s)=2\pi\lambda^{-2}\int_0^s\lambda_2(r)rdr.$$

Theorem 6.1. For a stationary, isotropic, orderly process:  $K(s) = \lambda^{-1} E$ [number of further events within distance s of an arbitrary event]

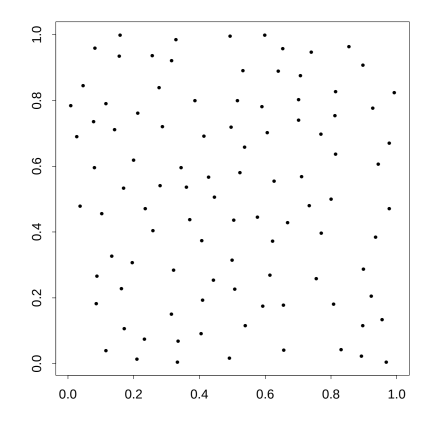
- gives a tangible interpretation of K(s)
- suggests a method of estimating K(s) from data

Theorem 6.2. For a homogeneous, planar Poisson process,

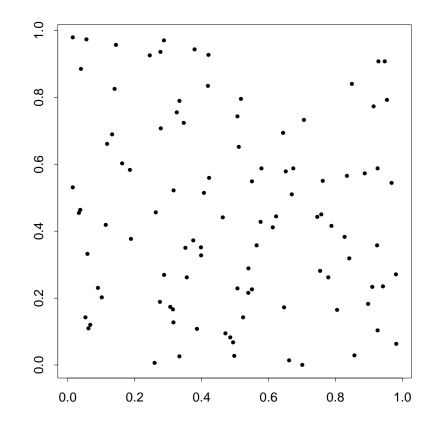
$$K(s) = \pi s^2$$

# Three pictures

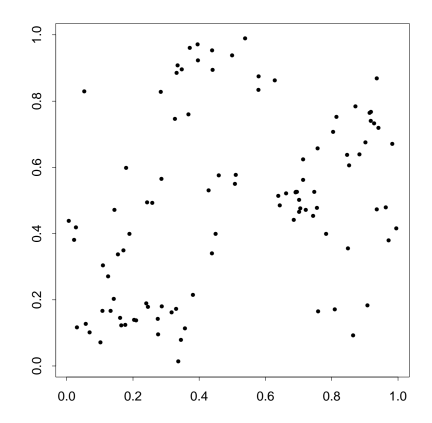
#### Regular



#### **Completely random**



#### Aggregated



## A useful property of the K-function

Def 6.5. A random thinning, P', of a point process P, is a point process whose events are a sub-set of the events of P generated by retaining or deleting the events of P in a series of mutually independent Bernoulli trials.

Theorem 6.3. K(s) is invariant to random thinning.

**Proof.** Exercise (use Theorem 6.1)

Implication: the interpretation of an estimated K-function is robust to incomplete ascertainment of events, provided the incompleteness is spatially neutral.

# Estimating the K-function

Data:  $x_i \in A : i = 1, \ldots, n$ 

Estimation of  $\lambda$ 

$$\hat{\lambda}=n/|A|$$

Estimation of K(s)

 $\lambda K(s) = E[$ number of further events within distance s of an arbitrary event]

1. Define  $E(s) = \lambda K(s)$ .

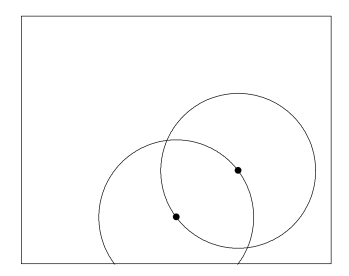
2. Let  $d_{ij}$  be the distance between the events  $x_i$  and  $x_j$ .

3. Define

$$\tilde{E}(s) = n^{-1} \sum_{i=1}^{n} \sum_{j \neq i} I(d_{ij} \leq s)$$

- 4. The estimator  $\tilde{E}(s)$  is negatively biased because we do not observe events outside A
- 5. Introduce weights,

 $w_{ij}$  = reciprocal of proportion of circumference of circle, centre  $x_i$  and radius  $d_{ij}$ , which is contained in A.



6. An edge-corrected estimator for E(s) is

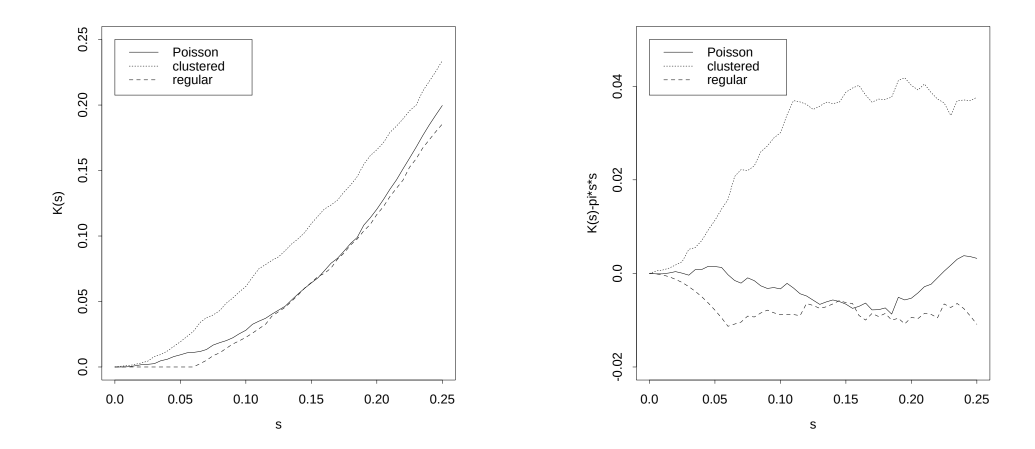
$$\hat{E}(s) = n^{-1} \sum_{i=1}^n \sum_{j \neq i} w_{ij} I(d_{ij} \leq s).$$

where  $I(\cdot)$  is the indicator function.

7. Since  $K(s) = E(s)/\lambda$ , define

$$egin{array}{rcl} \hat{K}(s)&=&\hat{E}(s)/\hat{\lambda}\ &=&n^{-2}|A|\sum_{i=1}^n\sum_{j
eq i}w_{ij}I(d_{ij}\leq s) \end{array}$$

#### Estimates $\hat{K}(s)$ for three simulated patterns



## **Bivariate K-functions**

 $\lambda_j : j = 1, 2$  denotes intensity of type j events.

