

Session 9.2: Spatio-temporal modelling for area data

Bayesian modelling for Spatial and Spatio-temporal data, Imperial College London

Learning Objectives

At the end of this session you should be able to:

- Explain how to extend spatial or temporal models to spatio-temporal models
- Fit Bayesian space-time models using R-INLA

The topics covered in this lecture can be found in Chapter 7 of the book **Spatial and Spatio-Temporal Bayesian Models with R-INLA**.

Outline

1. Temporal dependence
2. From space to space-time
3. Type of interactions

Temporal dependence

Temporal dependence

- Similarly to spatial dependence, it is sometimes necessary to model temporal dependence of data or of parameters:
- the weekly or monthly number of cases of many diseases exhibit often a seasonal pattern as well as short term dependence
- the underlying daily level of atmospheric pollutants, e.g. PM₁₀, will show strong correlation over consecutive days because their lifetime lasts over several days
- To the contrary of spatial models, there is a natural order to any time series data which is used in specifying the models.

Spatial patterns

1. Data

- Disease counts y_i in area i and stratum k , aggregated over a time period, $i = 1, \dots, N, k = 1, \dots, K$
- Population counts n_{ik} in area i and stratum k , aggregated over a time period
- Expected numbers $E_i = \sum_k n_{ik} r_k$, where r_k reference rate for stratum (age, gender,...)

2. Spatial smoothing using BYM model

$$y_i \sim \text{Poisson}(E_i \rho_i); \quad i = 1, \dots, N$$

$$\log \rho_i = b_0 + v_i + u_i$$

$$v_i \sim \text{Normal}(0, \sigma_v^2)$$

$$\mathbf{u} \sim \text{ICAR}(\mathbf{W}, \sigma_u^2) \rightarrow u_i | u_j, \quad j \neq i \sim \text{Normal} \left(\frac{\sum_j w_{ij} u_j}{\sum_j w_{ij}}, \sigma_u^2 / n_i \right)$$

with $w_{ij} = 1$ if areas i and j are neighbours, 0 otherwise

Temporal trends

1. Data

- Disease counts y_{tk} in time period t and stratum k , aggregated over space, $t = 1, \dots, T$ (equally-spaced time intervals), $k = 1, \dots, K$
- Population counts n_{tk} in time period t and stratum k , aggregated over space
- Expected numbers $E_t = \sum_k n_{tk} r_k$, where r_k reference rate for stratum (age, gender,...)

2. Temporal trends

$$y_t \sim \text{Poisson}(E_t \rho_t); \quad t = 1, \dots, T$$
$$\log \rho_t = ???$$

Temporal trends

1. Data

- Disease counts y_{tk} in time period t and stratum k , aggregated over space, $t = 1, \dots, T$ (equally-spaced time intervals), $k = 1, \dots, K$
- Population counts n_{tk} in time period t and stratum k , aggregated over space
- Expected numbers $E_t = \sum_k n_{tk} r_k$, where r_k reference rate for stratum (age, gender,...)

2. Temporal trends

$$y_t \sim \text{Poisson}(E_t \rho_t); \quad t = 1, \dots, T$$
$$\log \rho_t = b_0 + \beta t \quad \text{simple linear regression}$$

Temporal trends

1. Data

- Disease counts y_{tk} in time period t and stratum k , aggregated over space, $t = 1, \dots, T$ (equally-spaced time intervals), $k = 1, \dots, K$
- Population counts n_{tk} in time period t and stratum k , aggregated over space
- Expected numbers $E_t = \sum_k n_{tk} r_k$, where r_k reference rate for stratum (age, gender,...)

2. Temporal trends

$$\begin{aligned} y_t &\sim \text{Poisson}(E_t \rho_t); \quad t = 1, \dots, T \\ \log \rho_t &= b_0 + \beta t \quad \text{simple linear regression} \\ &= b_0 + \psi_t \quad \text{global temporal smoothing} \\ \psi_t &\sim \text{Normal}(0, \sigma_\psi^2) \end{aligned}$$

Temporal trends

1. Data

- Disease counts y_{tk} in time period t and stratum k , aggregated over space, $t = 1, \dots, T$ (equally-spaced time intervals), $k = 1, \dots, K$
- Population counts n_{tk} in time period t and stratum k , aggregated over space
- Expected numbers $E_t = \sum_k n_{tk} r_k$, where r_k reference rate for stratum (age, gender,...)

2. Temporal trends

$$y_t \sim \text{Poisson}(E_t \rho_t); \quad t = 1, \dots, T$$

$$\log \rho_t = b_0 + \beta t \quad \text{simple linear regression}$$

$$= b_0 + \psi_t \quad \text{global temporal smoothing}$$

$$\psi_t \sim \text{Normal}(0, \sigma_\psi^2)$$

$$= b_0 + \gamma_t \quad \text{local temporal smoothing}$$

$$\gamma_t \sim \text{distribution ?}$$

Temporal trends

1. Data

- Disease counts y_{tk} in time period t and stratum k , aggregated over space, $t = 1, \dots, T$ (equally-spaced time intervals), $k = 1, \dots, K$
- Population counts n_{tk} in time period t and stratum k , aggregated over space
- Expected numbers $E_t = \sum_k n_{tk} r_k$, where r_k reference rate for stratum (age, gender,...)

