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Introduction

Corona Virus

Introduction to Bayesian Inference

3

Bayesian inference is a branch of statistics that uses the Bayesian concept of probability and Bayes' theorem to investigate questions of stochastics.

Characteristic for Bayesian statistics is the consistent use of probability distributions or marginal distributions, whose form conveys the accuracy of the procedures or reliability of the data and the procedure.

The Bayesian concept of probability does not presuppose infinitely repeatable random experiments, so that Bayesian methods can be used even with small data sets. A small amount of data leads to a broad probability distribution, which is not strongly localised.

In the Bayesian approach, the parameters of interest are treated as random variables that are governed by their parameters, for instance the mean and standard deviation, and distributions.

Bayesian inference is an essential technique in mathematical statistics and the polar opposite of the frequentist approach, in which a hypothesis is tested without being assigned a probability.

In the Bayesian approach a *prior* distribution $\pi\left(\boldsymbol{\theta},\sigma^{2}\right)$ is introduced as part of the model. This distribution is intended to express a state of knowledge or ignorance about $\boldsymbol{\theta}$ and σ^{2} prior to obtaining the data. Using the prior distribution, the likelihood function $\pi\left(\boldsymbol{x}|\boldsymbol{\theta},\sigma^{2}\right)$, and the observed data \boldsymbol{x} , it is possible to calculate the probability distribution $\pi\left(\boldsymbol{\theta},\sigma^{2}|\boldsymbol{x}\right)$ of $\boldsymbol{\theta}$ and σ^{2} given the data \boldsymbol{x} . This distribution is called the *posterior* distribution of $\boldsymbol{\theta}$ and σ^{2} and is used to make inferences about the parameters [BT11, p. 6].

3.1 Preliminaries

This work follows strict notation rules to easily represent different elements such as matrices or graphs and contains frequently used abbreviations. These and some other basic concepts used in this work are introduced below. The notation follows the one used by Rue and Held [RH05, pp. 14–19].

3.1.1 Matrices and Vectors

Vectors and matrices are indicated by bold notation, such as \boldsymbol{x} and \boldsymbol{A} . The transpose of \boldsymbol{A} is denoted by \boldsymbol{A}^T . The element in the ith row and jth column of \boldsymbol{A} is referenced by A_{ij} . This notation is also used for vectors and x_i denotes the ith element of a vector. The vector $(x_i, x_{i+1}, ..., x_j)^T$ is abbreviated to $\boldsymbol{x}_{i:j}$. If the columns $\boldsymbol{A}_1, \boldsymbol{A}_2, ..., \boldsymbol{A}_m$ of a $n \times m$ matrix \boldsymbol{A} are stacked on top of each other, this is denoted by vec $(\boldsymbol{A}) = \left(\boldsymbol{A}_1^T, \boldsymbol{A}_2^T, ..., \boldsymbol{A}_m^T\right)$. Deleting rows and/or columns from \boldsymbol{A} creates a submatrix. If a submatrix of a $n \times n$ matrix \boldsymbol{A} can be obtained by removing rows and columns of the same index, it is called a principal submatrix. If this matrix can be obtained by deleting the last n-r rows and columns, it is called a leading principal submatrix of \boldsymbol{A} .

A diagonal $n \times n$ matrix **A** is denoted by diag (**A**) and has the following structure:

$$\operatorname{diag}\left(oldsymbol{A}
ight) = egin{pmatrix} A_{11} & & & & \ & \ddots & & \ & & A_{nn} \end{pmatrix}.$$

The identity matrix is denoted by I.

If $A_{ij} = 0$ for i < j or $A_{ij} = 0$ where i > j, then \boldsymbol{A} is called *upper triangular* and *lower triangular* respectively. The *bandwidth* of a matrix \boldsymbol{A} is defined as $\max\{|i-j|:A_{ij}\neq 0\}$. The *lower bandwidth* is given by $\max\{|i-j|:A_{ij}\neq 0\}$ and $i > j\}$. $|\boldsymbol{A}|$ denotes the *determinant* of a $n \times n$ matrix \boldsymbol{A} and is equal to the product of the eigenvalues of \boldsymbol{A} . The *rank* of \boldsymbol{A} , referenced by rank (\boldsymbol{A}) , is the number of linearly independent rows or columns of \boldsymbol{A} . The sum of the diagonal elements is called *trace* of \boldsymbol{A} , trace $(\boldsymbol{A}) = \sum_i A_{ii}$.

Finally, ' \odot ' denotes the element-wise multiplication of two matrices of size $n \times m$, ' \odot ' denotes the element-wise division and raising each element of a matrix \boldsymbol{A} to a scalar power uses the symbol ' \odot ' [RH05, pp. 14–15].

3.1.2 Lattice and Torus

 \mathcal{I}_n denotes a (regular) lattice (or grid) of size $n=(n_1,n_2)$ (in the two-dimensional case). \boldsymbol{x} can take values on \mathcal{I}_n and $x_{i,j}$ denotes the value of \boldsymbol{x} at location ij, for $i=1,...,n_1$ and $j=1,...,n_2$. For easier reading this will be shortened to x_{ij} . On an infinite lattice \mathcal{I}_{∞} , ij are numbered as $i=0,\pm 1,\pm 2,...$, and $j=0,\pm 1,\pm 2,...$

A lattice with cyclic or toroidal boundary conditions is referred to as *torus* and is denoted by \mathcal{I}_{∞} . The dimension is $\mathbf{n} = (n_1, n_2)$ (in the two-dimensional case) and all indices are modulus \mathbf{n} and run from 0 to $n_1 - 1$ or $n_2 - 1$. If a GMRF \mathbf{x} is defined on $\mathcal{I}_{\mathbf{n}}$, the toroidal boundary conditions imply that x_{-2,n_2} is equal to $x_{n_1-2,0}$ since $-2 \mod n_1$ is equal to $n_1 - 2$ and $n_2 \mod n_2$ is equal to 0.

An *irregular lattice* refers to a spatial configuration of regions i = 1, ..., n where the regions (mostly) have common boundaries, for instance the states of a nation [RH05, pp. 15–16]. An example of an irregular lattice is shown in Figure 3.1.

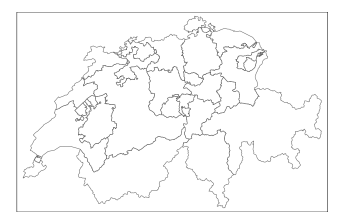


Fig. 3.1: The cantons of Switzerland, an example of an irregular lattice.

3.1.3 General Notation and Abbreviations

For $C \in \mathcal{I} = \{1, ..., n\}$ let $\boldsymbol{x}_C = \{x_i : i \in C\}$. -C denotes the set $\mathcal{I} - \mathcal{C}$ such that $\boldsymbol{x}_{-C} = \{x_i : i \notin C\}$. For two sets A and B, $A \setminus B = \{i : i \in A \text{ and } i \notin B\}$.

 $\pi\left(\cdot\right)$ denotes the density of its arguments, for example $\pi\left(\boldsymbol{x}\right)$ for the density of \boldsymbol{x} and $\pi\left(\boldsymbol{x}_{A}|\boldsymbol{x}_{-A}\right)$ for the conditional density of \boldsymbol{x}_{A} , given \boldsymbol{x}_{-A} . ' \sim ' is used when a variable is 'distributed' according to the law l.

The expected value is denoted by $\mathbb{E}[\cdot]$, the variance by $\text{Var}(\cdot)$, the covariance by

 $Cov(\cdot)$, the precision by $Prec(\cdot) = Cov(\cdot)^{-1}$, the correlation by $Corr(\cdot, \cdot)$ and a probability by $\mathbb{P}(\cdot)$ [RH05, p. 16].

3.1.4 Symmetric Positive Definite Matrices

An $n \times n$ matrix **A** is positive definite exactly if

$$\mathbf{x}^T \mathbf{A} \mathbf{x} > 0, \quad \forall \mathbf{x} \neq \mathbf{0}.$$

If A is also symmetric, it is called a symmetric positive definite (SPD) matrix. Only SPD matrices are considered and sometimes the notation A > 0 is used for an SPD matrix A.

An SPD matrix A has some of the following properties.

- 1. rank(A) = n.
- 2. |A| > 0.
- 3. $A_{ii} > 0$.
- 4. $A_{ii}A_{jj} A_{ij}^2 > 0$, for $i \neq j$.
- 5. $A_{ii} + A_{jj} 2|A_{ij}| > 0$ for $i \neq j$.
- 6. $\max A_{ii} > \max_{i \neq j} |A_{ij}|$.
- 7. A^{-1} is SPD.
- 8. All principal submatrices of A are SPD.

If A and B are SPD, A + B is also SPD, but the reverse is generally not true. Additionally, if AB = BA, AB is SPD.

The following conditions are all sufficient and necessary for a symmetric matrix A to be SPD:

- 1. All eigenvalues $\lambda_1, ..., \lambda_n$ of \boldsymbol{A} are strictly positive.
- 2. There exists such a matrix C that $A = CC^T$. If C is lower triangular, it is called the *Cholesky triangle* of A.
- 3. All leading principal submatrices have strictly positive determinants.

A sufficient, but not necessary condition for a (symmetrical) matrix to be SPD is the criterion of *diagonal dominance*:

$$A_{ii} - \sum_{j:j \neq i} |A_{ij}| > 0, \quad \forall i.$$

A $n \times n$ matrix ${\bf A}$ is called a *symmetric positive semidefinite* (SPSD) matrix. An SPSD matrix ${\bf A}$ is sometimes denoted ' ${\bf A} \ge 0$ ' [RH05, pp. 18–19].

3.2 Basic Concepts of Bayesian Theory

3.2.1 Bayes' Theorem

At the heart of Bayesian inference is *Bayes' theorem*, which describes the probability of an event given prior knowledge of factors that might influence the event. Let $\mathbf{x}^T = (x_1, ..., x_n)$ be a vector of n observations whose probability distribution $\pi(\mathbf{x}|\mathbf{\theta})$ depends on the values of k parameters $\mathbf{\theta}^T = (\theta_1, ..., \theta_k)$. Let $\pi(\mathbf{\theta})$ be the probability distribution of $\mathbf{\theta}$. Then

$$\pi\left(\boldsymbol{x}|\boldsymbol{\theta}\right)\pi\left(\boldsymbol{\theta}\right) = \pi\left(\boldsymbol{x},\boldsymbol{\theta}\right) = \pi\left(\boldsymbol{\theta}|\boldsymbol{x}\right)\pi\left(\boldsymbol{x}\right). \tag{3.1}$$

Given the observed data x, the conditional distribution of θ is

$$\pi\left(\boldsymbol{\theta}|\boldsymbol{x}\right) = \frac{p\left(\boldsymbol{x}|\boldsymbol{\theta}\right)\pi\left(\boldsymbol{\theta}\right)}{\pi\left(\boldsymbol{x}\right)}.$$
(3.2)

This last statement is known as Bayes' theorem [Bay63]. The *prior* distribution $\pi(\theta)$ contains knowledge about θ without knowledge of the data. $\pi(\theta|x)$ contains what is known about θ given knowledge of the data and is the *posterior* distribution of θ given x.