- $\lambda_j K_{ij}(s) =$  expected number of further type j events within distance s of an arbitrary type i event
  - if type j events are a homogeneous Poisson process, then

$$K_{jj}(s) = \pi s^2$$

• if type 1 and type 2 events are independent processes, then

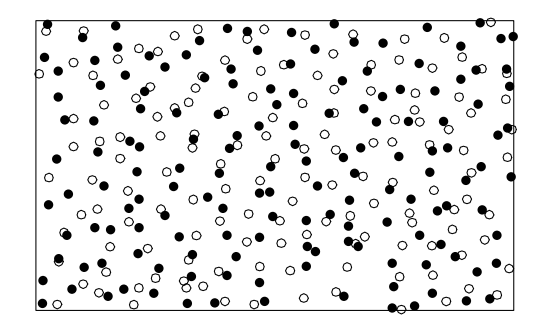
$$K_{12}(s) = \pi s^2$$

• if type 1 and type 2 events are a random labelling of a univariate process with K-function K(s), then

$$K_{11}(s) = K_{12}(s) = K_{22}(s) = K(s)$$

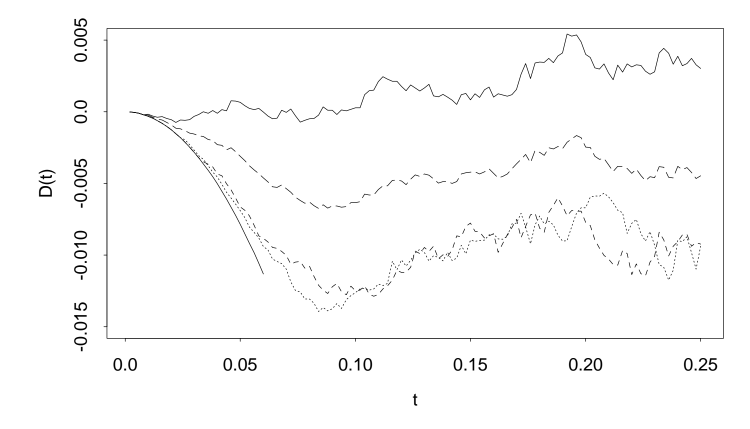
An example: displaced amacrine cells in rabbit retina

- type 1 events transmit information to the brain when a light goes on
- type 2 events transmit information to the brain when a light goes off
- interest is in discriminating between two developmental hypotheses:
  - 1. on and off cells are initially generated in separate layers which later fuse to form the mature retina
  - 2. on and off cells are initially undifferentiated in a single layer and acquire their distinct functionality at a later stage



Solid/open circles respectively identify on/off cells

#### Second-order properties:



Functions plotted are  $\hat{D}(t) = \hat{K}(t) - \pi t^2$  as follows: ---: on cells;  $\cdots :$  off cells; ---: all cells; ---: bivariate.

The parabola  $-\pi t^2$  is also shown as a solid line.

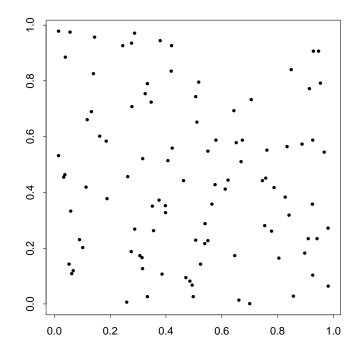
# **Computation with splancs**

```
#
# Exploratory analysis of amacrine cell data
#
library(splancs)
on<-scan("amacrines_on.data")</pre>
length(on)
on<-matrix(on,152,2,T)</pre>
off<-scan("amacrines_off.data")</pre>
length(off)
off<-matrix(off,142,2,T)
a<-1060/662
poly<-matrix(c(0,0,a,0,a,1,0,1),4,2,T)
par(pty="s",mfrow=c(1,1))
polymap(poly)
pointmap(on,add=T,pch=19,col="red")
pointmap(off,add=T,pch=19,col="blue")
```

```
?khat
s<-0.005*(0:51)
k.on<-khat(on,poly,s)
k.off<-khat(off,poly,s)
plot(s,k.on-pi*s*s,type="l",col="red",ylim=c(-0.015,0.005))
lines(s,k.off-pi*s*s,col="blue")
k.cross<-k12hat(on,off,poly,s)
lines(s,k.cross-pi*s*s)
```

# Three pictures re-visited

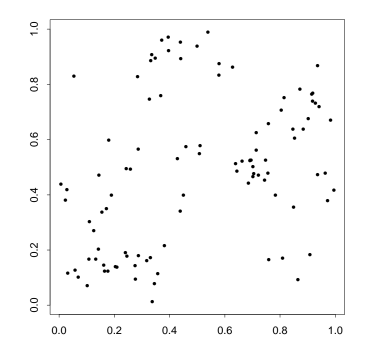
**Completely random** 



#### A Poisson process

- $N(A) \sim \operatorname{Poiss}(\lambda |A|)$
- conditional on N(A) = n, events  $x_i \sim \text{iid } U(A)$

#### Aggregated



#### A Cox process

- $\Lambda(x)$  a non-negative-valued spatial stochastic process
- conditional on  $\Lambda(x) = \lambda(x)$ , process is inhomogeneous Poisson (Cox, 1955)

Picture:  $\Lambda(x) = \sum g(x - X_i)$ 

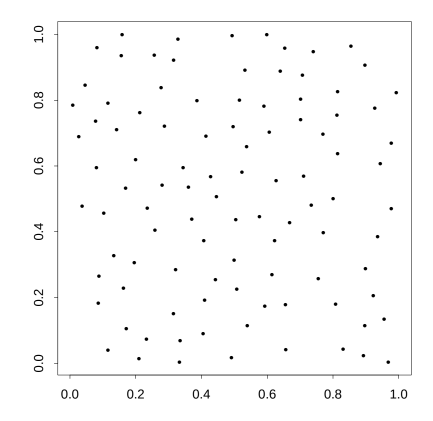
- $X_i: i = 1, 2, ...$  homogenous Poisson process
- $g(\cdot) = \text{bivariate Gaussian density, } N(0, \sigma^2 I)$

This process can also be interpreted as a Poisson cluster process in which:

- parents  $X_i$  form a homogeneous Poisson process
- number of offspring per parent are iid Poisson-distributed
- locations of offspring relative to their parent are iid with pdf  $g(\cdot)$
- observed pattern consists of offspring only