2. Temporal trends

$$y_t \sim \text{Poisson}(E_t \rho_t); \quad t = 1, \dots, T$$

$$\log \rho_t = b_0 + \beta t \quad \text{simple linear regression}$$

$$= b_0 + \psi_t \quad \text{global temporal smoothing}$$

$$\psi_t \sim \text{Normal}(0, \sigma_\psi^2)$$

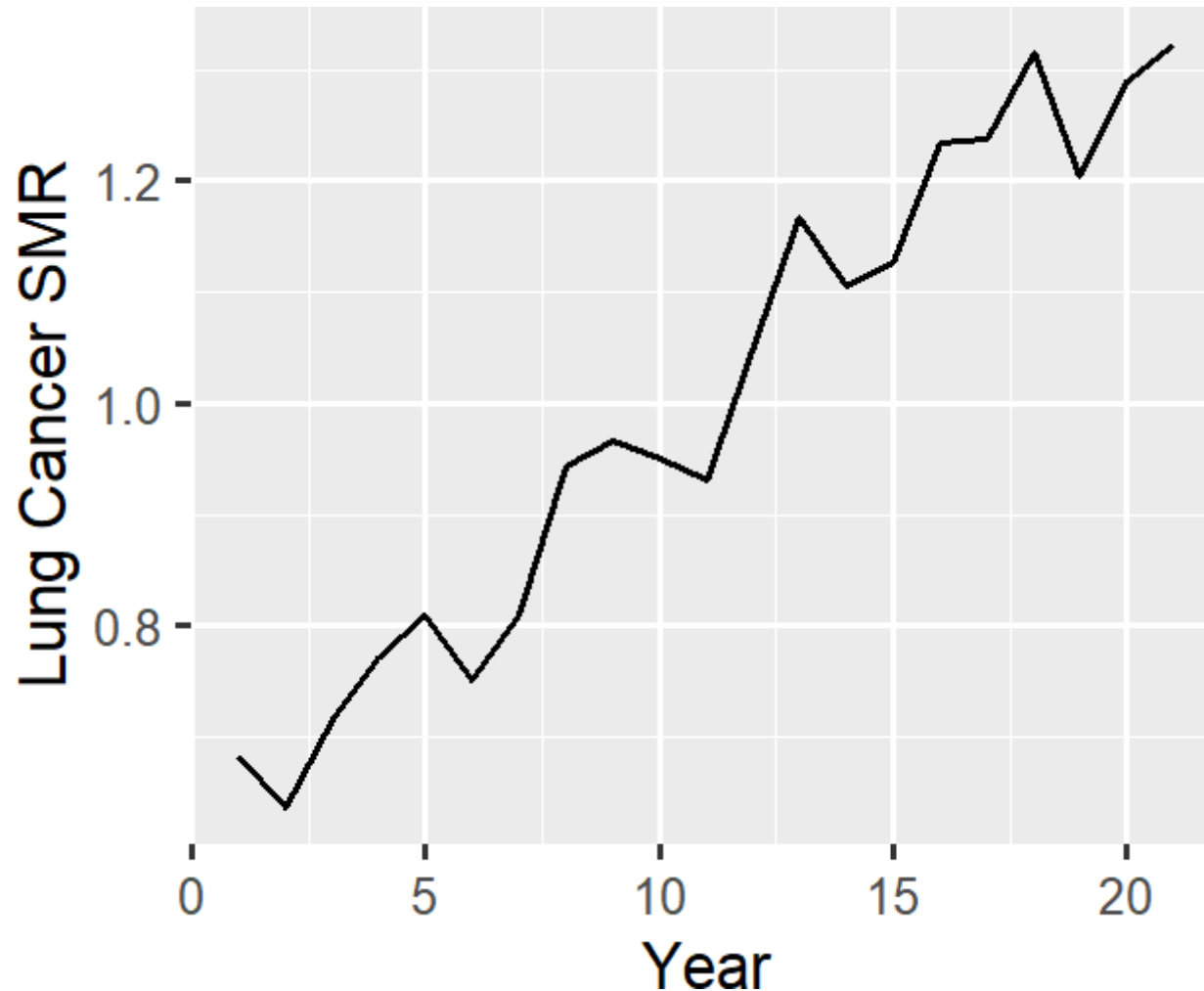
$$= b_0 + \gamma_t \quad \text{local temporal smoothing}$$

$$\gamma_t \sim \text{distribution ?}$$

$$= b_0 + \gamma_t + \psi_t \quad \text{global and local temporal smoothing}$$

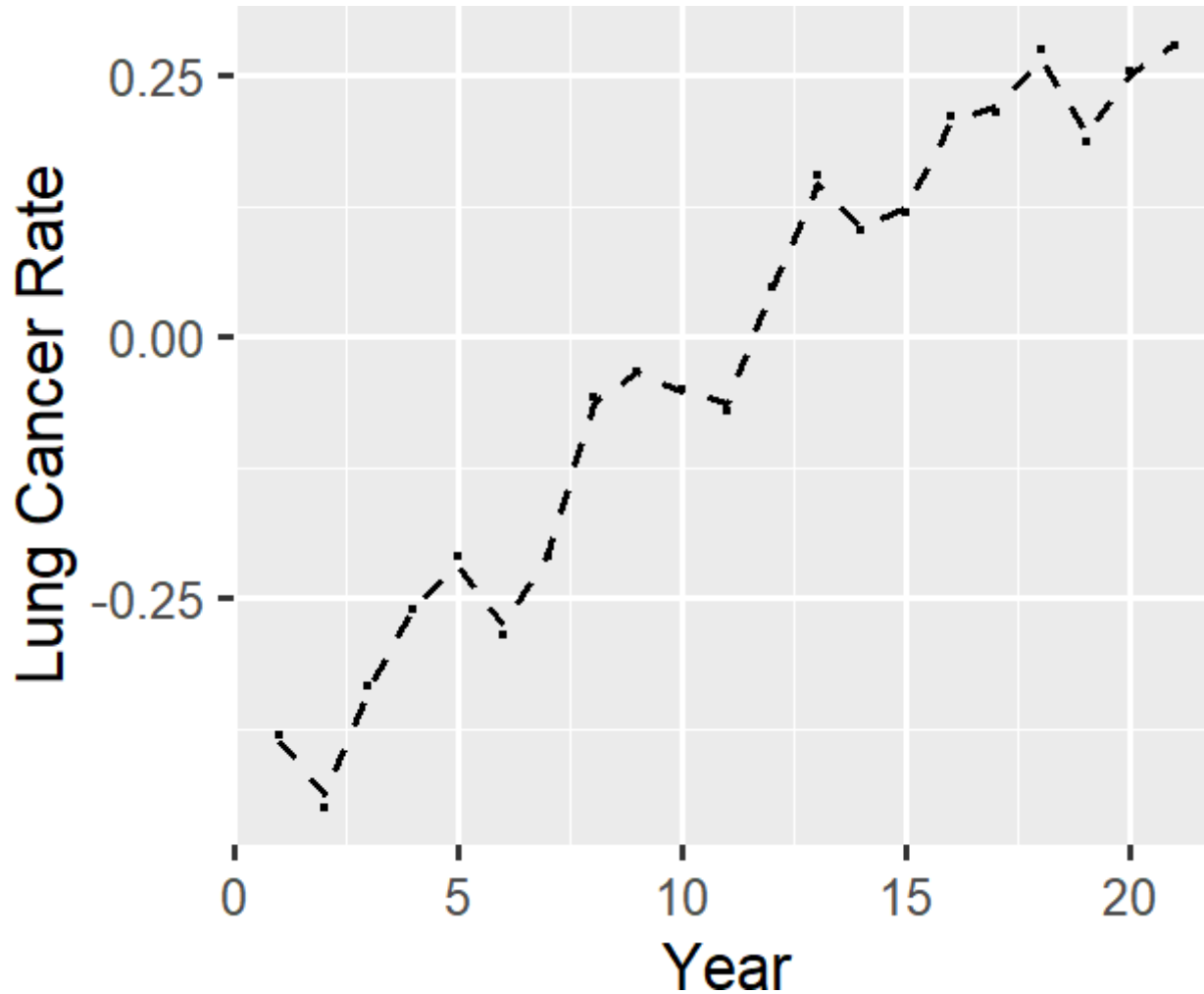
Ohio Lung cancer data

- Data on lung cancer in 88 counties of Ohio, 1968-1988
- Annual rates adjusted by gender and race

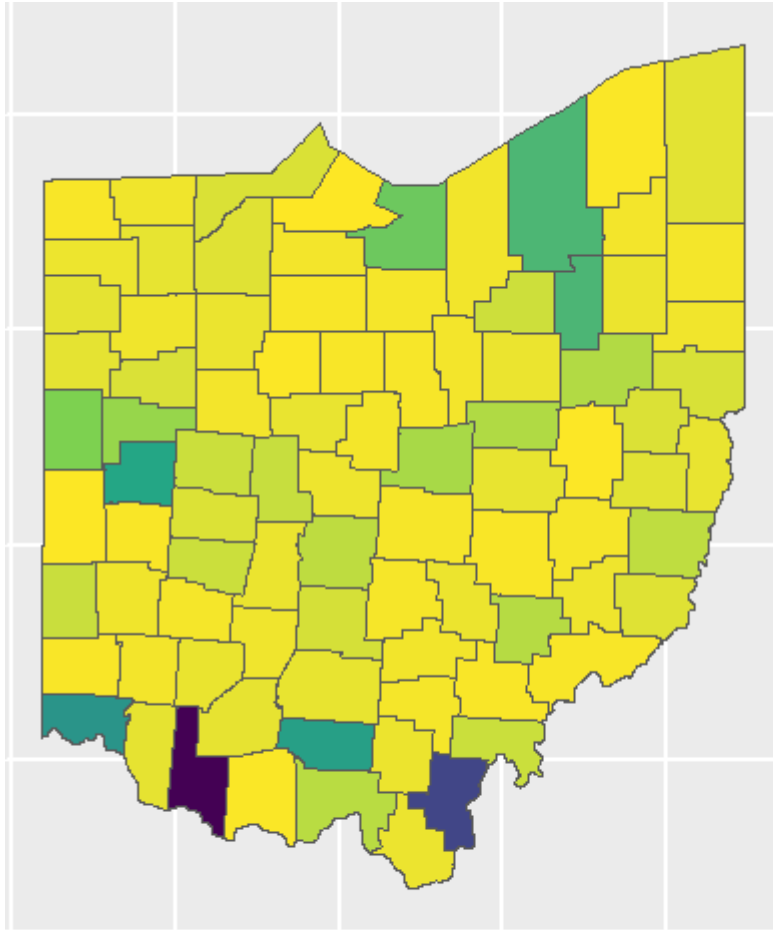


Ohio Lung cancer data: modelled temporal trend

- Smoothed temporal trend using random walk models (posterior mean) seen in the previous lecture