If $\pi(x|\theta)$ is considered as a function of θ instead of x, it is called the *likelihood* function of θ given x and can be written as $l(\theta|x)$. Thus Bayes' theorem can be written as

$$\pi\left(\boldsymbol{\theta}|\boldsymbol{x}\right) \propto l\left(\boldsymbol{\theta}|\boldsymbol{x}\right)\pi\left(\boldsymbol{\theta}\right).$$
 (3.3)

It is evident that the posterior distribution of θ given the data x is proportional to the product of the distribution of θ prior to observing the data and the likelihood function of θ given x. Therefore,

posterior distribution \propto likelihood \times prior distribution.

The data x modifies the prior knowledge of θ through the likelihood function, and thus can be be regarded as a representation of the information about θ derived from the data [BT11].

3.2.2 Conditional Independence

In probability theory, two random variables x and y are *independent* given a third variable z if and only if the occurrence of x and y in their conditional probability

distribution given z are independent events. To calculate the conditional density of x_A , given x_{-A} , the following statement will repeatedly be used,

$$\pi\left(\boldsymbol{x}_{A}|\boldsymbol{x}_{-A}\right) = \frac{\pi\left(\boldsymbol{x}_{A}, \boldsymbol{x}_{-A}\right)}{\pi\left(\boldsymbol{x}_{-A}\right)} \propto \pi\left(\boldsymbol{x}\right). \tag{3.4}$$

It follows that x and y are independent precisely when $\pi\left(x,y\right)=\pi\left(x\right)\pi\left(y\right)$, which is expressed by $x\perp y$. x and y are conditionally independent for a given z if and only if $\pi\left(x,y|z\right)=\pi\left(x|z\right)\pi\left(y|z\right)$ [Daw79]. The conditional independence can be easily validated with the help of the following factorisation criterion,

$$x \perp y | z \Longleftrightarrow \pi (x, y, z) = f(x, z) g(y, z), \qquad (3.5)$$

for some functions f and g, and for all z with $\pi(z) > 0$ [RH05, pp. 16–17].

3.2.3 Undirected Graphs

Undirected graphs are used to represent the conditional independence structure in a Gaussian Markov random field. An *undirected graph* $\mathcal G$ is defined as a tuple $\mathcal G=(\mathcal V,\mathcal E)$, where $\mathcal V$ contains all nodes in the graph and $\mathcal E$ is the set of edges $\{i,j\}$, with $i,j\in\mathcal V$ and $i\neq j$. For $\{i,j\}\in\mathcal E$ there exists an undirected edge from node i to node j in the other case such an edge does not exist. If $\{i,j\}\in\mathcal E$ $\forall i,j\in\mathcal V$ with $i\neq j$ a graph is *fully connected*. Most often $\mathcal V=\{1,2,...,n\}$ will be assumed, which is referred to as *labelled*. A simple example of an undirected graph is shown in Figure 3.2.



Fig. 3.2: An undirected labelled graph with 3 nodes, $V = \{1, 2, 3\}$ and $\mathcal{E} = \{\{1, 2\}, \{2, 3\}\}$.

The *neighbours* of node i are defined as all nodes in \mathcal{G} with an edge to node i,

$$ne(i) = \{j \in \mathcal{V} : \{i, j\} \in \mathcal{E}\}.$$

This definition can be extended to a set $A \subset \mathcal{V}$, where the neighbours of A are defined as

$$\operatorname{ne}\left(A\right)=\bigcup_{i\in A}\operatorname{ne}\left(i\right)\backslash A.$$

A path from i_1 to i_m is defined as a sequence of certain nodes in $\mathcal{V}, i_1, i_2, ..., i_m$, for which $(i_j, i_{j+1}) \in \mathcal{E}$ for j = 1, ..., m-1. Two nodes $i \notin C$ and $j \notin C$ are separated by a subset $C \subset \mathcal{V}$, if every path from i to j contains at least one node from C. Two disjoint sets $A \subset \mathcal{V} \notin C$ and $B \subset \mathcal{V} \notin C$ are separated by C, if all $i \in A$ and $j \in B$ are separated by C, that is, it is not possible to "wander" on the graph from somewhere in A and end somewhere in B without crossing C.

If i and j are neighbours in \mathcal{G} , this can be expressed by $i \stackrel{\mathcal{G}}{\sim} j$ or $i \sim j$ for the case where the graph is implicit. It follows that $i \sim j \iff j \sim i$.

Let A be a subset of \mathcal{V} . A subgraph \mathcal{G}^A is a graph restricted to A, i.e., the graph obtained after removing all nodes that do not belong to A and all edges where at least one node does not belong to A. $\mathcal{G}^A = \left\{ \mathcal{V}^A, \mathcal{E}^A \right\}$, where $\mathcal{V}^A = A$ and

$$\mathcal{E}^A = \{\{i, j\} \in \mathcal{A} \text{ and } \{i, j\} \in A \times A\}.$$

Let $\mathcal G$ be the graph in Figure 3.2 and $\mathcal A=\{2,3\}$, then $\mathcal V^A=\{2,3\}$ and $\mathcal E^A=\{\{2,3\}\}$ [RH05, pp. 17–18].

3.2.4 The Exponential Family

In statistics and probability theory, the *exponential family* is a parametric set of probability distributions of a specific form. The distribution of a random variable \boldsymbol{y} belongs to the exponential family if the discrete or continuous density with respect to a σ -finite measure of \boldsymbol{y} has the form

$$f(\boldsymbol{y}|\boldsymbol{\theta},\lambda) = \exp\left(\frac{\boldsymbol{y}^T\boldsymbol{\theta} - b(\boldsymbol{\theta})}{\lambda} + c(\boldsymbol{y},\lambda)\right),$$
 (3.6)

with $c(\boldsymbol{y}, \lambda) \geq 0$. $\boldsymbol{\theta} \in \Theta \subset \mathbb{R}^q$ is the *natural* or *canonical* parameter of the exponential family, while $\lambda > 0$ is a *dispersion* or *nuisance* parameter [HL81]. The natural parameter space Θ is the set of all $\boldsymbol{\theta}$ satisfying

$$0 < \int \exp\left(\left(\boldsymbol{y}^{T}\boldsymbol{\theta} - b(\boldsymbol{\theta})\right)/\lambda + c(\boldsymbol{y}, \lambda)\right) d\boldsymbol{y} < \infty$$
(3.7)

Moreover, $b(\theta)$ is a twice differentiable function and all moments of y exist. Specifically,

$$\mathbb{E}_{\boldsymbol{\theta}}(\boldsymbol{y}) = \mu(\boldsymbol{\theta}) = \frac{\partial b(\boldsymbol{\theta})}{\partial \boldsymbol{\theta}}$$
 (3.8)

$$\mathbb{E}_{\boldsymbol{\theta}}(\boldsymbol{y}) = \mu(\boldsymbol{\theta}) = \frac{\partial b(\boldsymbol{\theta})}{\partial \boldsymbol{\theta}}$$

$$\operatorname{Cov}_{\boldsymbol{\theta}}(\boldsymbol{y}) = \boldsymbol{\Sigma}(\boldsymbol{\theta}) = \lambda \frac{\partial^2 b(\boldsymbol{\theta})}{\partial \boldsymbol{\theta} \partial \boldsymbol{\theta}^T}.$$
(3.8)

The covariance matrix $\Sigma(\boldsymbol{\theta})$ is positive definite in Θ^0 , therefore $\mu:\Theta^0\to M=\mu\left(\Theta^0\right)$ is injective. By substituting the inverse function $theta(\mu)$ into $\frac{\partial^2 b(\theta)}{\partial \theta \partial \theta^T}$, the variance function

$$v(\mu) = \frac{\partial^2 b(\boldsymbol{\theta}(\mu))}{\partial \boldsymbol{\theta} \partial \boldsymbol{\theta}^T}$$
 (3.10)

is given and the covariance can be written as

$$Cov_{\boldsymbol{\theta}}(\boldsymbol{y}) = \lambda v(\mu). \tag{3.11}$$

Important members of the exponential family are the normal, binomial, Poisson, gamma and multivariate normal distribution [FT13, p. 433].

The Multivariate Normal Distribution

The density of a normally distributed random variable $\mathbf{x} = (x_1, ..., x_n)^T$, $n < \infty$ with mean vector $\boldsymbol{\mu}$ $(n \times 1)$ and SPD covariance matrix $\boldsymbol{\Sigma}$ $(n \times n)$ is

$$\pi\left(\boldsymbol{x}\right) = \left(2\pi\right)^{-n/2} |\mathbf{\Sigma}|^{-1/2} \exp\left(-\frac{1}{2}\left(\boldsymbol{x}-\boldsymbol{\mu}\right)^T \mathbf{\Sigma}^{-1}\left(\boldsymbol{x}-\boldsymbol{\mu}\right)\right), \ \boldsymbol{x} \in \mathbb{R}^n$$
 (3.12)

Here, $\mu_i = \mathbb{E}[x_i]$, $\Sigma_{ij} = \text{Cov}(x_i, x_j)$, $\Sigma_{ii} = \text{Var}(x_i) > 0$ and $\text{Corr}(x_i, x_j) = 0$ $\Sigma_{ij}/(\Sigma_{ii}\Sigma_{jj})^{1/2}$. This is written as $\boldsymbol{x} \sim \mathcal{N}(\boldsymbol{\mu}, \boldsymbol{\Sigma})$. For $n=1, \mu=0$ and $\Sigma_{11}=1$ the standard normal distribution is obtained.