Bartlett, 1964

### Regular



#### An inhibitory process

- events  $\mathcal{X} = \{x_1, ..., x_n\}$  in spatial region A
- $LR(\mathcal{X}) =$  likelihood ratio for  $\mathcal{X}$  wrt Poisson process of unit intensity
- non-negative-valued interaction function  $h(u): u \ge 0$

$$LR(\mathcal{X}) \propto eta^n \prod_{j 
eq i} h(||x_i - x_j||)$$

**Picture:** 

$$h(u) = \left\{egin{array}{ccc} 0 & : & u < \delta \ 1 & : & u \geq \delta \end{array}
ight.$$

## Poisson processes

• completely defined by their intensity function  $\lambda(x)$ 

 $- \ N(A) \sim \mathrm{Poiss}\left(\int_A \lambda(x) dx 
ight)$ 

- conditional on N(A) = n, events  $x_i : i = 1, ..., n$  are independent random sample from distribution with pdf  $f(x) \propto \lambda(x)$
- Log-likelihood function,

$$L( heta) = \sum_{i=1}^n \log \lambda(x_i; heta) - \int_A \lambda(x; heta) dx$$

• independence property often unrealistic, but may be a useful approximation

### Cox processes

- a Cox process is an inhomogeneous Poisson process with stochastic intensity  $\Lambda(x)$
- useful class of models for environmentally driven processes
- even more useful when environmental covariates can explain part of the variation in  $\Lambda(x)$

Cox, 1955

Link to continuous spatial variation (geostatistics)

Cox process:  $[\Lambda][\mathcal{X}|\Lambda]$ Geostatistical model: [S][Y|S] Cox processes: moment properties

Assume  $\Lambda(x)$  stationary with mean  $\lambda$  and covariance function  $\gamma(u)$ , then:

- $\lambda = \text{intensity}$
- $\gamma(u) = \text{covariance density}$

$$K(s)=\pi s^2+2\pi\lambda^{-2}\int_0^s\gamma(u)udu$$

## Cox processes: model-fitting

- likelihood generally intractable (except by Monte Carlo)
- ad hoc estimation by matching theoretical and empirical second moments (not entirely satisfactory)

$$D( heta) = \int_0^s w(u) \{\hat{K}(u) - K(u; heta)\}^2 du$$

Møller and Waagepetersen, 2004

# Pairwise interaction point processes (PIPPs)

- defined by their likelihood ratio wrt Poisson process
- useful for modelling inhibitory interactions between events
- can be derived as continuous limit of Poisson MRF models on a regular lattice

Besag, Milne and Zachary, 1982

• problematic for modelling attractive interactions (recall similar reservation wrt auto-Poisson model)

### **PIPPs:** formulation

- events  $\mathcal{X} = \{x_1, ..., x_n\}$  in spatial region A
- $LR(\mathcal{X}) =$  likelihood ratio for  $\mathcal{X}$  wrt Poisson process of unit intensity
- non-negative-valued interaction function  $h(u): u \ge 0$

$$LR(\mathcal{X}) \propto eta^n \prod_{j 
eq i} h(||x_i - x_j||)$$

- process well-defined if  $h(u) \leq 1$  for all u
- h(u) = 1 for all u gives homogeneous Poisson process

### **PIPP's: model-fitting**

Conditional intensity at x, given  $\mathcal{X} = \{x_1, ..., x_n\}$  in  $A - \{x\}$ ,

$$\lambda(x|\mathcal{X}) = eta \prod_{i=1}^n h(||x_i - x||)$$

- MCMC scheme for simulating realisations operates by alternating between:
  - adding event according to pdf  $f(x) \propto \lambda(x|\mathcal{X})$
  - deleting event at random

Ripley (1979) - note date!

• likelihood evaluation requires Monte Carlo methods

• pseudo-likelihood:

- treats  $\lambda_c(\cdot)$  as if unconditional intensity, hence

$$L( heta) = \sum_{i=1}^n \log \lambda_c(x_i | \mathcal{X} - \{x_i\}; heta) - \int_A \lambda(x | \mathcal{X}; heta) dx$$

- gives good starting values for Monte Carlo inference

Link to discrete spatial variation (Markov random fields)

# **Computation using spatstat**

```
#
# fitting a pairwise interaction point process to the
#amacrine "on" cells
#
library(spatstat)
library(splancs)
#
xy.on<-matrix(scan("amacrines_on.data"),152,2,T)
xy<-xy.on
?ppp
xy.ppp<-ppp(xy[,1],xy[,2],xrange=c(0,1060),yrange=c(0,662))</pre>
```

```
?ppm
?quadscheme
Q<-quadscheme(xy.ppp,nd=c(80,56))
#
# 80 by 56 quadrature grid gives approximate convergence of
# non-parametric estimate
#
stuff<-ppm(Q,interaction=PairPiece(r=20*(1:10)),</pre>
                               correction="Ripley")
h.nonparam.on<-c(0,0.0589,0.2857,0.6922,0.9524,1.0087,
                    0.9468,0.9230,0.8553,0.8415)
u.nonparam < -20 * (0:9) + 10
par(mfrow=c(1,1))
plot(u.nonparam,h.nonparam.on,type="l",xlab="r",ylab="h(u)")
```

## **PIPPs:** Monte Carlo likelihood

Likelihood function for PIPP with parameter  $\theta$  and data  $\mathcal{X}$  can always be written as

$$\ell(\theta) = a(\theta) LR(\mathcal{X}, \theta)$$

Circumvent intractability of normalising constant  $a(\theta)$  as follows:

• Write

$$egin{aligned} a( heta)^{-1} &= \int LR(\mathcal{X}, heta)d\mathcal{X} \ &= \int LR(\mathcal{X}, heta) imes rac{a( heta_0)}{a( heta_0)} imes rac{LR(\mathcal{X}, heta_0)}{LR(\mathcal{X}, heta_0)}d\mathcal{X} \end{aligned}$$

• Define  $r(\mathcal{X}, \theta, \theta_0) = LR(\mathcal{X}, \theta)/LR(\mathcal{X}, \theta_0)$ , then

$$egin{aligned} a( heta)^{-1} &=& a( heta_0)^{-1} \int r(\mathcal{X}, heta, heta_0) \ell(\mathcal{X}, heta_0) d\mathcal{X} \ &=& a( heta_0)^{-1} \mathrm{E}_{ heta_0}[r(\mathcal{X}, heta, heta_0)] \end{aligned}$$

• Since  $\theta_0$  is arbitrary, it follows that for any value  $\theta_0$ , the MLE  $\hat{\theta}$  maximises

$$L(\theta) = \log LR(\mathcal{X}, \theta) - \log E_{\theta_0}[r(\mathcal{X}, \theta, \theta_0)]$$

which in turn can be approximated by

$$L^*( heta) = \log LR(\mathcal{X}, heta) - \log \left\{ s^{-1} \sum_{j=1}^s r(\mathcal{X}_j, heta, heta_0)] 
ight\},$$