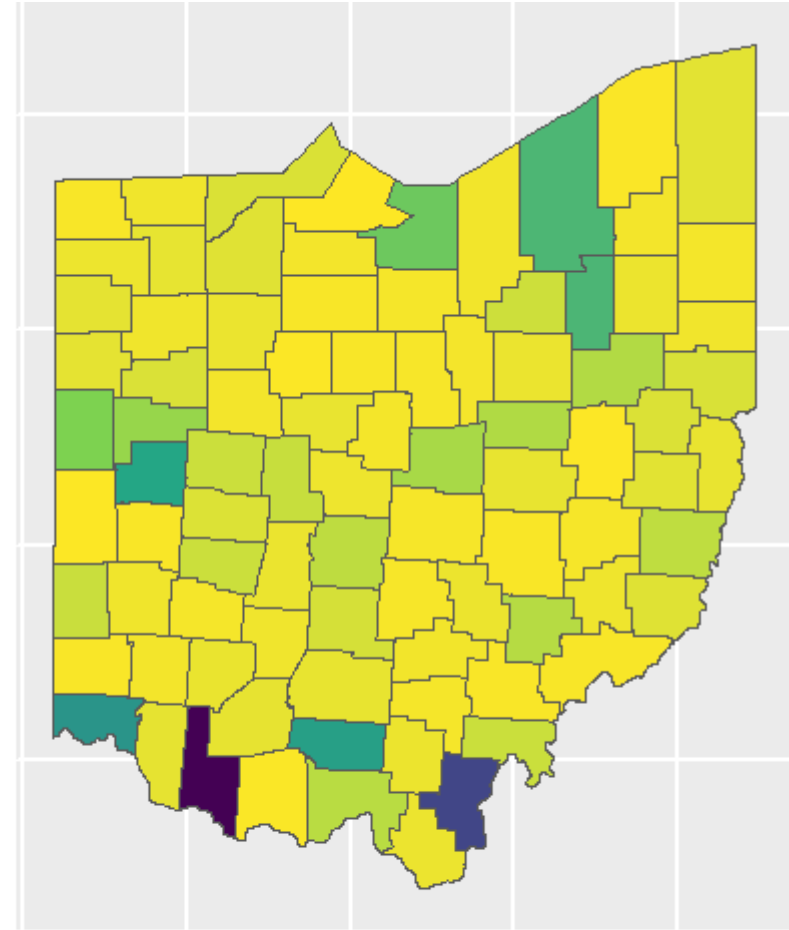


Ohio Lung cancer spatial pattern (no temporal dimension)