 $m{x}$ is now split up into $m{x} = \left(m{x}_{m{A}}^T, m{x}_{m{B}}^T\right)$ and $m{\mu}$ and $m{\Sigma}$ are divided accordingly:

$$m{\mu} = egin{pmatrix} m{\mu}_{m{A}} \ m{\mu}_{m{B}} \end{pmatrix} \quad ext{ and } \quad m{\Sigma} = egin{pmatrix} m{\Sigma}_{m{A}m{A}} & m{\Sigma}_{m{A}m{B}} \ m{\Sigma}_{m{B}m{A}} & m{\Sigma}_{m{B}m{B}} \end{pmatrix}.$$

Some basic properties of the multivariate normal distribution are the following.

- 1. $x_A \sim \mathcal{N}(\mu_A, \Sigma_{AA})$.
- 2. $\Sigma_{AB} = 0$ precisely when x_A and x_B are independent.

3. The conditional distribution $\pi\left(\pmb{x_A}|\pmb{x_B}\right)$ is $\mathcal{N}\left(\pmb{\mu_{A|B}}, \pmb{\Sigma_{A|B}}\right)$, where

$$egin{align} m{\mu_{A|B}} &= m{\mu_A} + m{\Sigma_{AB}} m{\Sigma_{BB}}^{-1} \left(m{x_B} - m{\mu_B}
ight) \ ext{and} \ m{\Sigma_{A|B}} &= m{\Sigma_{AA}} - m{\Sigma_{AB}} m{\Sigma_{BB}}^{-1} m{\Sigma_{BA}}. \end{split}$$

4. If
$$x \sim \mathcal{N}(\mu, \Sigma)$$
 and $x' \sim \mathcal{N}(\mu', \Sigma')$ are independent,
then $x + x' \sim \mathcal{N}(\mu + \mu', \Sigma + \Sigma')$ [RH05, pp. 19–20].

3.2.6 Skewness

Skewness is a statistical indicator that describes the type and strength of the asymmetry of a probability distribution. It shows whether and how strongly the distribution is skewed to the right (right-skewed, left-skewed, negative skewness) or to the left (left-skewed, right-skewed, positive skewness). Any non-symmetrical distribution is called skewed.

The skewness of a random variable X is the third standardized moment, defined as

$$\operatorname{Skew}[X] = \mathbb{E}\left[\left(\frac{X-\mu}{\sigma}\right)^{3}\right] = \frac{\mathbb{E}\left[\left(X-\mu\right)^{3}\right]}{\left(\mathbb{E}\left[\left(X-\mu\right)^{3/2}\right]\right)^{2}} = \frac{\mu_{3}}{\sigma^{3}},\tag{3.13}$$

with μ_3 the third central moment and σ the standard deviation [DS11; Wil44].

3.2.7 Kurtosis

Kurtosis is a measure of the slope of a probability distribution of a random variable. Distributions with low kurtosis scatter relatively evenly; for distributions with high kurtosis, the scatter results more from extreme but rare events.

The kurtosis of a random variable X is the fourth standardized moment, defined as

$$\operatorname{Kurt}[X] = \mathbb{E}\left[\left(\frac{X-\mu}{\sigma}\right)^4\right] = \frac{\mathbb{E}\left[\left(X-\mu\right)^4\right]}{\left(\mathbb{E}\left[\left(X-\mu\right)^2\right]\right)^2} = \frac{\mu_4}{\sigma^4},\tag{3.14}$$

with μ_4 the fourth central moment and σ the standard deviation [DeC97; Wil44].

3.3 Prior Selection

3.3.1 Conjugate Priors

One property of exponential families is that they have conjugate priors [DY79], which is an important property in Bayesian statistics. If the posterior distribution $\pi(\theta|\mathbf{x})$ and the prior distribution $\pi(\theta)$ belong to the same probability distribution family, the prior and posterior distributions are called *conjugate* distributions. Furthermore, the prior for the likelihood function $\pi(\mathbf{x}|\theta)$ is called the *conjugate prior*. These priors were first discussed and formalised by Raifa and Schlaifer in 1961 [RS61].

The construction of a conjugate prior is done by factorising the likelihood function into two parts. One part must be independent of the parameter(s) of interest but can be dependent on the data, while the other factor is a function that depends on the parameter(s) of interest and is dependent on the data only through the sufficient statistics. The family of conjugate priors is by definition proportional to the second factor. The posterior distribution resulting from the conjugate prior is itself a member of the same family as the conjugate prior [RS61]. In cases where the prior and posterior distributions are part of the same family, the prior is said to be closed under sampling. Furthermore, since the data are only incorporated into the posterior distribution through the sufficient statistics, there exist relatively simple formulas for updating the prior into the posterior [Fin97]. An example of the construction of a conjugated prior is provided in the appendix.

A drawback of conjugated priors is that the a priori known information about μ may be insufficient for determining both parameters or may be inconsistent with the structure imposed by conjugacy [RMR10]. Moreover, these priorities can be too restrictive and not every belief about the priori can be described [Irw05].

Thus, although conjugate priors are easy to handle both mathematically and computationally [Irw05], they are not often used in practice because of these drawbacks.

3.3.2 Jeffreys' Prior

As conjugate priors depend on information from the data, they are so-called *informative priors*. *Jeffreys' prior* [Jef46] is characterised in comparison to other a priori distributions by the fact that it is invariant to a reparametrisation of the model parameters. Due to this invariance, the Jeffreys' a priori distribution is also called *non-informative*.

Let $\phi = \phi\left(\theta\right)$ be a one-to-one transformation of θ and $I\left(\cdot\right)$ be the Fisher information, then

$$I_{\phi}\left(\phi\right) = I_{\theta}\left(\theta\right) \left(\frac{\partial \theta}{\partial \phi}\right)^{2}.$$
(3.15)

If some principle of choice led to $\pi\left(\theta\right)$ as a non-informative prior for θ , the same principle should lead to

$$\pi_{\phi}\left(\phi\right) = \pi_{\theta}\left(\theta\right) \left| \frac{\partial \theta}{\partial \phi} \right| \tag{3.16}$$

as a non-informative prior for ϕ . The desired "invariance" is then given by

$$\pi_{\phi}\left(\phi\right) \propto \sqrt{I_{\phi}\left(\phi\right)} = \sqrt{I_{\theta}\left(\theta\right)} \left| \frac{\partial \theta}{\partial \phi} \right| \propto \pi_{\theta}\left(\theta\right) \left| \frac{\partial \theta}{\partial \phi} \right|$$
 (3.17)

[BT11, pp. 42–43]. While Jeffreys' priors work well for single parameter models, they struggle when it comes to models with multidimensional parameters as these priors can have poor convergence properties [Jor10].

3.3.3 Penalised Complexity Priors

One issue when selecting the prior distribution of a particular parameter is that it is not always intuitive when it comes to understanding and interpreting this distribution, something that is essential to ensure that it behaves as intended by the user. This problem can be addressed by using *penalised complexity priors*, which is a methodology that penalises the complexity of model components in relation to deviation from simple base model formulations.

PC priors provide a systematic and unified approach to calculating priority distributions for parameters of model components by using an inherited nested structure. This structure contains two models, the base model and a flexible version of the model. The first of the two is generally characterised by a fixed value of the relevant parameter, while the second version is considered a function of the random parameter. By penalising the deviation from the flexible model to the fixed base model, the PC prior is calculated [Sim+17].

3.3.3.1 The Principles Behind PC Priors

Four main principles should be followed to calculate priors in a consistent way and to understand their properties.

Support to Occam's Razor

Let $\pi(x|\xi)$ denote the density of a model component x and ξ the parameter to which a prior distribution is to be assigned. The base model is characterised by a density $\pi(x|\xi=\xi_0)$, where ξ_0 is a fixed value. The prior for ξ should be such that proper shrinkage is given to ξ_0 . The simplicity of the model is therefore prioritised over the complexity of the model, preventing overfitting [Sim+17].

Penalisation of Model Complexity

Let $f_1 = \pi\left(x|\xi\right)$ and $f_0\left(x|\xi=\xi_0\right)$ denote the flexible model and the base model respectively. The complexity of f_1 compared to f_0 is characterised using the Kullback-Leibler divergence [KL51] to calculate a measure of complexity between the two models,

$$KLD\left(f_{1}||f_{2}\right) = \int f_{1}\left(x\right)\log\left(\frac{f_{1}\left(x\right)}{f_{0}\left(x\right)}\right)dx. \tag{3.18}$$

This can be used to measure the information that is lost when f_1 is approximated by the simpler model f_0 . For multinormal densities with zero mean, the calculation simplifies to

$$KLD(f_1||f_0) = \frac{1}{2} \left(trace \left(\Sigma_0^{-1} \Sigma_1 \right) - n - \ln \left(\frac{|\Sigma_1|}{|\Sigma_0|} \right) \right), \tag{3.19}$$

where $f_i \sim \mathcal{N}\left(0, \mathbf{\Sigma}_i\right)$, i=0,1, while n represents the dimension. For easier interpretation, the Kullback-Leibler divergence is transformed into a unidirectional distance measure

$$d(\xi) = d(f_1||f_0) = \sqrt{2\text{KLD}(f_1||f_0)}$$
(3.20)

which can be interpreted as a measure of distance from f_1 to f_0 [Sim+17].

Constant Rate Penalisation

The derivation of the PC prior is based on a system of constant rate penalisation, given by

$$\frac{\pi_{d}\left(d\left(\xi\right)+\delta\right)}{\pi_{d}\left(d\left(\xi\right)\right)}=r^{\delta},\qquad d\left(\xi\right),\delta\geq0. \tag{3.21}$$

 $r \in (0,1)$ represents the constant decay rate and thus implies that the relative change in the priority distribution for $d(\xi)$ is independent of the actual distance. Therefore, $d(\xi)$ is exponentially distributed with density $\pi(d(\xi)) = \lambda \exp(-\lambda d(\xi))$ and rate

 $\lambda = -\ln{(r)}$. By a standard variable change transformation, the corresponding PC prior for ξ is given [Sim+17].

User-Defined Scaling

Since λ characterises the shrinkage properties of the prior, it is important that the rate can be chosen in an intuitive and interpretable way. One possibility is to determine λ by including a probability statement of tail events, for example

$$\mathbb{P}\left(Q\left(\xi\right) > U\right) = \alpha,\tag{3.22}$$

where U represents an assumed upper bound for an interpretable transformation $Q(\xi)$ and α denotes a small probability [Sim+17].