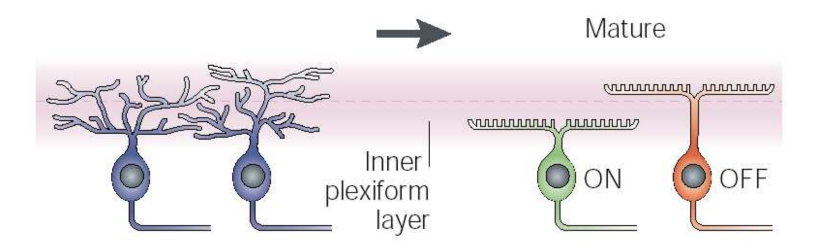
where  $\mathcal{X}_j : j = 1, ..., s$  are simulated with  $\theta = \theta_0$ 

#### Algorithm

- 1. Pick starting value  $\theta_0$  (eg maximum pseudo-likelihood estimate), and number of simulations s
- 2. Maximise resulting  $L^*(\theta)$  to give  $\theta = \tilde{\theta}$
- 3. Set  $\theta_0 = \tilde{\theta}$ , increase s and repeat

# **Example: displaced amacrine cells**

Biology (as of 2004)



Diggle, Eglen and Troy, 2006

### **Bivariate pairwise interaction point processes**

**Bivariate data** 

 $X_1 = \{x_{1i} : i = 1, ..., n_1\} \qquad X_2 = \{x_{2i} : i = 1, ..., n_2\}$ 

**Bivariate pairwise interaction model** 

 $f(X_1, X_2) \propto P_{11}P_{22}P_{12}$ 

$$egin{array}{rcl} P_{11} &=& \prod_{i=2}^{n_1} \prod_{j=1}^{i-1} h_{11}(||x_{1i}-x_{1j}||) \ P_{22} &=& \prod_{i=2}^{n_2} \prod_{j=1}^{i-1} h_{22}(||x_{2i}-x_{2j}||) \ P_{12} &=& \prod_{i=1}^{n_1} \prod_{j=1}^{n_2} h_{12}(||x_{1i}-x_{2j}||) \end{array}$$

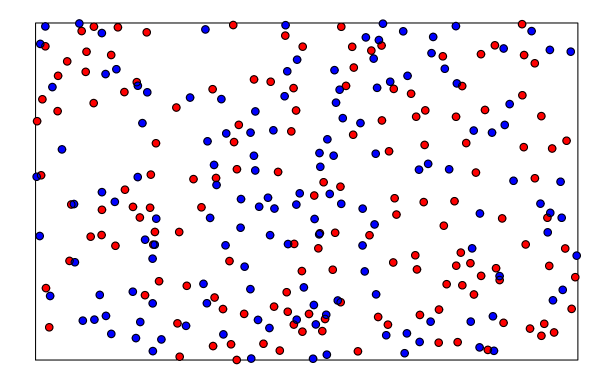
### **Parametric family of interaction functions**

$$h(u; heta) = \left\{egin{array}{ccc} 0 & : & u \leq \delta \ 1 - \exp[-\{(u-\delta)/\phi\}^lpha] & : & u > \delta \end{array}
ight.$$

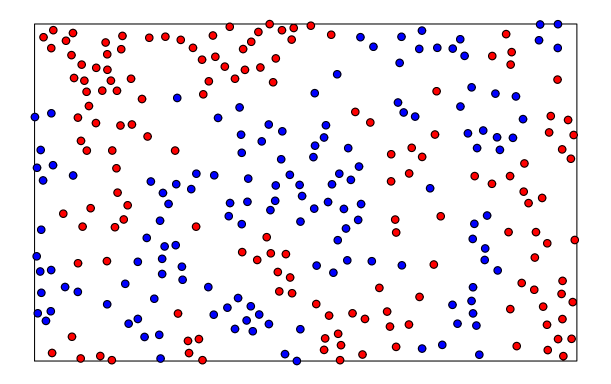
**Special cases** 

- Simple inhibition:  $\phi \to 0$
- Independence:  $h_{12}(u) = 1$
- Functional independence:  $h_{12}(\cdot)$  simple inhibitory

Marginal behaviour depends on  $h_{12}(\cdot)$ 



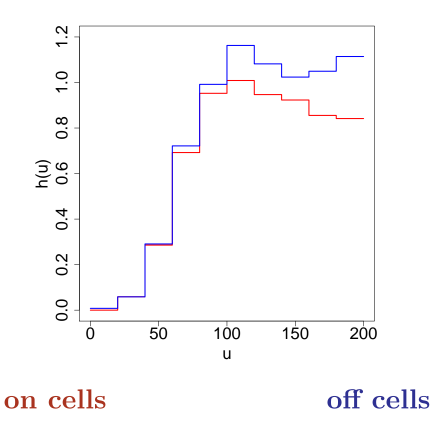
 $\delta_{12} = 0$  (independence)



 $\delta_{12} = 50$  (mutually inhibitory)

## Parametric analysis of the amacrine cells

Non-parametric estimates of h(u) obtained by fitting step-function model using maximum pseudo-likelihood



#### Fitted univariate models

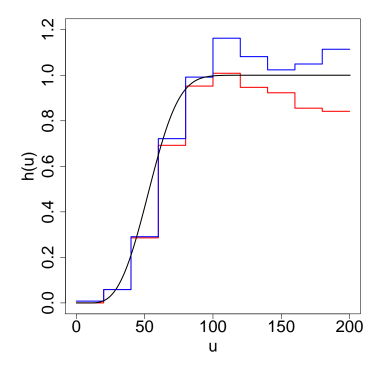
$$h(u; heta) = \left\{egin{array}{ccc} 0 &: & u \leq \delta \ 1 - \exp[-\{(u-\delta)/\phi\}^lpha] &: & u > \delta \end{array}
ight.$$

- Likelihood ratio statistic for common marginal parameters:  $D=1.36\sim\chi^2_2$  p=0.507
- Pooled Monte Carlo MLEs

Parameter	Estimate	Std Error	Correlation
$\phi$	49.08	2.51	
lpha	2.92	0.25	-0.06

Treat  $\delta$  as known (physical size of cells)