SMR

3	6	9	12
---	---	---	----



p_i

3	6	9	12
---	---	---	----

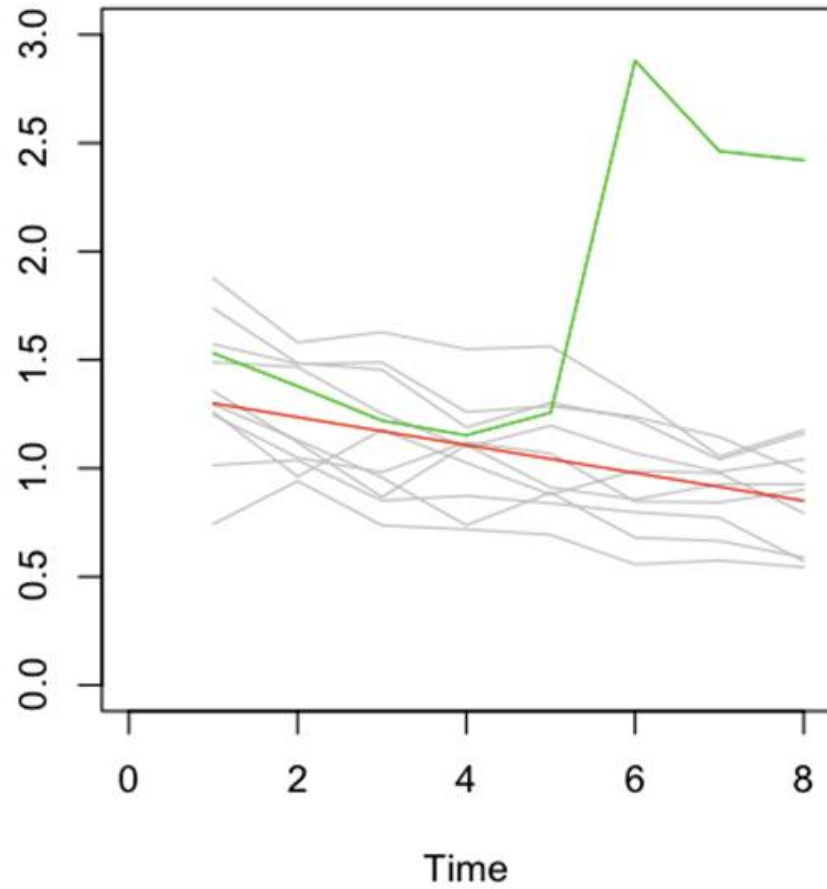
From space to space-time

Disease mapping: Extending space to space-time

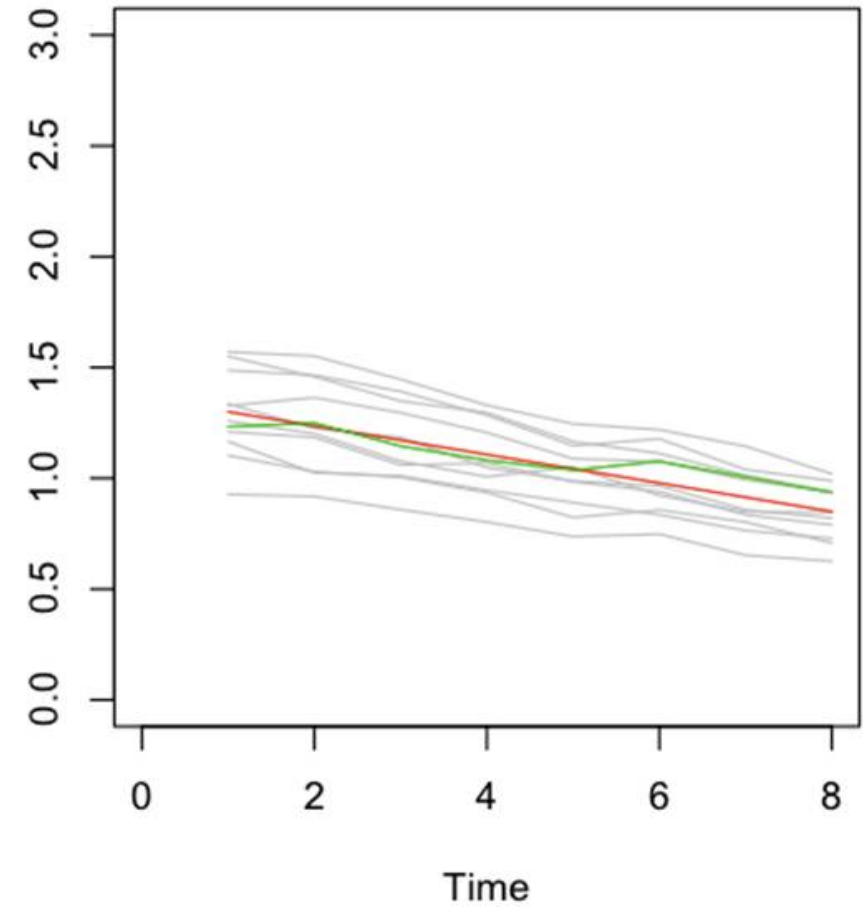
- Disease mapping is usually carried out on aggregated data over a time period
- Rather than suppressing the time dimension, it can be interesting to use models that combine the space and time dimension
- The stability (or not) of the spatial pattern can aid interpretation
- The specific space-time components of the model can potentially pinpoint unusual/emerging hazards
- Data:
 - y_{it} and
 - E_{it} : the observed and expected number of cases in area i at time t calculated as $E_{it} = \sum_k n_{itk} r_k$, where r_k reference rate for stratum (age, gender,...)

Schematic representation I

Raw SMR



Smoothed RR



Linear spatio-temporal model

- A simple model assumes a linear effect of time:

$$\log \rho_{it} = b_0 + u_i + v_i + \beta \times t$$

- Main spatial effect $u_i + v_i$
- Main linear trend β (global time effect)

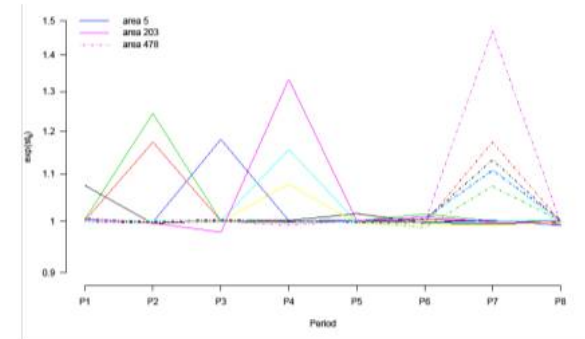
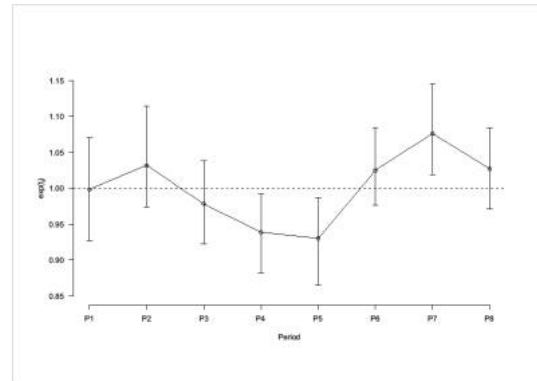
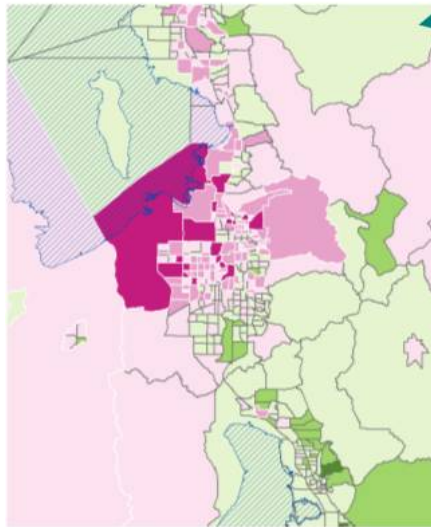
- A differential effect δ_i can be added

$$\log \rho_{it} = b_0 + u_i + v_i + (\beta + \delta_i) \times t$$

- If $\delta_i < 0$ then the area-specific trend is less steep than the mean trend, whilst $\delta_i > 0$ implies that the area-specific trend is steeper than the mean trend.

Noise model: Poisson/Binomial

Latent structure: Space + Time + Interactions



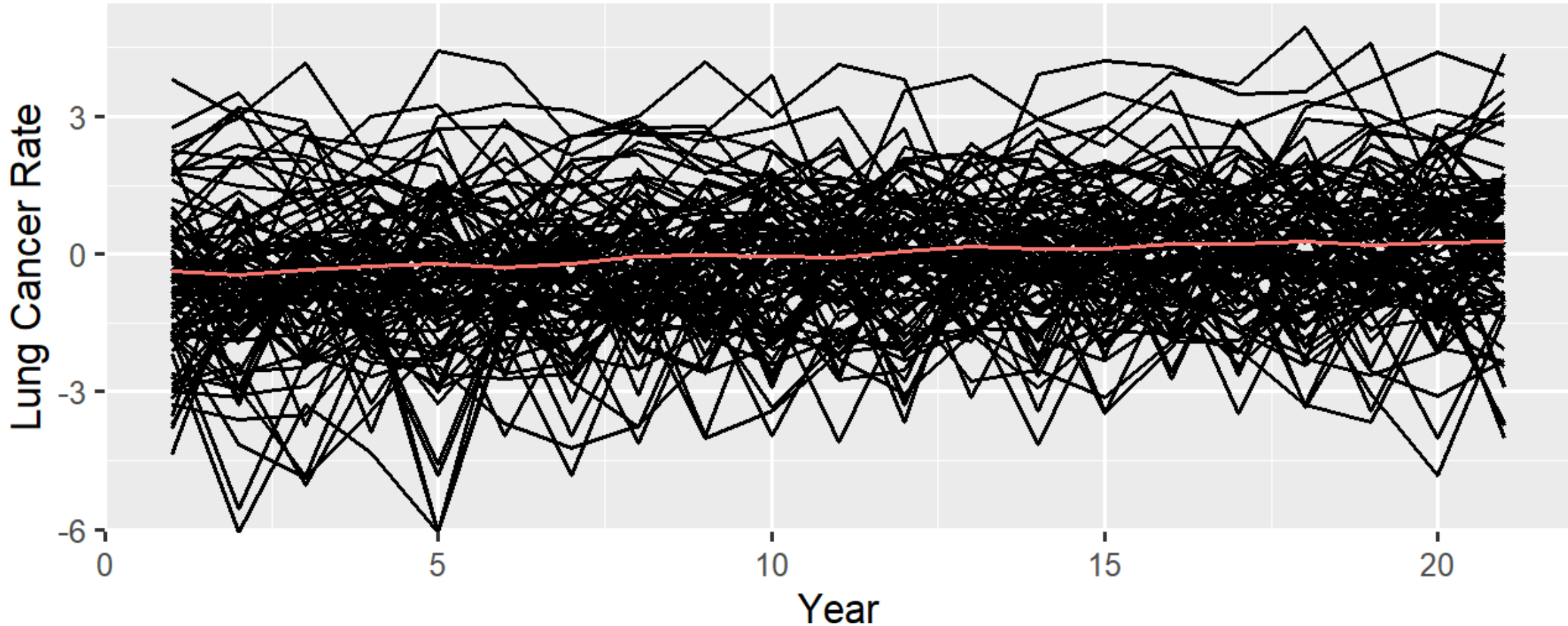
joint Bayesian estimation

Inference

Ohio Lung cancer - temporal SMRs

Log SMR time trends in each county

- Slightly increasing trend but lots of variation across the counties



Simple linear spatio-temporal model (Model 1)

$$y_{it} \sim \text{Poisson}(E_{it}\rho_{it})$$
$$\log \rho_{it} = b_0 + \beta * t + u_i + v_i$$

where

- b_0 overall log RR in Ohio over the 21-year period
- $v_i \sim \text{Normal}(0, \sigma_v^2)$ spatially unstructured RE
- $\mathbf{u} \sim \text{ICAR}(\mathbf{W}, \sigma_u^2)$ spatially structured RE
- $\exp(\beta)$ is the change in the RR associated with a 1-year increase in time

```
> formula.mod1 = y ~ 1 + f(county,model="bym",  
+                           graph=Ohio.adj) + year
```