3.4 Markov-Chain-Monte-Carlo-Methods

Markov chain Monte Carlo methods, also referred to as MCMC methods, are a set of algorithms that enable sampling from probability distributions based on the construction of Markov chains. After a sufficient number of iterations, the stationary distribution of a Markov chain can be taken as the desired distribution, with the quality of this distribution improving as the number of iterations increases. Most of the time, the construction of such a chain is relatively simple; the real challenge is to determine how many steps are needed before convergence towards the stationary distribution is achieved. MCMC methods are mostly used to compute numerical approximations of multidimensional integrals, for instance in Bayesian statistics or computational biology. The two main concepts used in MCMC methods are Monte Carlo integration and the aforementioned Markov chains, hence the name Markov Chain Monte Carlo.

3.4.1 Monte Carlo Integration

Monte Carlo integration is a technique that uses the generation of random numbers for numerical computation of definite integrals and is especially useful for higher-dimensional integrals. The problem the method addresses is the computation of the integral

$$\mathbb{E}_f[h(X)] = \int_X h(x)f(x)dx. \tag{3.23}$$

The integral can be approximated by using a sample $(X_1, ..., X_m)$ generated from f and calculating the arithmetic mean

$$\overline{h}_m = \frac{1}{m} \sum_{j=1}^m h(x_j).$$
(3.24)

According to the Strong Law of Large Numbers, \overline{h}_m is likely to converge to $\mathbb{E}_f[h(X)]$. When the expectation of h^2 under f is finite, the convergence speed of \overline{h}_m can be assessed. The variance too can be estimated from the sample $(X_1,...,X_n)$ through

$$v_m = \frac{1}{m^2} \sum_{j=1}^{m} \left[h(x_j) - \overline{h}_m \right]^2.$$
 (3.25)

For m large,

$$\frac{\overline{h}_{m} - \mathbb{E}_{f} \left[h \left(X \right) \right]}{\sqrt{v_{m}}} \tag{3.26}$$

is approximately distributed as a $\mathcal{N}(0,1)$ variable. This can be used for constructing a convergence test and to calculate confidence bounds for the approximation of $\mathbb{E}_f\left[h\left(X\right)\right]$ [RC13, pp. 83–84]. The term Monte Carlo was first used in 1949 by Metropolis and Ulam to describe a method dealing with problems related to "integro-differential equations that occur in various branches of the natural sciences" [MU49].

3.4.2 Markov Chains

Markov chains are stochastic processes that aim to provide the probability of the occurrence of future events. A Markov chain is defined by the fact that even if only a limited history is known, predictions about future developments can be made just as reliably as if the entire history of a process were known. Thus, the probability of moving from the current state to any state depends only on the current state of the chain. These probabilities are defined by a *transition kernel*, which is a function K on $\mathcal{X} \times \mathcal{B}(\mathcal{X})$, such that

- i. $\forall x \in \mathcal{X}, K(x, \cdot)$ is a probability measure;
- ii. $\forall A \in \mathcal{B}(\mathcal{X}), K(\cdot, A)$ is measureable.

In the discrete case, the transition kernel is a matrix K with elements

$$\mathbb{P}_{xy} = \mathbb{P}\left(X_n = y | X_{n-1} = x\right), \quad x, y \in \mathcal{X}.$$

If \mathcal{X} is continuous, the kernel denotes the conditional density $K\left(x,x^{T}\right)$ of the transition $K\left(x,\cdot\right)$,

$$\mathbb{P}\left(X \in A|x\right) = \int_{A} K\left(x, x^{T}\right) dx^{T}.$$

Given a transition kernel K, a sequence $X_0, X_1, ..., X_t$ of random variables is a *Markov chain* (X_n) , if, for any t, the conditional distribution of X_t given the previous states is the same as the distribution of X_t given the last state, x_{t-1} ,

$$\mathbb{P}(X_{t+1} \in A | x_0, x_1, x_2, ..., x_t) = \mathbb{P}(X_{t+1} \in A | x_t)$$

$$= \int_{A} K(x_t, dx). \tag{3.27}$$

These chains were first introduced by Markov in 1906 [Mar06]. Markov chains can have certain properties that affect their long-term behaviour and are of particular importance for MCMC algorithms. In sections 3.4.2.1– 3.4.2.5, some of them will be introduced.

3.4.2.1 Irreducibility

Irreducibility is critical to the construction of Markov chain Monte Carlo algorithms, as it ensures the convergence of such an algorithm. A Markov chain is *irreducible* if all states communicate, that is, for all states i and j the probability of getting from i to j in finite time is true positive.

Formally speaking, given a measure φ , a Markov chain (X_n) with transition kernel K(x,y) is φ -irreducible, if, for every $A \in B(\mathcal{X})$ with $\varphi(A) > 0$, there exists n such that $K^n(x,A) \forall x \in \mathcal{X}$. The chain is *strongly* φ -irreducible if $n=1 \forall$ measurable A [RC13, pp. 213–214].

3.4.2.2 Periodicity

The behaviour of a Markov chain can sometimes be limited by deterministic constraints on the transitions from X_n to X_{n+1} . For discrete chains, the *period* of a state $w \in \mathcal{X}$ is defined as.

$$d(w) = \text{g.c.d. } \{m \ge 1; K^m(w, w) > 0\},\$$

with g.c.d the greatest common denominator. If a Markov chain is irreducible, the transition matrix can be written as a block matrix

$$\mathbf{P} = \begin{pmatrix} 0 & \mathbf{D}_1 & 0 & \dots & 0 \\ 0 & 0 & \mathbf{D}_2 & \dots & 0 \\ \vdots & \vdots & \ddots & & \\ \mathbf{D}_d & 0 & 0 & & 0 \end{pmatrix}, \tag{3.28}$$

It is evident that at every d-th step there is a return to the initial group. There exists only one value for the period when a chain is irreducible. If this value is 1, the irreducible chain is aperiodic [RC13, pp. 217–218].

3.4.2.3 Transience and Recurrence

To guarantee an acceptable approximation of a simulated model, a Markov chain needs to have good stability properties. Irreducibility is not strong enough to ensure that the trajectory of (X_n) enters A often enough. This leads to the formalisation of recurrence and transience.

In a finite space \mathcal{X} , a state $w \in \mathcal{X}$ is transient if it is finitely often visited and recurrent if it is almost certainly infinitely often visited.

For irreducible chains, these two properties are properties of the chain, not of a particular state [RC13, pp. 218–219].

3.4.2.4 Ergodicity

When looking at a Markov chain (X_n) from a temporal point of view, it is essential to establish to what the chain is converging. A natural candidate for the limiting distribution is the stationary distribution π which leads to the need to define sufficient conditions on (X_n) for X_n to be asymptotically distributed according to π . There are several conditions that can be imposed on the convergence of P^n , the distribution of X_n to π . The most fundamental and important is that of *ergodicity*, that is, independence of initial conditions.

If a Markov chain (X_n) is both aperiodic and positive recurrent, it is called an *ergodic* Markov chain [RC13, pp. 231–234].

3.4.2.5 Stationary distribution

A chain (X_n) is more stable if the marginal distribution of X_n is independent of n. This is a requirement for the existence of a probability distribution π such that $X_{n+1} \sim \pi$ if $X_n \sim \pi$. Markov chain Monte Carlo methods rely on the fact that this condition can be satisfied.

A σ -finite measure π is *invariant* for the transition kernel $K(\cdot,\cdot)$ if

$$\pi(B) = \int_{\mathcal{X}} K(x, B) \pi(dx), \quad \forall B \in \mathcal{B}(\mathcal{X}).$$

This distribution is referred to as *stationary* if π is a probability measure, as $X_0 \sim \pi$ implies that $X_n \sim \pi$ is $\forall n$. An irreducible Markov chain has a stationary distribution precisely if it is positively recurrent. The distribution is then given by

$$\pi_x = (\mathbb{E}_x \left[\tau_x \right])^{-1}, \quad x \in \mathcal{X},$$
(3.29)

where $\mathbb{E}_x [\tau_x]$ can be interpreted as the average number of transitions between two passages in x.

In practice, the stationary distributions are often of special interest. If these distributions are defined as the starting distribution of X_0 , then all following distributions of the states X_n for any n are equal to the starting distribution. The interesting question here is when such distributions exist and when any distribution converges against a stationary distribution of this kind [RC13, pp. 223–224].

3.4.3 The Metropolis-Hastings Algorithm

Having established the basics of MCMC methods, one of the best known MCMC algorithms, the Metropolis-Hastings algorithm, is introduced next. The algorithm is based on the Metropolis algorithm, which was developed to simulate the states of a system according to the Boltzmann distribution, with the newest state always depending on the previous state [Met+53].

The Metropolis-Hastings algorithm is a procedure for drawing random samples from a probability distribution from which direct sampling is difficult if a function proportional to the *target density* f is known. This function q(y|x) is called the *proposal density* and must be easy to simulate in order for the Metropolis-Hastings algorithm to be implementable. Moreover, it must be either explicitly present or *symmetric*, meaning q(x|y) = q(y|x).

The Metropolis-Hastings algorithm of a target density f and proposal density q produces a Markov chain $\left(X^{(t)}\right)$ by the following transition.

Algorithm 1 The Metropolis-Hastings Algorithm

Given $f(\boldsymbol{x})$ and $q(\boldsymbol{y}|\boldsymbol{x})$

- 1: Initialisation: Choose arbitrary x_t as the first sample
- 2: **for** each iteration t **do**
- 3: Generate $Y_t \sim q\left(\boldsymbol{y}|x^{(t)}\right)$
- 4: Take

$$X^{(t+1)} = \begin{cases} Y_t & \text{with probability } \mathbb{P}\left(x^{(t)}, Y_t\right) \\ x^{(t)} & \text{with probability } 1 - \mathbb{P}\left(x^{(t)}, Y_t\right) \end{cases}$$

where

$$\mathbb{P}(x,y) = \min \left\{ \frac{f(\mathbf{y})}{f(\mathbf{x})} \frac{q(\mathbf{x}|\mathbf{y})}{q(\mathbf{y}|\mathbf{x})}, 1 \right\}.$$
(3.30)

 $\mathbb{P}(x,y)$ is the *Metropolis-Hastings* acceptance probability.

The algorithm always accepts values y_t that lead to an increase in the ratio $f\left(y_t\right)/q\left(y_t|x^{(t)}\right)$ compared to the previous value $f\left(x^{(t)}\right)/q\left(x^{(t)}|y_t\right)$. In the symmetric case, the acceptance probability simplifies to

$$\mathbb{P}\left(x,y\right) = \min\left\{\frac{f\left(\boldsymbol{y}\right)}{f\left(\boldsymbol{x}\right)}, 1\right\}$$

[Has70].