#### Goodness-of-fit



A bivariate model for the amacrine cells

Likelihood ratio tests

• statistical independence vs functional independence

$$D = 5.30 \sim \chi_1^2$$
  $p = 0.021$ 

• functional independence vs general bivariate

$$D = 0.30 \sim \chi_2^2$$
  $p = 0.861$ 

• 95% confidence interval for  $\delta_{12}$ 

$$2.3 \le \delta_{12} < 5.0$$

#### Goodness-of-fit

- $\hat{K}_{ij}(s)$  estimate from data
- $\bar{K}_{ij}(s)$  mean of estimates from 99 simulations of model
- three test statistics:

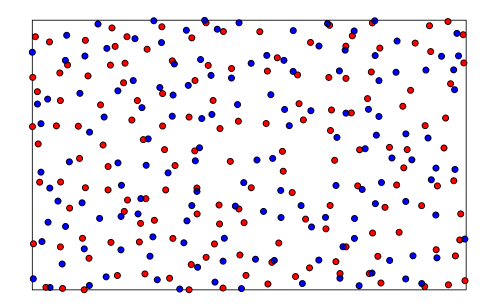
$$T_{ij} = \sum_{s=1}^{150} [\{\hat{K}_{ij}(s) - \bar{K}_i(s)\}/s]^2$$

#### Results

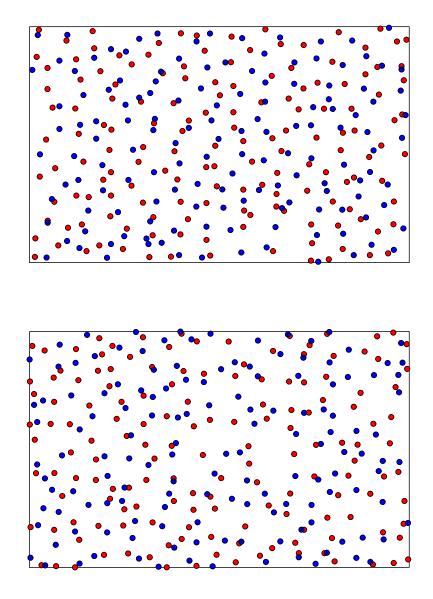
 $T_{11}, \quad p = 0.11 \,\, ({
m on \ cells}) \ T_{22}, \quad p = 0.05 \,\, ({
m off \ cells}) \ T_{12}, \quad p = 0.25 \,\, ({
m dependence})$ 

Bonferroni:  $p \leq 0.15$ 

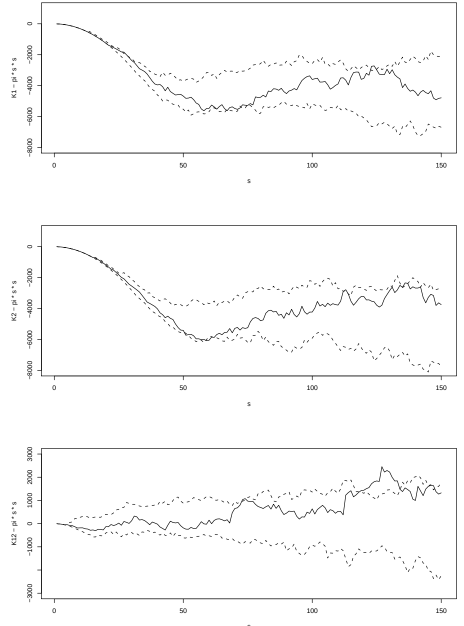
#### Realisation of fitted model, $\delta_{12} = 5$ (functional independence)



#### Data and realisation of fitted model



#### K-functions and simulation envelopes



# 7. Spatio-temporal modelling

- spatial time series
- spatio-temporal point processes
- case-studies

## **Classification of spatio-temporal data?**

Some possiblities:

- geostatistical:  $(x_i, t_i, Y_i) : i = 1, ..., n; (x_i, t_i) \in \mathbb{R}^2 \times \mathbb{R}^+$
- regular lattice:  $Y_{ijt}$ : i = 1, ..., n; j = 1, ..., m; t = 1, ..., T (spatially discrete)
- spatial time series:  $(x_i, Y_{it}) : i = 1, ..., n; t = 1, ..., T$ (spatially discrete or spatially continuous)
- point process:  $(x_i, t_i) : i = 1, ..., n$
- various hybrids

## Spatial time series

$$(Y_{it}, x_i): i = 1, ..., n; t = 1, ..., T$$

- spatially discrete sample from a spatially continuous phenomenon
- a common situation in practice, e.g. environmental monitoring networks
- implicit assumption that data are spatially sparse but temporally dense

### Spatial time series: model specification

- 1. Direct specification:  $\operatorname{Cov}\{Y(x,t),Y(x',t')\}=\sigma^2
  ho(u,v),\ u=||x-x'||,\,v=|t-t'|$ 
  - (a) separable:  $ho(u,v) = 
    ho_s(u)
    ho_t(v)$
  - (b) non-separable:  $\rho(u, v) \neq \rho_s(u)\rho_t(v)$
- 2. Conditioning on the past:
  - $Y_t = \{Y_t(x) : x \in \mathbb{R}^2\}$
  - model  $Y_t$  conditional on  $\{Y_s : s < t\}$

Natural starting point for modelling,

 $[Y_t | \{Y_s : s < t\}] = [Y_t | Y_{t-1}]$ 

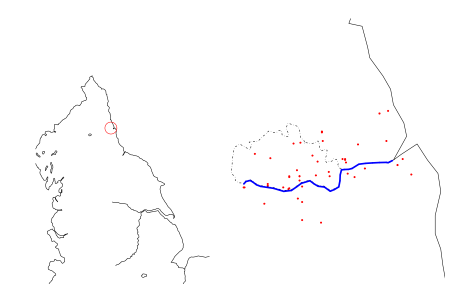
Separability implies that  $[Y_t(x)|Y_{t-1}] = [Y_t(x)|Y_{t-1}(x)]$ 

# The PAMPER study

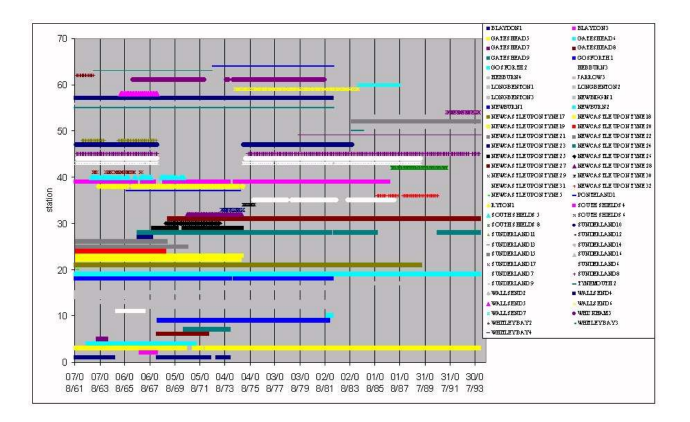
Goal: Construct predictions of black smoke levels, S(x, t), over thirty-year period