Log-linear temporal model with unstructured temporal RE (Model 2)

$$y_{it} \sim \text{Poisson}(E_{it}\rho_{it})$$
$$\log \rho_{it} = b_0 + \beta * t + u_i + v_i + \psi_t$$

where

- b_0 overall log RR in Ohio over the 21-year period
- $v_i \sim \text{Normal}(0, \sigma_v^2)$ spatially unstructured RE
- $\mathbf{u} \sim \text{ICAR}(\mathbf{W}, \sigma_u^2)$ spatially structured RE
- $\exp(\beta)$ is the change in the RR associated with a 1-year increase in time
- $\psi_t \sim \text{Normal}(0, \sigma_\psi^2)$ temporally unstructured RE

```
> year2 = ohio.data$year
> formula.mod2 = y ~ 1 + f(county,model="bym",
+                          graph=ohio.adj) +
+                          f(year2,model="iid") +
+                          year
```

Ohio lung cancer - comparison models 1 and 2

```
> mod1 = inla(data=ohio.data,formula=formula.mod1, E=E, family="poisson")
> mod2 = inla(data=ohio.data,formula=formula.mod2, E=E, family="poisson")
> # Posterior mean and 95% CI for Model 1
> mod1$summary.fixed
```

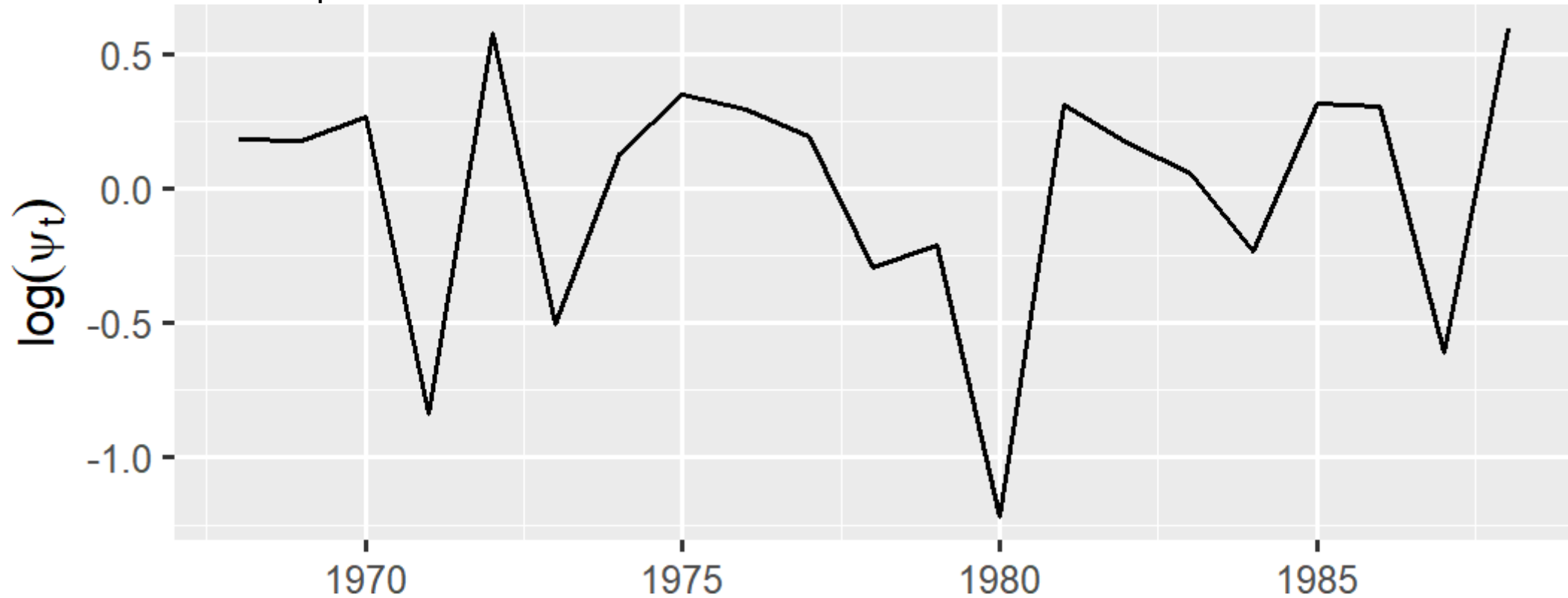
	mean	sd	0.025quant	0.5quant	0.975quant	mode	kld
(Intercept)	-0.98839017	0.1198803206	-1.22400175	-0.98837507	-0.75286303	-0.98834459	2.312841e-09
year	0.02612126	0.0005862819	0.02497147	0.02612126	0.02727105	0.02612126	5.527797e-11

```
> #Posterior mean and 95% CI for Model 2
> mod2$summary.fixed
```

	mean	sd	0.025quant	0.5quant	0.975quant	mode	kld
(Intercept)	-1.00051649	0.24837671	-1.49021595	-1.00050788	-0.5108657	-1.0004908	1.127221e-08
year	0.02042516	0.01702402	-0.01327892	0.02042535	0.0541282	0.0204257	1.358019e-08

Ohio lung cancer - comparison models 1 and 2

Let's look at the temporal random effect in model 2



Some temporal structure which suggests that the linear term is not capturing the temporal trend well

Simple additive space-time structure version 1 (Model 3)

This model assumes that the space time variations can be captured by the superimposition of a BYM spatial model and a structured time trend

$$y_{it} \sim \text{Poisson}(E_{it}\rho_{it})$$
$$\log \rho_{it} = b_0 + b_i + \gamma_t$$

- $b_i = v_i + u_i$
- $v_i \sim \text{Normal}(0, \sigma_v^2)$ spatially unstructured RE
- $\mathbf{u} \sim \text{ICAR}(\mathbf{W}, \sigma_u^2)$ spatially structured RE
- $\gamma_t \sim \text{RW}(1)$ temporally structured RE with variance parameter σ_γ^2
- Assuming a BYM2 specification, there are 3 hyperparameters
 - τ_b , the marginal precision of b_i
 - ϕ , the proportion of spatial variability explained by the spatially structured component
 - σ_γ^2 , the conditional variance of the RW(1) modelling the time trend

```
> formula.mod3 = y ~ 1 + f(county,model="bym",  
+                          graph=Ohio.adj) +  
+                          f(year,model="rw1")
```

```
> mod3 = inla(data=ohio.data,formula=formula.mod3, E=E, family="poisson")
```