If the Markov chain starts with a value $x^{(0)} > 0$, then $f\left(x^{(t)}\right) > 0 \ \forall t \in \mathbb{N}$ since the

values of y such that $f(y_t)=0$ will all be rejected by the algorithm. As the number of iterations t increases, the distribution of saved states $x_0,...,x_t$ will converge towards the target density $f(\boldsymbol{x})$ [RC13, pp. 270–275].

3.5 Latent Gaussian Models and INLA

In recent years, a growing amount of georeferenced data has become available, leading to an increased need for appropriate statistical modeling to handle large and complex datasets. Bayesian hierarchical models have proven to be effective in capturing complex stochastic structures in spatial processes. A large proportion of these models are based on latent Gaussian models, a subclass of structured additive regression models. These models include Integrated Nested Laplace Approximations (INLA), which are a class of models used to approximate the posterior marginals of a latent Gaussian field. These approximations work by reformulating the regression model as a three-part hierarchical model. The parts are as follows:

- 1. Approximation of the posterior marginal of θ by using the Laplace approximation.
- 2. Computation of the (simplified) Laplace approximation for selected values of θ to improve the Gaussian approximation.
- 3. Combination of the first two parts by numerical integration.

 θ is a vector of hyperparameters. The hyperparameters θ can be, for example, the variance in the Gaussian likelihood or the shape parameter in the likelihood of the gamma distribution. In the case of latent fields, they can be, for instance, dispersion parameters or spatial correlation parameters. [RMC09]

3.5.1 Notation and Basic Properties

For structured additive regression models, the distribution of the response variable y_i is assumed to be a member of the exponential family, with the mean μ_i linked to a structured additive predictor η_i by a link function $g(\cdot)$ such that $g(\mu_i) = \eta_i$. The predictor η_i takes into account the effect of multiple covariates in an additive way,

$$\eta_i = \alpha + \sum_{j=1}^{n_f} f^{(j)}(u_{ji}) + \sum_{k=1}^{n_\beta} \beta_k z_{ki} + \epsilon_i, \qquad i = 1, ..., n$$
(3.31)

[Sto85].

The $\left\{f^{(j)}\left(\cdot\right)\right\}$ s are unknown functions of the covariates u, while the $\left\{\beta_k\right\}$ s represent the linear effect of the covariates z and the ϵ_i s are unstructured terms. Latent Gaussian models assign a Gaussian prior to α , $\left\{f^{(j)}\left(\cdot\right)\right\}$ and $\left\{\epsilon_i\right\}$. In the following \boldsymbol{x} shall denote the vector of all latent Gaussian variables $\left(\left\{\eta_i\right\}, \alpha, \left\{f^{(j)}\right\}\right)$ and $\left\{\beta_k\right\}$)

and θ the vector of hyperparameters.

The conditional density $\pi\left(\boldsymbol{x}|\theta_{1}\right)$ is Gaussian with an assumed zero mean and precision matrix $\boldsymbol{Q}\left(\theta_{1}\right)$. The Gaussian density $\mathcal{N}\left(\mu,\boldsymbol{\Sigma}\right)$ with mean μ and covariance $\boldsymbol{\Sigma}$ at configuration \boldsymbol{x} is denoted by $\mathcal{N}\left(\boldsymbol{x};\mu,\boldsymbol{\Sigma}\right)$. For simplicity, $\{\eta_{i}\}$ has been included instead of $\{\epsilon_{i}\}$.

The distribution for the n_d observational variables $y = \{y_i : i \in \mathcal{I}\}$ is denoted by $\pi(\boldsymbol{y}|\boldsymbol{x},\theta_2)$ and is assumed conditionally independent given \boldsymbol{x} and θ_2 . Let $\boldsymbol{\theta} = \begin{pmatrix} \theta_1^T, \theta_2^T \end{pmatrix}^T$ with $\dim(\boldsymbol{\theta}) = m$. For non-singular $\boldsymbol{Q}(\boldsymbol{\theta})$ the posterior is given by

$$\pi\left(\boldsymbol{x},\boldsymbol{\theta}|\boldsymbol{y}\right) \propto \pi\left(\boldsymbol{\theta}\right) \pi\left(\boldsymbol{x}|\boldsymbol{\theta}\right) \prod_{i \in I} \pi\left(y_i|x_i,\boldsymbol{\theta}\right)$$

$$\propto \pi\left(\boldsymbol{\theta}\right) |\boldsymbol{Q}\left(\boldsymbol{\theta}\right)|^{1/2} \exp\left[-\frac{1}{2}\boldsymbol{x}^T \boldsymbol{Q}\left(\boldsymbol{\theta}\right) \boldsymbol{x} + \sum_{i \in I} \log\left\{\pi\left(y_i|x_i,\boldsymbol{\theta}\right)\right\}\right]. \tag{3.32}$$

Most latent Gaussian models satisfy two basic properties:

- 1. The latent field x is of large dimension, $n \approx 10^2 10^5$. Therefore, the latent field is a Gaussian Markov random field with sparse precision matrix $Q(\theta)$.
- 2. The number of hyperparameters, m, is small, $m \leq 6$.

In most cases, both properties are required to produce fast inference, and thus these will be assumed to be true for the remainder of this work [RMC09].

3.5.2 Applications for Latent Gaussian Models

Latent Gaussian models can be employed in a vast range of different domains, in fact most structured Bayesian models are of this particular form. Some of these domains are presented below.

3.5.2.1 Regression Models

Bayesian generalised linear models correspond to the linear relationship $\eta_i = \alpha + \sum_{k=1}^{n_\beta} \beta_k z_{ki}$ [DGM00]. Either the linear relationship of the covariates can be relaxed through the $f(\cdot)$ terms [FT13], random effects can be introduced through them or both. Smooth covariate effects are frequently modeled using penalised spline models [LB04] or random walk models [FT13], continuous indexed spline models [RH05] or Gaussian processes [CGW05]. The incorporation of random effects allows for the consideration of overdispersion caused by unobserved heterogeneity or correlation

in longitudinal data and can be introduced by defining $f(u_i) = f_i$ and $\{f_1\}$ to be independent, zero mean and Gaussian [FL01].

3.5.2.2 Dynamic Models

Temporal dependence can be introduced by using i in (3.31) as temporal index t and defining $f(\cdot)$ and \mathbf{u} such that $f(u_t) = f_t$. Both a discrete-time and a continuous-time autoregressive model can be modeled by $\{f_t\}$. Furthermore, a seasonal effect or the latent process of a structured time series model can be modeled [KG96]. Alternatively, a smooth temporal function in the same sense as for regression models can be represented by $\{f_t\}$.

3.5.2.3 Spatial and Spatio-Temporal Models

Similar to the previous type of model, spatial dependence can be modeled by a spatial covariate \boldsymbol{u} such that $f(u_s) = f_s$, where s denotes the spatial location or region s. The stochastic model for f_s is constructed to promote spacial smooth realisations of some sort. Popular models of this type include the Besag-York-Mollié [BYM91] model with extensions for regional data, continuous indexed Gaussian models [BCG14] and texture models [Mar+01]. The dependence between spatial and temporal covariates can be achieved either by using a spatio-temporal covariate (s,t) or a corresponding spatio-temporal Gaussian field [KW03].

Often the final model consists of a sum of several components, e.g. a spatial component, random effects and both linear and smooth effects of some covariates. In order to separate the effects of the different components in (3.31), sometimes linear or sum-to-zero constraints can be imposed [RMC09, pp. 319–321].

3.5.3 The MCMC Approach to Inference

The usual approach to inference for latent Gaussian models involves the previously introduced Markov chain Monte Carlo methods. Due to several factors, these methods may perform poorly when applied to such models. One factor is the interdependence of the components of the latent field \boldsymbol{x} while another is that $\boldsymbol{\theta}$ and \boldsymbol{x} are highly dependent on each other, especially for large n. The first of these problems can potentially be overcome by constructing a joint proposal based on a Gaussian approximation of the full conditional of \boldsymbol{x} [Gam97], while the second problem requires, at least in part, a joint update of $\boldsymbol{\theta}$ and \boldsymbol{x} . There are several

proposals to solve these shortcomings, but MCMC sampling continues to show poor computational speed [RMC09, p. 322].

3.5.4 Gaussian Random Fields

Let $\mathbf{s} = (s_1, ..., s_n)^T$ be a vector of locations. A Gaussian random field (GRF)

$$\left\{ Z(s) : s \in D \subset \mathbb{R}^2 \right\} \tag{3.33}$$

is a set of random variables where the observations occur in a continuous domain and where each finite set of random variables follows a multivariate normal distribution. A random process $Z(\cdot)$ is strictly stationary if it is invariant to shifts, i.e., if for each set of locations and each $h \in \mathbb{R}^2$ the distribution of $Z(s) = (Z(s_1), ..., Z(s_n))$ is equal to that of $Z(s+h) = (Z(s_1+h), ..., Z(s_n+h))$. A less constraining requirement is given by second-order stationarity. Under this condition, the process has a constant mean value

$$\mathbb{E}\left[\boldsymbol{Z(s)}\right] = \mu, \qquad \forall s \in D, \tag{3.34}$$

and the covariances depend only on the differences between locations

$$Cov(\mathbf{Z(s)}, \mathbf{Z(s+h)}) = C(\mathbf{h}), \quad \forall \mathbf{s} \in D, \forall \mathbf{h} \in \mathbb{R}^2.$$
 (3.35)

Furthermore, if the covariances depend only on the distances between the locations and not on the directions, the process is called isotropic. Else, the process is anisotropic. An intrinsically stationary process has a constant mean value and satisfies

$$\operatorname{Var}\left(Z\left(s_{i}\right)-Z\left(s_{j}\right)\right)=2\gamma\left(s_{i}-s_{j}\right), \quad \forall s_{i}, s_{j}. \tag{3.36}$$

 $2\gamma\left(\cdot\right)$ is the variogram and $\gamma\left(\cdot\right)$ is called the semivariogram [Cre15]. Under the assumption of intrinsic stationarity, the constant-mean assumption implies

$$2\gamma(\mathbf{h}) = \text{Var}(\mathbf{Z}(\mathbf{s} + \mathbf{h}) - \mathbf{Z}(\mathbf{s})) = \mathbb{E}\left[(\mathbf{Z}(\mathbf{s} + \mathbf{h}) - \mathbf{Z}(\mathbf{s}))^2 \right],$$

and the estimation of the semivariogram can be obtained using the empirical semivariogram as follows:

$$2\widehat{\gamma}\left(\boldsymbol{h}\right) = \frac{1}{|N\left(\boldsymbol{h}\right)|} \sum_{N\left(\boldsymbol{h}\right)} \left(Z\left(s_{i}\right) - Z\left(s_{j}\right)\right)^{2}, \tag{3.37}$$

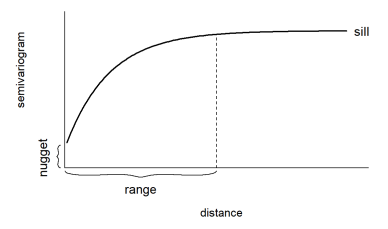


Fig. 3.3: A typical semivariogram

where $N(\mathbf{h}) = \{(s_1, s_j) : s_i - s_j = \mathbf{h}, i, j = 1, ..., n\}$ denotes the number of pairs and $|N(\mathbf{h})|$ the number of distinct pairs. For isotropic processes, the semivariogram is a function of distance $h = ||\mathbf{h}||$.