Available data:

• monitored black smoke levels from spatially discrete monitoring network



#### • monitors are only active intermittently



# Modelling strategy

Two-stage approach:

- 1. model temporal variation in spatially averaged black smoke levels
- 2. model residual spatio-temporal variation about temporal average

### Model for temporal variation in spatially averaged black smoke

 $Y_t$  = spatially averaged black smoke at time t

Model needs to take account of:

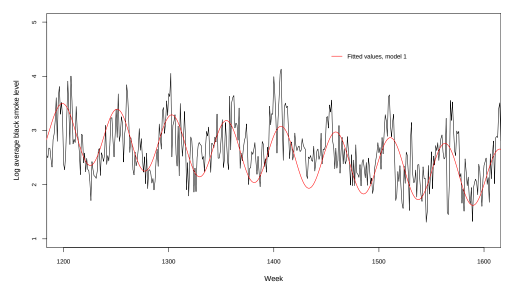
- long-term (decreasing) trend
- seasonal variation

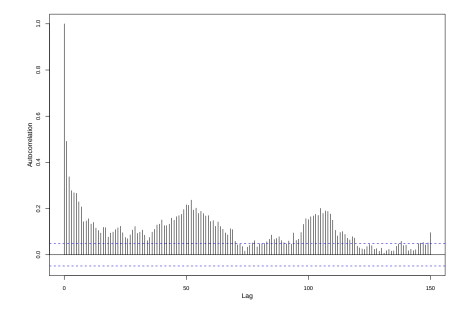
Classical regression model for  $Y_t$  is

$$\log P_t = \alpha + \beta t + \sum_{k=1}^r \{A_k \cos(k\omega t) + B_k \sin(k\omega t)\} + Z_t$$

Case r = 1 gives pure sinusoid, r = 2, 3, ... allows non-sinusoidal seasonal patterns

#### Static fit and residual autocorrelation structure





### Model for temporal variation in spatially averaged black smoke (continued)

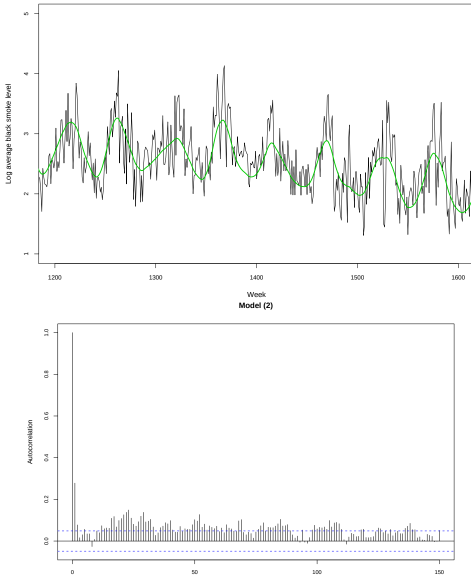
Classical model fails because seasonal pattern is stochastic. Dynamic model:

 $\log P_t = \alpha + \beta t + \{A_t \cos(\omega t) + B_t \sin(\omega t)\} + Z_t$ 

$$egin{array}{rcl} A_t &=& A_{t-1}+\epsilon_t \ B_t &=& B_{t-1}+\delta_t \end{array}$$

Allows locations and magnitudes of seasonal peaks and troughs to vary between years

#### Dymamic fit and residual autocorrelation structure



# Model for spatio-temporal variation in residuals

$$Y_t(x) = \log \hat{P}_t + S(x,t) + Z_t(x)$$

- S(x,t) = spatio-temporally correlated (?) random field
- $Z_t(x)$  = mutually independent measurement errors

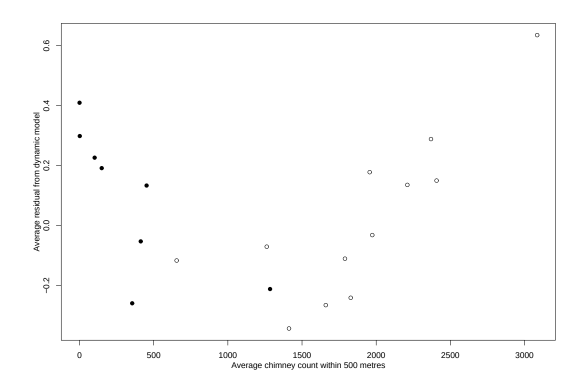
### **Constructed covariates**

- where does the spatio-temporal correlation come from?
- look for possible surrogate measures which:
  - are available at all locations and times
  - correlate well with measured black smoke concentrations at monitored locations

#### Monitored black smoke vs domestic chimney density

Important interactions with:

- non-residential/residential land-use (solid/open circles)
- clean-air act (staggered implementation)



# PAMPER analysis: discussion points

- 1. temporal takes precedence over spatial
- 2. construction of spatially continuous explanatory variables assists prediction of spatio-temporally continuous exposure surface
- 3. and may eliminate residual spatio-temporal correlation

### Spatio-temporal point processes: Cox process models

1. Unobserved stochastic intensity,

 $\Lambda(x,t) = ext{non-negative-valued stochastic process}$ 

2. Conditional on  $\Lambda(x,t) = \lambda(x,t), \forall x,t$ , point process is Poisson with intensity  $\lambda(x,t)$ 

Useful class of models for:

- environmentally driven processes
- aggregated point patterns
- empirical prediction

### Real-time disease surveillance

Data: daily calls to NHS direct Model: log-Gaussian Cox process

$$egin{array}{rcl} \Lambda(x,t)&=&\lambda_0(x)\mu_0(t)\exp\{S(x,t)\}\ S(x,t)&\sim&\mathrm{SGP}\{-0.5\sigma^2,\sigma^2,
ho(u,v)\} \end{array}$$

Goal: real-time mapping of  $P\{S(x,t) > c\}$  for pre-specified c Diggle, Rowlingson and Su (2005)