Simple additive space-time structure version 2 (Model 4)

$$y_{it} \sim \text{Poisson}(E_{it}\rho_{it})$$
$$\log\rho_{it} = b_0 + u_i + v_i + \psi_t + \gamma_t$$

where

- b_0 overall log RR in Ohio over the 21-year period
- $v_i \sim \text{Normal}(0, \sigma_v^2)$ spatially unstructured RE
- $\mathbf{u} \sim \text{ICAR}(\mathbf{W}, \sigma_u^2)$ spatially structured RE
- $\gamma_t \sim \text{RW}(1)$ temporally structured RE with variance parameter σ_γ^2
- $\psi_t \sim \text{Normal}(0, \sigma_\psi^2)$ temporally unstructured RE

```
> formula.mod4 = y ~ 1 + f(county,model="bym",  
+                       graph=Ohio.adj) +  
+                       f(year,model="rw1") +  
+                       f(year2,model="iid")
```

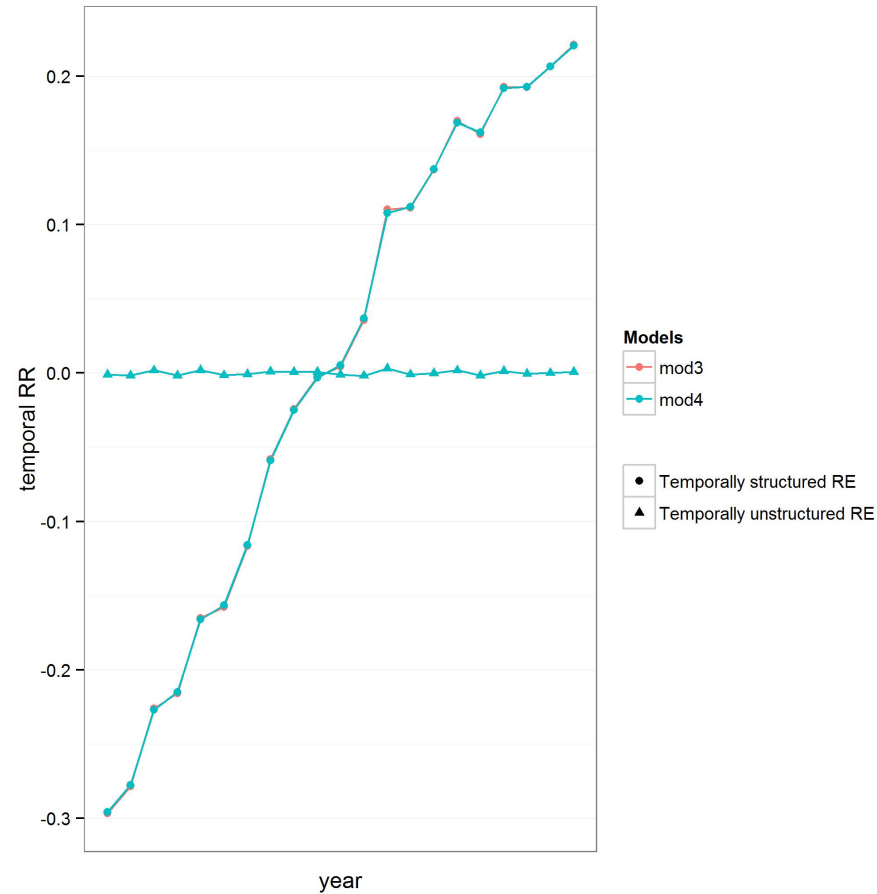
```
> mod4 = inla(data=ohio.data,formula=formula.mod4, E=E, family="poisson")
```

Ohio lung cancer - comparison models 3 and 4

Posterior means of the temporal RE

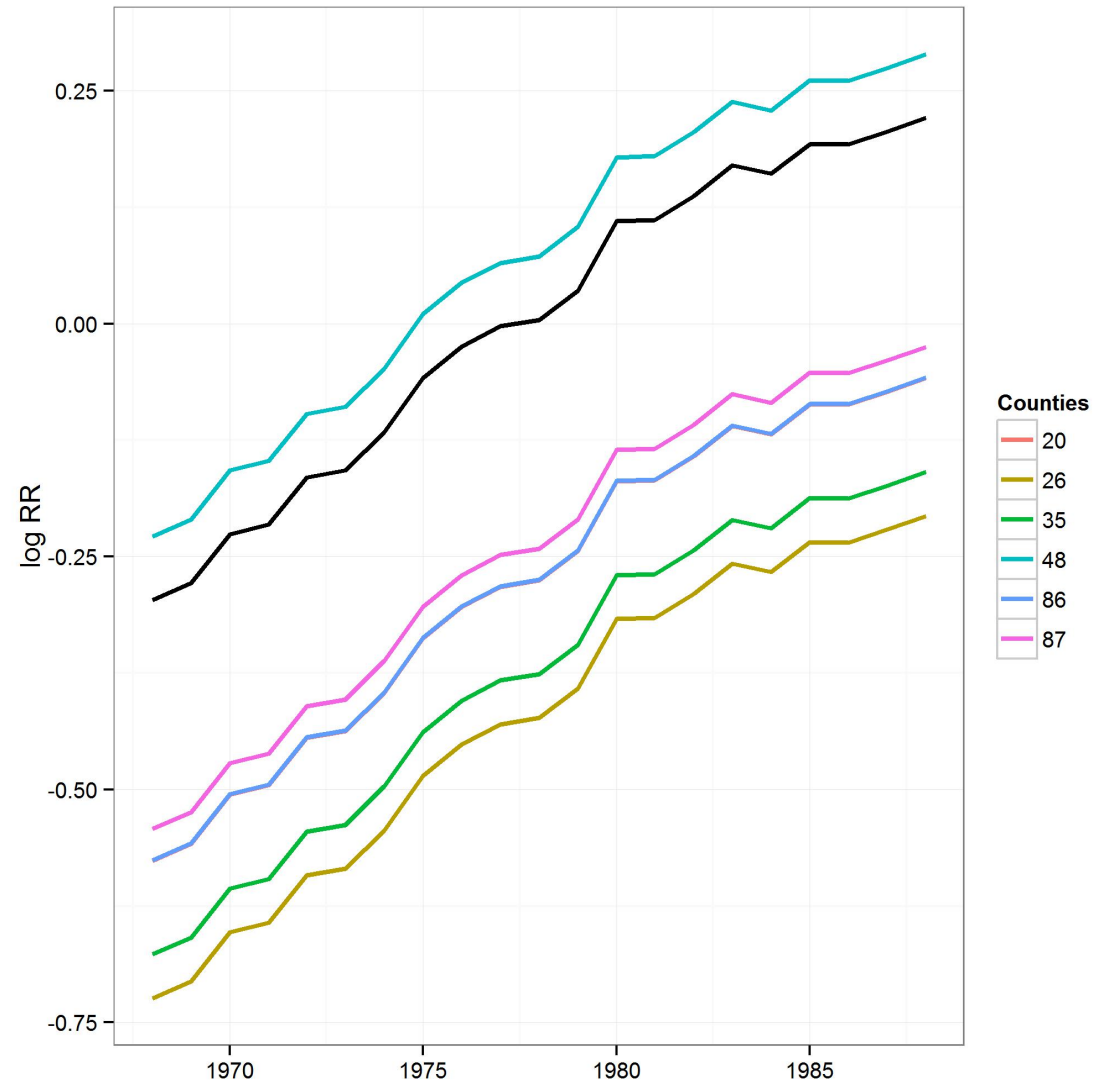
Model 3 = γ_t

Model 4 = $\psi_t + \gamma_t$



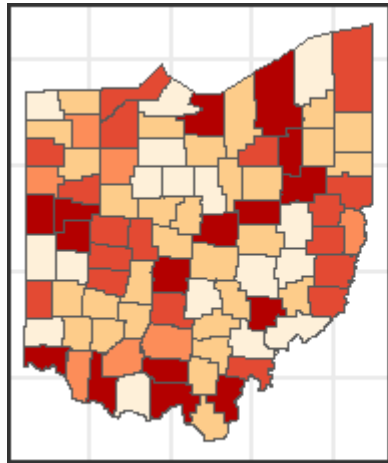
Ohio lung cancer - area-specific temporal RR