Plotting the empirical semivariogram against the separation distance conveys essential information regarding the continuity and spatial variability of the process. Given relatively short distances, the semivariogram tends to be small but increases with distance, indicating the similarity of observations in close proximity. The semivariogram levels off to a nearly constant value, also called the sill, as the separation distance increases, indicating a decrease in spatial dependence with distance within the range and no spatial correlation outside the range, which is reflected in a nearly constant variance. If there is a discontinuity or a vertical jump at the origin, the process has a nugget effect, which is often due to a measurement error, but may also be indicative of a spatially discontinuous process. The empirical semivariogram is an exploratory tool useful for assessing whether data exhibit spatial correlation. Furthermore, it can be compared to a Monte Carlo envelope of empirical semivariograms calculated from random permutations of the data while keeping the locations fixed [DRC03]. If the empirical semivariogram lies outside the Monte Carlo envelope with increasing distance, this is an indication of spatial correlation.

These graphs are again taken from "Geospatial Health Data: Modeling and Visualization with R-INLA and Shiny" by Paula Moraga [Mor19]. An example of a semivariogram is shown in Figure 3.3. This graph is taken from "Geospatial Health Data: Modeling and Visualization with R-INLA and Shiny" by Paula Moraga [Mor19].

The dependence structure of a GRF is given by the covariance matrix, which is constructed from a covariance function. Matern models and exponential functions are conventionally used for this purpose [Gel+10]. For the locations $s_i, s_j \in \mathbb{R}^2$ the exponential covariance function is given by

$$Cov(Z(s_i), Z(s_j)) = \sigma^2 \exp(-\kappa ||s_i - s_j||),$$
 (3.38)

where the distance between the locations s_i and s_j is denoted by $||s_i - s_j||$, the variance of the spatial field is given by σ^2 , while $\kappa > 0$ controls the rate at which the correlation decays as the distance increases.

The Matérn family represents a flexible class of covariance functions that arises naturally in a variety of scientific fields [GG06]. The Matérn covariance function is written as

$$Cov(Z(s_i), Z(s_j)) = \frac{\sigma^2}{2^{\nu-1}\Gamma(\nu)} (\kappa ||s_i - s_j||)^{\nu} K_{\nu} (\kappa ||s_i - s_j||).$$
 (3.39)

 σ^2 denotes the marginal variance of the spatial field, $K_{\nu}\left(\cdot\right)$ represents the modified Bessel function of second kind and order $\nu>0$, where ν is an integer. The mean square differentiability of the process is determined by ν and is usually fixed since it is difficult to identify in applications. For $\nu=0.5$, this covariance function is the equivalent of the exponential covariance function. $\kappa>0$ is related to the range ρ , which is defined as the distance at which there is approximately no correlation between two given points, $\rho=\sqrt{8\nu}/\kappa$ to be exact [Cam+13]. Examples of these two covariance functions are shown in Figure 3.4. These graphs are again taken from "Geospatial Health Data: Modeling and Visualization with R-INLA and Shiny" by Paula Moraga [Mor19].

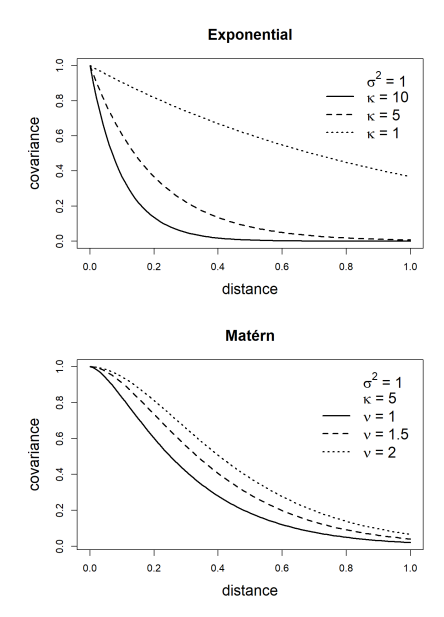


Fig. 3.4: Covariance functions corresponding to exponential and Matérn models.

3.5.5 Gaussian Markov Random Fields

3.5.5.1 Definition of GMRFs

Let $\boldsymbol{x}=(x_1,...,x_n)^T$ be normally distributed with mean $\boldsymbol{\mu}$ and covariance matrix $\boldsymbol{\Sigma}$. Let $\mathcal{G}=(\mathcal{V},\mathcal{E})$, where $\mathcal{V}=\{1,...,\}$ and \mathcal{E} be such that there is no edge between nodes i and j exactly when $x_i\perp x_j|\boldsymbol{x}_{ij}$. Then \boldsymbol{x} is a Gaussian Markov random field (GMRF) with respect to \mathcal{G} .

Since μ does not affect the pairwise conditional independence properties of x, this information is 'hidden' in Σ . Hence,

$$x_i \perp x_j | x_{ij} \Longleftrightarrow Q_{ij} = 0.$$

Therefore, the non-zero pattern of Q determines \mathcal{G} , i.e. whether x_i and x_j are conditionally independent, and can be derived from Q. If Q is a fully dense matrix, then \mathcal{G} is fully connected, implying that any normal distribution with SPD covariance matrix is a GMRF and vice versa.

The elements of Q are used for conditional interpretations. For any GMRF with respect to G = (V, E) with mean μ and precision matrix Q > 0,

$$\mathbb{E}\left[x_{i}|\boldsymbol{x}_{-i}\right] = \mu_{i} - \frac{1}{Q_{ii}} \sum_{j:j \sim i} Q_{ij} \left(x_{j} - \mu_{j}\right), \tag{3.40}$$

$$\operatorname{Prec}\left(x_{i}|\boldsymbol{x}_{-i}\right) = Q_{ii} \quad \text{and} \tag{3.41}$$

$$\operatorname{Corr}(x_i, x_j | \boldsymbol{x}_{ij}) = -\frac{Q_{ij}}{\sqrt{Q_{ii}Q_{jj}}}, \quad i \neq j$$
(3.42)

[RH05, p. 21].

On the main diagonal of ${\bf Q}$ are the conditional precisions of x_i given ${\bf x}_{-i}$ are placed, while the other elements, when scaled appropriately, provide information about the conditional correlation between x_i and x_j given ${\bf x}_{ij}$. Since ${\rm Var}\,(x_i) = \Sigma_{ii}$ and ${\rm Corr}\,(x_i,x_j) = \Sigma_{ij}/\sqrt{\Sigma_{ii}\Sigma_{jj}}$, the information about the marginal variance of x_i and the marginal correlation between x_i and x_j is given by ${\bf \Sigma}$. The marginal interpretation provided by the correlation matrix is intuitive and informative, as the scope of the interpretation is reduced from an n-dimensional distribution to a one-or two-dimensional distribution. ${\bf Q}$ is difficult to interpret marginally because either ${\bf x}_{-i}$ or ${\bf x}_{ij}$ would have to be integrated out of the joint distribution parameterized with respect to ${\bf Q}$. ${\bf Q}^{-1} = {\bf \Sigma}$ by definition, and in general Σ_{ii} depends on each element in ${\bf Q}$ and vice versa [RH05, pp. 20–23].

3.5.5.2 Markov Properties of GMRFs

One property of GMRFs is that more information regarding conditional independence can be extracted from \mathcal{G} . The following three properties are equivalent. The *pairwise Markov property*:

$$x_i \perp x_j | \boldsymbol{x}_{ij}$$
 if $\{i, j\} \notin \mathcal{E}$ and $i \neq j$.

The local Markov property:

$$x_i \perp \boldsymbol{x}_{-\{i, \text{ne}(i)\}} | \boldsymbol{x}_{\text{ne}(i)} \quad \forall i \in \mathcal{V}.$$

The *global Markov property*:

$$oldsymbol{x}_A \perp oldsymbol{x}_B | oldsymbol{x}_C$$

for all disjoint sets A, B and C where A and B are non-empty and separated by C. Illustrations for these properties are shown in Figure 3.5, Figure 3.6 and Figure 3.7. These illustrations are taken from Rue and Held [RH05, pp. 23–24].

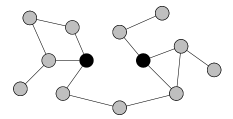


Fig. 3.5: The pairwise Markov property; the black nodes are conditionally independent given the light gray nodes.

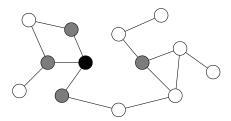


Fig. 3.6: The local Markov property; the black nodes and white nodes are conditionally independent given the dark gray nodes.

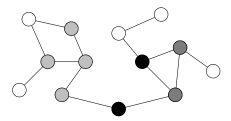


Fig. 3.7: The global Markov property; the dark gray and light gray nodes are globally independent given the black nodes.

3.5.5.3 Conditional Properties of GMRFs

An essential result of GMRFs is the conditional distribution for a subset x_a given x_{-A} . Here the canonical parameterisation proves useful, since by definition it can be easily updated by successive conditioning.