Animation at www.lancaster.ac.uk/staff/diggle

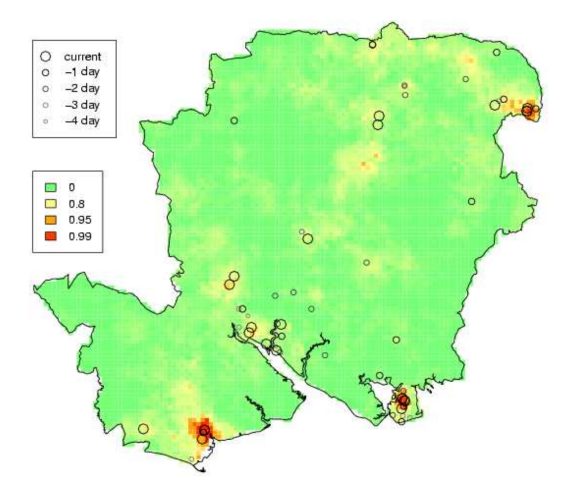
### **Spatial prediction**

- plug-in for estimated model parameters
- MCMC to generate samples from conditional distribution of S(x, t) given data up to time t
- choose critical threshold value c > 1
- map empirical exceedance probabilities,

 $p_t(x) = P\left(\exp\{S(x,t)\} > c|\text{data}\right)$ 

• web-reporting with daily updates

# **Spatial prediction : results for 6 March 2003**



c = 2

### Spatio-temporal point processes: conditional intensity models

 $\mathcal{H}_t = \text{ complete history (locations and times of events)}$ 

 $\lambda(x, t | \mathcal{H}_t) =$  conditional intensity (hazard) for new event at location x, time t, given history  $\mathcal{H}_t$ 

Useful class of models for:

- processes involving interactions amongst events
- aggregated or regular point patterns
- mechanistic modelling

2001 foot-and-mouth epidemic in Cumbria:

www.lancaster.ac.uk/staff/diggle

### Likelihood analysis

Log-likelihood for data  $(x_i, t_i) \in A \times [0, T] : i = 1, ..., n$ , with  $t_1 < t_2 < ... < t_n$ , is

$$L( heta) = \sum_{i=1}^n \log \lambda(x_i, t_i | \mathcal{H}_{t_i}) - \int_0^T \int_A \lambda(x, t | \mathcal{H}_t) dx dt$$

Rarely tractable, but Monte Carlo methods available in special cases (eg log-Gaussian Cox processes)

### Partial likelihood analysis

Data  $(x_i, t_i) \in A \times [0, T] : i = 1, ..., n; t_1 < t_2 < ... < t_n$ 

Condition on locations  $x_i$  and times  $t_i$ Derive log-likelihood for observed ordering 1, 2, ..., n

Need to distinguish between:

- Spatially discrete set of potential points
- Spatially continuous set of potential points

### **Partial Likelihood Formulation**

- Condition on the locations  $x_i$  and times  $t_i$
- $\mathcal{R}_i$ : the risk set at time  $t_i$
- Partial log-likelihood  $L_p(\theta) = \sum_{i=1}^n \log p_i$
- Spatially discrete  $\rightarrow \mathcal{R}_i = \{i, i+1, ..., n\}$

$$p_i = rac{\lambda(x_i, t_i | \mathcal{H}_{t_i})}{\sum_{j \geq i} \lambda(x_j, t_i | \mathcal{H}_{t_i})}$$

• Spatially continuous  $\rightarrow \mathcal{R}_i \equiv A$ 

$$p_i = rac{\lambda(x_i, t_i | \mathcal{H}_{t_i})}{\int_A \lambda(x, t_i | \mathcal{H}_{t_i}) dx}$$

# The 2001 UK FMD epidemic

- First confirmed case 20 February 2001
- Approximately 140,000 at-risk farms in the UK (cattle and/or sheep)
- Outbreaks in 44 counties, epidemic particularly severe in Cumbria and Devon
- Last confirmed case 30 September 2001
- Consequences included:
  - more than 6 million animals slaughtered (4 million for disease control, 2 million for "welfare reasons")
  - estimated direct cost  $\pounds 8$  billion

# Progress of the epidemic in Cumbria

• Animation

# Progress of the epidemic in Cumbria

#### • Animation

- predominant pattern is of transmission between near-neighbouring farms
- but also some apparently spontaneous outbreaks?
- qualitatively similar pattern in Devon

# Questions

- What factors affected the spread of the epidemic?
- How effective were control strategies in limiting the spread?

# A model for the FMD epidemic (after Keeling et al, 2001)

Notation

- $\mathcal{H}_t = \text{history of process up to } t -$
- $\lambda(x, t | \mathcal{H}_t) = \text{conditional intensity}$
- $\lambda_{jk}(t) = \text{rate of transmission from farm } j$  to farm k

Farm-specific covariates for farm i

- $n_{1i}$  = number of cows
- $n_{2i}$  = number of sheep

Transmission kernel

$$f(u) = \exp\{-(u/\phi)^\kappa\} + \rho$$

At-risk indicator for transmission of infection

 $I_{jk}(t) = 1$  if farm k not infected and not slaughtered by time t, and farm j infected and not slaughtered by time t

**Reporting delay** 

Simplest assumption is that reporting date is infection date plus  $\tau$  (latent period of disease plus reporting delay if any)

**Resulting statistical model** 

$$\lambda_{jk}(t) = \lambda_0(t) A_j B_k f(||x_j - x_k||) I_{jk}(t)$$

$$\lambda_0(t)= ext{ arbitrary}$$

$$A_j = (\alpha n_{1j} + n_{2j})$$

$$B_k = (\beta n_{1k} + n_{2k})$$

#### Fitting the model

• rate of infection for farm k at time t is

$$\lambda_k(t) = \sum_j \lambda_{jk}(t)$$

• partial likelihood contribution from ith case is

$$p_i = \lambda_i(t_i) / \sum_k \lambda_k(t_i)$$

• fix  $\tau = 5$ ,  $\kappa = 0.5$ , estimate remaining parameters by maximising partial likelihood

# **FMD** results

**Common parameter values in Cumbria and Devon?** 

Likelihood ratio test:  $\chi_4^2 = 2.98$ 

Parameter estimates

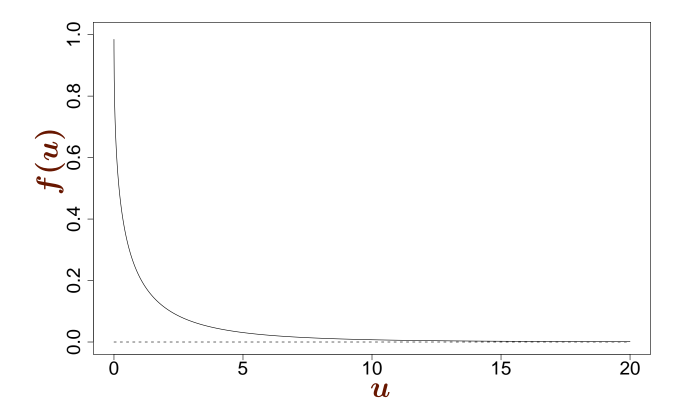
$$(\hat{lpha},\hat{eta},\hat{\phi},\hat{
ho})=(4.92,30.68,0.39,9.9 imes10^{-5})$$

But note that likelihood ratio test rejects  $\rho = 0$ .