Posterior means of the temporal RE γ_t (model 3)

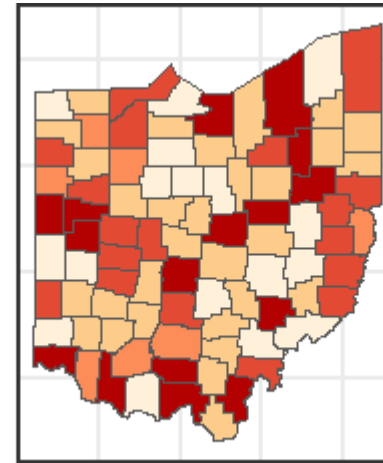
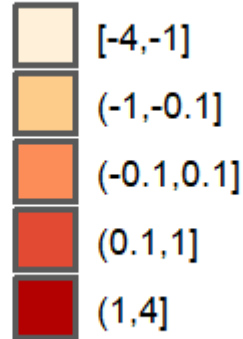


Ohio lung cancer - spatial patterns (all models)

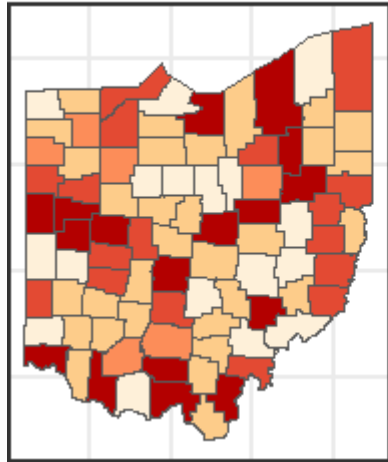
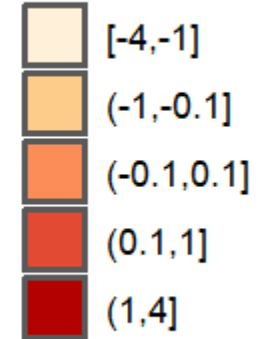
Posterior means of the spatial random effects for the 4 different models



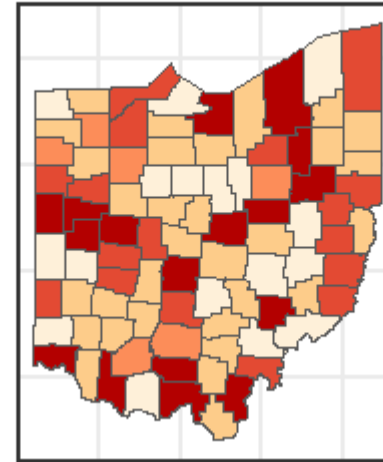
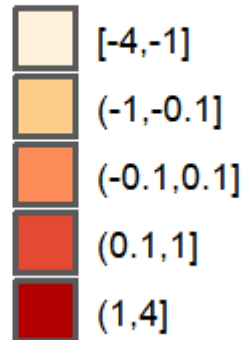
model 1



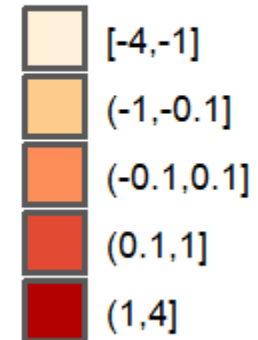
model 2



model 3



model 4



Similar patterns with the 4 models

Space-time model with exchangeable interactions (Model 5)

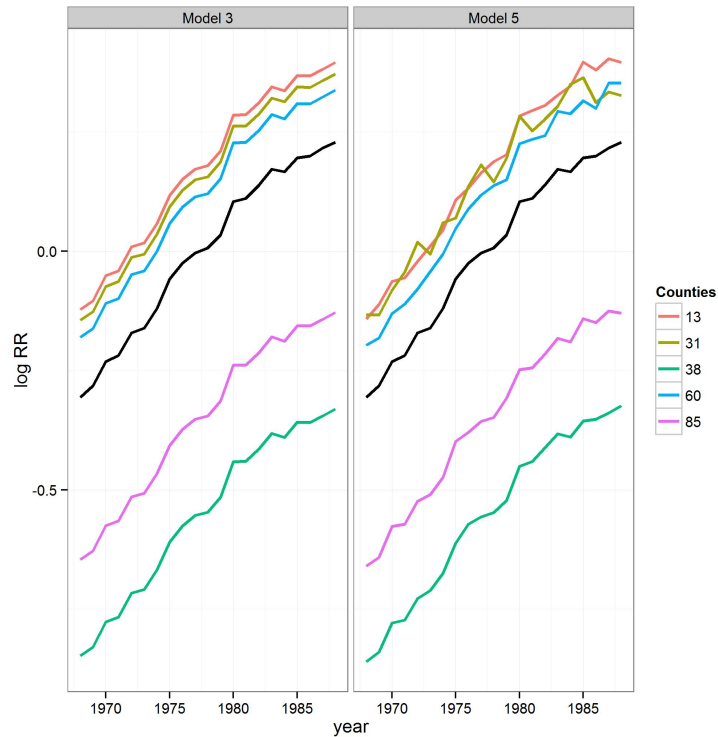
$$\begin{aligned}y_{it} &\sim \text{Poisson}(\mathbf{E}_{it}\rho_{it}) \\ \log \rho_{it} &= b_0 + b_i + \gamma_t + \psi_t + \delta_{it} \\ b_i &= v_i + u_i \\ \gamma_t &\sim \text{RW}(1) \\ \psi_t &\sim \text{N}(0, \sigma_\psi^2) \\ \delta_{it} &\sim \text{Normal}(0, \sigma_\delta^2)\end{aligned}$$

The simple additive model 4 can be extended by including **space-time interactions parameters**, δ_{it} , modelled as exchangeable random effects, that capture departure from the additive structure

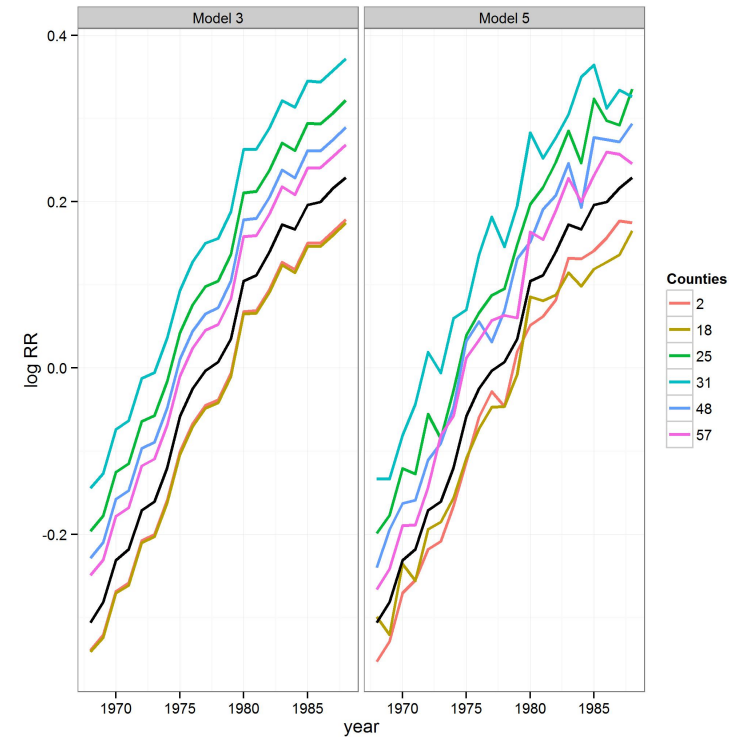
Interpretation of the interactions in model 5