By splitting the indices into the non-empty sets A and B, of which the latter is equal to -A,

$$\boldsymbol{x} = \begin{pmatrix} \boldsymbol{x}_A \\ \boldsymbol{x}_B \end{pmatrix}. \tag{3.43}$$

The mean and the precision are divided accordingly,

$$\mu = \begin{pmatrix} \mu_A \\ \mu_B \end{pmatrix}, \text{ and } \mathbf{Q} = \begin{pmatrix} \mathbf{Q}_{AA} & \mathbf{Q}_{AB} \\ \mathbf{Q}_{BA} & \mathbf{Q}_{BB} \end{pmatrix}.$$
 (3.44)

The conditional distribution of $x_A|x_B$ is then a GMRF with respect to the subgraph \mathcal{G}^A with mean $\mu_{A|B}$ and precision matrix $\mathbf{Q}_{A|B} > 0$, where

$$\mu_{A|B} = \mu_A - Q_{AA}^{-1} Q_{AB} (x_B - \mu_B)$$
 (3.45)

and

$$Q_{A|B} = Q_{AA}.$$

Thus, the explicit knowledge of $\mathbf{Q}_{A|B}$ is available through \mathbf{Q}_{AA} , i.e. no calculation is required to obtain the conditional precision matrix. Moreover, the conditional mean depends only on the values of $\boldsymbol{\mu}$ and \mathbf{Q} in $A \cup \operatorname{ne}(A)$, since $Q_{ij} = 0 \ \forall j \ \text{inne}(i)$. For successive conditioning, the canonical parameterisation for GMRF is useful. A GMRF \boldsymbol{x} with respect to $\mathcal G$ and canonical parameters \boldsymbol{b} and $\mathbf Q > 0$ has the density

$$\pi\left(\boldsymbol{x}\right)\propto\exp\left(-1rac{1}{2}\boldsymbol{x}^{T}\boldsymbol{Q}\boldsymbol{x}+\boldsymbol{b}^{T}\boldsymbol{x}
ight).$$

The precision matrix is Q and the mean is $\mu = Q^{-1}b$. The canonical parameterisation is written as

$$oldsymbol{x} \sim \mathcal{N}_C \left(oldsymbol{b}, oldsymbol{Q}
ight)$$
 .

Furthermore,

$$\mathcal{N}\left(\boldsymbol{\mu}, \boldsymbol{Q}^{-1}\right) \Longleftrightarrow \mathcal{N}_{C}\left(\boldsymbol{Q}\boldsymbol{\mu}, \boldsymbol{Q}\right).$$

If the indices are partitioned into two non-empty sets A and B and x, b and Q are partitioned as in (3.43) and (3.44), then

$$\boldsymbol{x}_A | \boldsymbol{x}_B \sim \mathcal{N}_C \left(\boldsymbol{b}_A - \boldsymbol{Q}_{AB} \boldsymbol{x}_B, \boldsymbol{Q}_{AA} \right).$$
 (3.46)

Let $m{y}|m{x} \sim \mathcal{N}\left(m{x}, m{P}^{-1}\right)$ and $m{x} \sim \mathcal{N}_{C}\left(m{b}, m{Q}\right)$, then

$$x|y \sim \mathcal{N}_C(b + Py, Q + P)$$
. (3.47)

This allows the calculation of conditional densities with multiple sources of conditioning, e.g. conditioning on observed data and a subset of variables. Therefore, the canonical parameterisation can be repeatedly updated without explicitly calculating the mean until it is actually needed. The computation of the mean requires the solution of $Q\mu = b$, but only matrix-vector products are needed for updating the canonical parameterisation [RH05, pp. 25–27].

3.5.5.4 Specification Through Full Conditionals

Alternatively, a GMRF can be specified by the full conditionals $\{\pi(x_i|\boldsymbol{x}_{-i})\}$ in place of $\boldsymbol{\mu}$ and \boldsymbol{Q} . Suppose the full conditionals are given as normals with

$$\mathbb{E}\left[x_i|\boldsymbol{x}_{-i}\right] = \mu_i - \sum_{j:j\sim i} \beta_{ij} \left(x_j - \mu_j\right) \quad \text{and}$$
 (3.48)

$$\operatorname{Prec}\left(x_{i}|\boldsymbol{x}_{-i}\right) = \kappa_{i} > 0 \tag{3.49}$$

for i=1,...,n, for μ , κ and some $\{\eta_{ij}, i \neq j\}$. Evidently, \sim is implicitly defined by the non-zero terms of $\{\beta_{ij}\}$. For there to exist a joint density $\pi\left(\boldsymbol{x}\right)$ leading to these full conditional distributions, these full conditionals must be consistent. Since \sim is symmetric, it follows that if $\beta_{ij} \neq 0$, then $\beta_{ji} \neq 0$. If the entries of the precision matrix are chosen such that

$$Q_{ii} = \kappa_i$$
, and $Q_{ij} = \kappa_i \beta_{ij}$

and Q must be symmetrical, i.e.,

$$\kappa_i \beta_{ij} = \kappa_j \beta_{ji},$$

then \boldsymbol{x} is a GMRF with respect to a labelled graph $\mathcal{G}=(\mathcal{V},\mathcal{E})$ with mean $\boldsymbol{\mu}$ and precision matrix $\boldsymbol{Q}=(Q_{ij})$ [RH05, p. 27].

3.5.5.5 Multivariate GMRFs

A *multivariate GMRF* (MGMRF) is a multivariate extension of a GMRF that has proven useful in applications. Let x be a GMRF with respect to \mathcal{G} , then the Markov property implies that

$$\pi(x_i|\mathbf{x}_{-i}) = \pi(x_i|\{x_j: j \sim i\}).$$

 x_i is the value related to node i. Often the nodes have physical interpretations such as an administrative region of a country, which can be used to define the neighbours of node i. Let each of the n nodes have an associated vector \mathbf{x}_i of dimension p, resulting in a GMRF of size np. Such a GMRF is denoted by $\mathbf{x} = \left(\mathbf{x}_1^T, ..., \mathbf{x}_n^T\right)^T$. The Markov property with respect to the nodes is preserved, i.e.,

$$\pi\left(\boldsymbol{x}_{i}|\boldsymbol{x}_{-i}\right) = \pi\left(\boldsymbol{x}_{i}|\left\{\boldsymbol{x}_{j}: j \sim i\right\}\right),$$

where \sim is with respect to the same graph \mathcal{G} . Let $\boldsymbol{\mu} = \left(\boldsymbol{\mu}_1^T,...,\boldsymbol{\mu}_n^T\right)^T$ be the mean of \boldsymbol{x} , where $\mathbb{E}\left[\boldsymbol{x}_i\right] = \boldsymbol{\mu}_i$, and $\widetilde{\boldsymbol{Q}} = \left(\widetilde{\boldsymbol{Q}}_{ij}\right)$ its precision matrix, where each element of the matrix is a $p \times p$ matrix.

It follows that

$$oldsymbol{x}_i \perp oldsymbol{x}_j | oldsymbol{x}_{-ij} \Longleftrightarrow \widetilde{oldsymbol{Q}}_{ij} = oldsymbol{0}.$$

Formally, a random vector $\boldsymbol{x} = \left(\boldsymbol{x}_1^T,...,\boldsymbol{x}_n^T\right)^T$ with $\dim\left(\boldsymbol{x}_i\right) = p$, is called a MGMRF $_p$ with respect to $\mathcal{G} = \left(\mathcal{V} = \left\{1,....,n\right\},\mathcal{E}\right)$ with mean $\boldsymbol{\mu}$ and precision matrix $\widetilde{\boldsymbol{Q}} > 0$, exactly when its density has the form

$$\pi\left(\boldsymbol{x}\right) = \left(\frac{1}{2\pi}\right)^{np/2} \left|\widetilde{\boldsymbol{Q}}\right|^{1/2} \exp\left(-\frac{1}{2}\left(\boldsymbol{x} - \boldsymbol{\mu}\right)^{T} \widetilde{\boldsymbol{Q}}\left(\boldsymbol{x} - \boldsymbol{\mu}\right)\right)$$
$$= \left(\frac{1}{2}\right)^{np/2} \left|\widetilde{\boldsymbol{Q}}\right|^{1/2} \exp\left(-\frac{1}{2}\sum_{ij}\left(\boldsymbol{x}_{i} - \boldsymbol{\mu}_{i}\right)^{T} \widetilde{\boldsymbol{Q}}_{ij}\left(\boldsymbol{x}_{j} - \boldsymbol{\mu}_{j}\right)\right)$$

and

$$\tilde{\boldsymbol{Q}}_{ij} \neq 0 \Longleftrightarrow \{i,j\} \in \mathcal{E} \, \forall \, i \neq j.$$

A MGMRF_p is equivalent to a GMRF of dimension np with identical mean vector and precision matrix. Therefore, all results valid for a GMRF are also valid for a MGMRF_p, with modifications, since the graph for a MGMRF_p has size n and is defined with respect to $\{x_i\}$, while for a GMRF it has size np and is defined with respect to $\{x_i\}$.

The interpretation of \widetilde{Q}_{ii} and \widetilde{Q}_{ij} can be derived from the full conditional $\pi(x_i|x_{-i})$. The extensions of (3.40) and (3.41) are

$$\mathbb{E}\left[\boldsymbol{x}_{i}|\boldsymbol{x}_{-i}\right] = \boldsymbol{\mu}_{i} - \widetilde{\boldsymbol{Q}}_{ii}^{-1} \sum_{j:j \sim i} \widetilde{\boldsymbol{Q}}_{ij} \left(\boldsymbol{x}_{j} - \boldsymbol{\mu}_{j}\right)$$
(3.50)

$$\operatorname{Prec}(\boldsymbol{x}_{i}|\boldsymbol{x}_{-i}) = \widetilde{\boldsymbol{Q}}_{ii}. \tag{3.51}$$

In some applications, the full conditionals

$$\mathbb{E}\left[\boldsymbol{x}_{i}|\boldsymbol{x}_{-i}\right] = \boldsymbol{\mu}_{i} - \sum_{i:j\sim i} \boldsymbol{\beta}_{ij} \left(\boldsymbol{x}_{j} - \boldsymbol{\mu}_{j}\right)$$
(3.52)

$$\operatorname{Prec}\left(\boldsymbol{x}_{i}|\boldsymbol{x}_{-i}\right)=\boldsymbol{\kappa}_{i}>0,\tag{3.53}$$

In some applications, the full conditionals are used to define the MGMRF_p, for given $p \times p$ -matrices $\{\beta_{ij}, i \neq j\}$, $\{\kappa_i\}$, and vectors μ_i . Again, \sim is implicitly defined by the non-zero matrices $\{\kappa_i\}$. Similar requirements as for p=1 apply to the existence of the joint density: $\kappa_i\beta_{ij} = \beta_{ij}^T\kappa_j$ for $i \neq j$ and $\tilde{Q} > 0$. The $p \times p$ elements of \tilde{Q} are

$$\widetilde{m{Q}}_{ij} = egin{cases} m{\kappa}_i m{eta}_{ij} & i
eq j \ m{\kappa}_i & i = j \end{cases};$$

therefore $\tilde{\mathbf{Q}} > 0 \Longleftrightarrow (\mathbf{I} + (\boldsymbol{\beta}_{ij})) > 0$ [RH05, pp. 29–30].