#### **Standard errors**

Available via usual asymptotic argument, but numerical estimates of information matrix unreliable?

# Fitted transmission kernel



Qualitatively similar to estimate given in Keeling et al (2001)

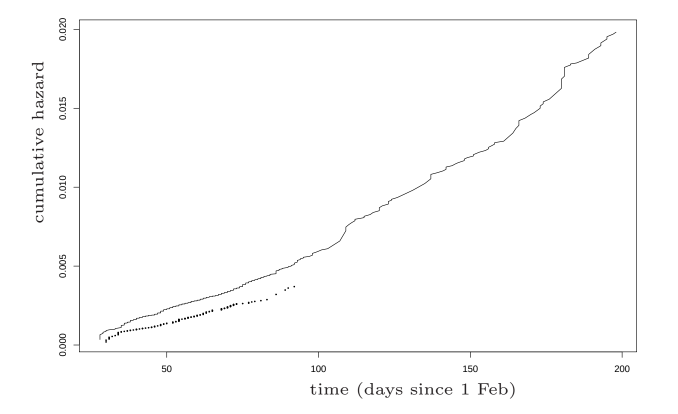
# Estimating $\lambda_0(t)$

 $egin{aligned} \lambda_{ij}(t) &=& \lambda_0(t)
ho_{ij}(t) \ && 
ho(t) &=& \sum_i \sum_j I_{ij}(t)
ho_{ij}(t) \ && \Lambda(t) &=& \int_0^t \lambda_0(u) du \end{aligned}$ 

Nelson-Aalen estimator

$$\hat{\Lambda}_0(t) = \int_0^t \hat{
ho}(u)^{-1} dN(u) = \sum_{i:t_i \le t} \hat{
ho}(t_i)^{-1}$$

#### Nelson-Aalen estimates for Cumbria (solid line) and Devon (dotted line)

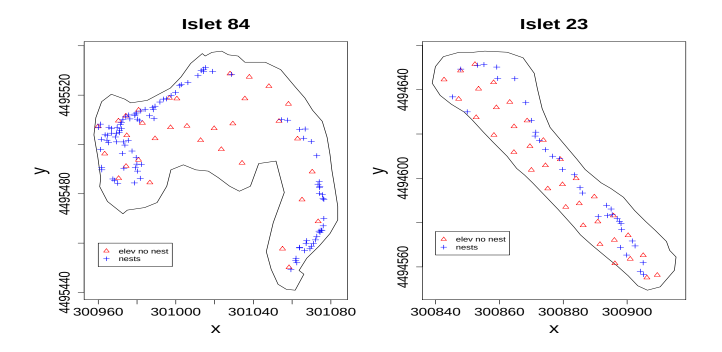


**Diggle**, **2006** 

# Nesting colonies of common terns



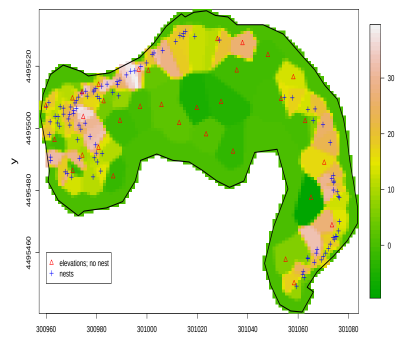
#### Islets 23 and 84



Coast boundaries (-), spatial locations of the nests (+), and other locations for which elevation is recorded  $(\triangle)$  for islets 84 (left panel) and 23 (right panel)

Approximation of elevation surface

Approximate elevation surface z(x) for islet 84 based on all available elevations and assuming piece-wise constant z(x) within Voronoi tiles



#### **Conditional intensity**

 $\lambda(\mathbf{x}, t | \mathcal{H}_t) = \lambda_0(t) \exp\{\beta z(\mathbf{x})\}g(\mathbf{x}, t_i | \mathcal{H}_t)$ 

- $g(\mathbf{x}, t | \mathcal{H}_t)$  models dependence on locations of earlier nests
- $\beta z(\mathbf{x})$  models log-linear effect of elevation

#### Two models for $g(\cdot)$

• 
$$\mathcal{M}_1$$
:  
 $g(\mathbf{x}, t | \mathcal{H}_t) = h\left(\min_{j: t_j < t} (||\mathbf{x}_j - \mathbf{x}||)\right)$   
•  $\mathcal{M}_2$ :  
 $g(\mathbf{x}, t | \mathcal{H}_t) = \prod_{j: t_j < t} h(||\mathbf{x} - \mathbf{x}_j||)$ 

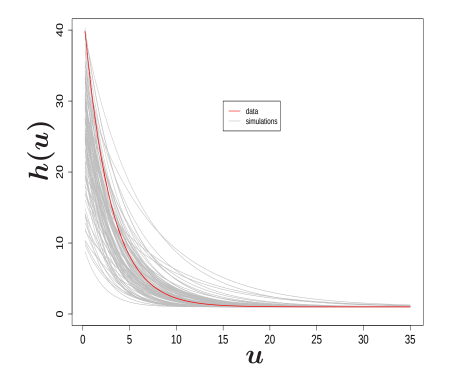
$$h(u) = \left\{egin{array}{cc} 0, & u \leq d_0 \ 1 + heta \exp\left(-rac{u-d_0}{\phi}
ight), & u > d_0 \end{array}
ight.$$

#### Results

- likelihood ratio tests favour model  $\mathcal{M}_1$
- highly significant effect of elevation  $\hat{\beta} = 0.05, SE = 0.0006, p << 0.001$

### Monte Carlo interval estimation

Envelope of estimates  $\hat{h}(u)$  from 99 simulations of fitted model



Diggle, Kaimi and Abellana, 2010

# Conclusions

- spatio-temporal data-sets becoming widely available
- different problems require different modelling strategies
- temporal should often take precedence over spatial
- routine implementation is an important consideration when exploring many different models