- The interactions δ_{it} allow to highlight unusual temporal trends
- Rules based on the posterior probabilities $p(\delta_{it} > 0)$ for at least 1 time t
- Ohio lung cancer: 6 counties with unusual temporal trends

"usual" temporal trends



unusual temporal trends



INLA code for model with interaction (Model 5)

- In INLA is very easy to include the interaction in the formula environment
- We need to specify an index for the interaction, i.e. for each combination of area/time

$$\log \rho_{it} = b_0 + b_i + \gamma_t + \psi_t + \delta_{it}$$

$$b_i = v_i + u_i$$

$$\gamma_t \sim \text{RW}(1)$$

$$\psi_t \sim N(0, \sigma_\psi^2)$$

$$\delta_{it} \sim \text{Normal}(0, \sigma_\delta^2)$$

```
> formula.intI = y ~ + f(county,model="bym",  
+                       graph=Ohio.adj) +  
+                       f(year,model="rw1") +  
+                       f(year2,model="iid") +  
+                       f(area.year,model="iid")
```


Types of interactions

Another example: Birth weight in Georgia

- Count of babies weighing less than 2500g in 159 counties in Georgia (US)
- Period 2000 – 2010



Different types of interactions

$$\begin{aligned}y_{it} &\sim \text{Poisson}(E_{it}\rho_{it}) \\ \log \rho_{it} &= b_0 + b_i + \gamma_t + \psi_t + \delta_{it} \\ b_i &= v_i + u_i \\ \gamma_t &\sim \text{RW}(1) \\ \psi_t &\sim N(0, \sigma_\psi^2)\end{aligned}$$

Characteristics of ST interaction

Interaction	Parameters	Rank
I	v and ψ	nT
II	v and γ	$n(T-1)$ for RW1, $n(T-2)$ for RW2
III	u and ψ	$(n-1)T$
IV	u and γ	$(n-1)(T-1)$ for RW1, $(n-1)(T-2)$ for RW2

- We will learn how to develop Type I interaction in INLA

How to model interactions

- Data in R format

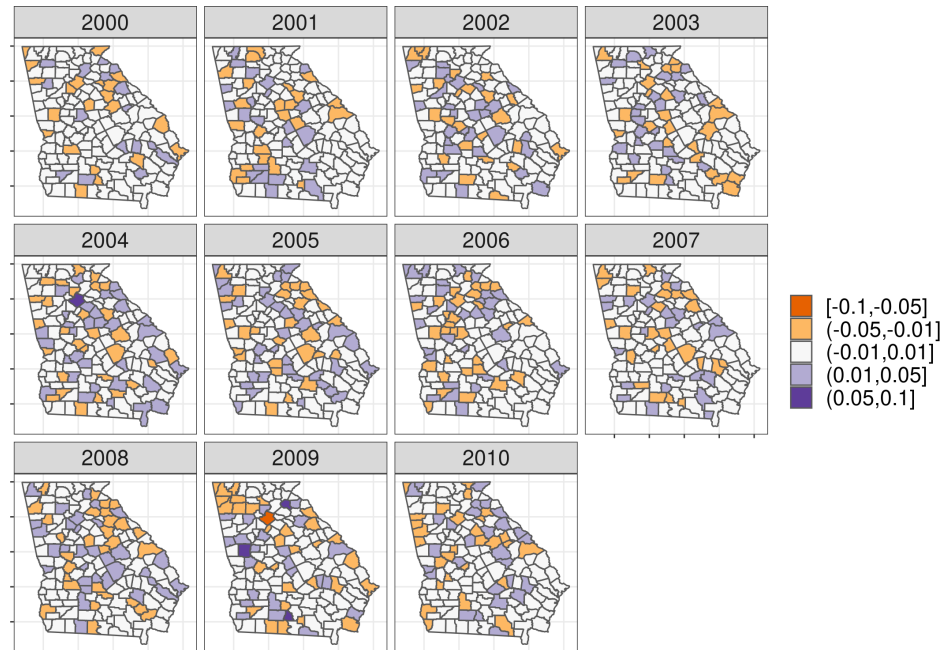
```
# A tibble: 1,749 × 8
  ID.area  Obs ID.year  Exp    y    E NAME  ID.area.year
  <int> <int>   <dbl> <dbl> <int> <dbl> <chr>      <int>
1      1    20      1  25.8   20  25.8 Appling      1
2      1    24      2  25.1   24  25.1 Appling      2
3      1    25      3  22.8   25  22.8 Appling      3
4      1    31      4  25.2   31  25.2 Appling      4
5      1    24      5  24.9   24  24.9 Appling      5
6      1    40      6  26.7   40  26.7 Appling      6
7      1    29      7  26.0   29  26.0 Appling      7
8      1    35      8  27.2   35  27.2 Appling      8
9      1    26      9  25.1   26  25.1 Appling      9
10     1    25     10  24.5   25  24.5 Appling     10
# ... with 1,739 more rows
```

- We need to make sure to have an index for area (ID.area), one for time (ID.year) and one for the interaction (ID.area.year)

Type I interaction in INLA

Let's adopt Type I interaction, which assumes that the two unstructured effects v_i and ψ_t interact

```
> Georgia.adj = "Georgia.graph"  
> ID.year2 = data_INLA$ID.year  
> formula.intI = y ~ + f(ID.area,model="bym2",  
+                       graph=Georgia.adj) +  
+                       f(ID.year,model="rw1") +  
+                       f(ID.year2,model="iid") +  
+                       f(ID.area.year,model="iid")
```



Kronecker product

- For the interactions of type II-IV we will need to use the **Kronecker product** to specify the dependencies. It is a matrix multiplication which returns a block matrix.

For instance

$$\begin{bmatrix} 1 & 2 \\ 3 & 4 \end{bmatrix} \otimes \begin{bmatrix} 0 & 5 \\ 6 & 7 \end{bmatrix} = \begin{bmatrix} 1 \begin{bmatrix} 0 & 5 \\ 6 & 7 \end{bmatrix} & 2 \begin{bmatrix} 0 & 5 \\ 6 & 7 \end{bmatrix} \\ 3 \begin{bmatrix} 0 & 5 \\ 6 & 7 \end{bmatrix} & 4 \begin{bmatrix} 0 & 5 \\ 6 & 7 \end{bmatrix} \end{bmatrix}$$

- There is a function in R which does exactly that: `kron(a,b)`
- We will not go through type II-IV of interactions, as they are pretty complex and also they can take a **very long** time to run
- If you are interested in knowing more how to set interactions using the Kronecker product in INLA see Goicoa, Adin, Ugarte, and Hodges (2018)

Summary

- Increase quality of datasets that are both spatially and temporally indexed
- Advanced methods to deal with this type of data
- Allow to interpret the stability (or not) of the spatial patterns

References

Goicoa, T., A. Adin, M. Ugarte, et al. (2018). "In spatio-temporal disease mapping models, identifiability constraints affect PQL and INLA results". In: *Stochastic Environmental Research and Risk Assessment* 32.3, pp. 749-770.