3.5.6 Integrated Nested Laplace Approximation

An alternative to MCMC methods that is both less computationally intensive and suitable for performing approximate Bayesian inference in latent Gaussian models is *Integrated nested Laplace Approximation* (INLA). The basis of INLA is the use of a combination of analytical approximations and numerical algorithms for sparse matrices to approximate the posterior distribution using closed-form expressions. This speeds up inference and circumvents problems of sample convergence and mixing, making it suitable for fitting large data sets or exploring other models [RMC09].

INLA can be used for all models of the following form,

$$y_i | \boldsymbol{x}, \boldsymbol{\theta} \sim \pi \left(y_i | x_i, \boldsymbol{\theta} \right), \qquad i = 1, ..., n,$$

 $\boldsymbol{x} | \boldsymbol{\theta} \sim \mathcal{N} \left(\mu \left(\boldsymbol{\theta} \right), \boldsymbol{Q} \left(\boldsymbol{\theta} \right)^{-1} \right),$
 $\boldsymbol{\theta} \sim \pi \left(\boldsymbol{\theta} \right).$

As introduced in subsection 3.5.1, y are the observed data, x is a Gaussian field, θ represents the hyperparameters, while $\mu(\theta)$ and $Q(\theta)$ denote the mean and precision matrix respectively. To ensure fast inference, the dimension of the hyperparameter vector θ should be small, since the approximations are computed by numerical integration over the hyperparameter space.

In most cases, the observations y_i are assumed to belong to the exponential family with mean $\mu_i = g^{-1}\left(\eta_i\right)$. As shown in equation (3.31), η_i accounts for the effects of several covariates in an additive way, which makes it suitable for a wide range of models, including spatial and spatio-temporal models, since $\left\{f^{(j)}\right\}$ can take very different forms.

Let $\mathbf{x} = (\alpha, \{\beta_k\} | \theta \sim \mathcal{N}(\mu(\theta), \mathbf{Q}(\theta)^{-1}))$ be the vector of latent Gaussian variables, and let $\boldsymbol{\theta}$ be the vector of hyperparameters, which are not required to be Gaussian. INLA calculates accurate and fast approximations for the posterior marginals of the components of the latent Gaussian variables

$$\pi\left(x_{i}|\boldsymbol{y}\right), \qquad i=1,...,n,$$

as well as the posterior marginals for the hyperparameters of the latent Gaussian model

$$\pi\left(\theta_{i}|\boldsymbol{y}\right), \qquad j=1,...,\dim\left(\boldsymbol{\theta}\right).$$

For each element x_i of \boldsymbol{x} the posterior marginals are given by

$$\pi(x_i|\boldsymbol{y}) = \int \pi(x_i|\boldsymbol{\theta}, \boldsymbol{y}) \,\pi(\boldsymbol{\theta}|\boldsymbol{y}) \,d\boldsymbol{\theta}, \qquad (3.54)$$

and the posterior marginal for the hyperparameters can be expressed by

$$\pi\left(\theta_{j}|\boldsymbol{y}\right) = \int \pi\left(\boldsymbol{\theta}|\boldsymbol{y}\right) d\boldsymbol{\theta}_{-j}.$$
 (3.55)

 $\pi\left(x_i|\boldsymbol{y}\right)$ is approximated by combining analytical approximations to the full conditionals $\pi\left(x_i|\boldsymbol{\theta},\boldsymbol{y}\right)$ and $\pi\left(\boldsymbol{\theta}|\boldsymbol{y}\right)$ and numerical integration routines to integrate out $\boldsymbol{\theta}$. Similarly, $\pi\left(\theta_j|\boldsymbol{y}\right)$ is approximated by approximating $\pi\left(\boldsymbol{\theta}|\boldsymbol{y}\right)$ and integrating out $\boldsymbol{\theta}_{-j}$. In particular, the posterior density of $\boldsymbol{\theta}$ is obtained through Gaussian approximation for the posterior of the latent field, $\widetilde{\pi}_G\left(\boldsymbol{x}|\boldsymbol{\theta},\boldsymbol{y}\right)$, evaluated at the posterior mode, $x^*\left(\boldsymbol{\theta}\right) = \arg\max_{\boldsymbol{x}} \pi_G\left(\boldsymbol{x}|\boldsymbol{\theta},\boldsymbol{y}\right)$,

$$\widetilde{\pi}\left(\boldsymbol{\theta}|\boldsymbol{y}\right) \propto \frac{\pi\left(\boldsymbol{x},\boldsymbol{\theta},\boldsymbol{y}\right)}{\widetilde{\pi}_{G}\left(\boldsymbol{x}|\boldsymbol{\theta},\boldsymbol{y}\right)}\bigg|_{\boldsymbol{x}=\boldsymbol{x}^{*}\left(\boldsymbol{\theta}\right)}.$$
 (3.56)

Next, the following nested approximations are constructed,

$$\widetilde{\pi}\left(x_{i}|\boldsymbol{y}\right) = \int \widetilde{\pi}\left(x_{i}|\boldsymbol{\theta},\boldsymbol{y}\right) \widetilde{\pi}\left(\boldsymbol{\theta}|\boldsymbol{y}\right) d\boldsymbol{\theta}, \qquad \widetilde{\pi}\left(\theta_{j}|\boldsymbol{y}\right) = \int \widetilde{\pi}\left(\boldsymbol{\theta}|\boldsymbol{y}\right) d\boldsymbol{\theta}_{-j}. \tag{3.57}$$

Finally, these approximations are numerically integrated with respect to θ

$$\widetilde{\pi}(x_i|\boldsymbol{y}) = \sum_k \widetilde{\pi}(x_i|\theta_k, \boldsymbol{y}) \widetilde{\pi}(\theta_k|\boldsymbol{y}) \times \Delta_k,$$
(3.58)

$$\widetilde{\pi}\left(\theta_{j}|\boldsymbol{y}\right) = \sum_{l} \widetilde{\pi}\left(\theta_{l}^{*}|\boldsymbol{y}\right) \times \Delta_{l}^{*}, \tag{3.59}$$

with Δ_k and Δ_l^* representing the area weights corresponding to θ_k and θ_l^* . To obtain the approximations for the posterior marginals for the x_i 's condit

To obtain the approximations for the posterior marginals for the x_i 's conditioned on selected values of θ_k and $\widetilde{\pi}\left(x_i|\theta_k, \boldsymbol{y}\right)$, a Gaussian, Laplace or simplified Laplace approximation can be used. Using a Gaussian approximation derived from $\widetilde{\pi}_G\left(\boldsymbol{x}|\boldsymbol{\theta},\boldsymbol{y}\right)$ is the simplest and fastest solution, but in some situations it produces errors in the location and is unable to capture skewness behaviour. Therefore, the Laplace approximation is favoured over the Gaussian approximation, although it is relatively expensive. The simplified Laplace approximation is associated with lower costs and addresses inaccuracies of the Gaussian approximation in terms of location and skewness in a satisfactory manner [Mor19].

3.6 Bayesian Spatial Models

3.6.1 The Besag-York-Mollie Model

[BYM91]

3.6.1.1 The BYM2 Model

[Sim+17]

3.6.2 Besags' Proper Spatial Model

[Bes74]

3.6.3 The Leroux Model

[LLB00]

3.7 Goodness-of-Fit indicators

The goodness of fit indicates "how well" an estimated model can explain a set of observations. Measures of goodness of fit allow a statement to be made about the discrepancy between the theoretical values of the random variables under investigation, which are expected or predicted on the basis of the model, and the values actually measured.

The goodness of fit of a model to available data can be assessed with the help of statistical tests or suitable ratios.

3.7.1 The Akaike Information Criterion

The historically oldest criterion was proposed in 1973 by Hirotsugu Akaike (1927-2009) as an information criterion and is known today as the Akaike information criterion (AIC). The AIC is one of the most frequently used criteria for model selection in the context of likelihood-based inference.

Let the population contain the distribution of a variable with unknown density function p. The maximum likelihood estimation assumes a known distribution with an unknown parameter θ , hence the density function can be written as $q\left(\theta\right)$. The Kullback-Leibler divergence is used as a distance measure between p and $q\left(\widehat{\theta}\right)$ with $\widehat{\theta}$ the estimated parameter from the maximum likelihood estimation. The better the maximum likelihood model, the small the Kullback-Leibler divergence $D\left(P||Q\right)$.

For a maximum likelihood model with a p-dimensional parameter vector $\hat{\theta}$, the Akaike information criterion is defined as

$$AIC = -2l\left(\hat{\boldsymbol{\theta}}_{ML}\right) + 2p,\tag{3.60}$$

with l the log-likelihood function [Aka74].

3.7.2 The Deviance Information Criterion

In statistics, the deviance information criterion, or DIC for short, is a measure (criterion) for the prediction error of a model. This measure is an information criterion and belongs to the environment of the Bayesian method for model comparisons. The smaller the deviance information criterion, the better the model fit. The deviance information criterion can be regarded as the Bayesian equivalent of the Akaike

information criterion.

The deviance is defined as

$$D(\boldsymbol{\theta}) = -2\log(l(\boldsymbol{y}|\boldsymbol{\theta})) + C, \tag{3.61}$$

with y the data, θ the unknown parameters of the model and l the likelihood function. C is a constant that cancels out in all calculations that compare different models and therefore it does not need to be known [NW72].

The DIC is given by

$$DIC = D\left(\overline{\boldsymbol{\theta}}\right) + 2p_D, \tag{3.62}$$

with

$$p_D = \overline{D(\boldsymbol{\theta})} - D(\overline{\boldsymbol{\theta}}), \qquad (3.63)$$

where $\bar{\theta}$ is the expected value of θ [Spi+14].

3.7.3 The Watanabe-Akaike Information Criterion

The Watanabe-Akaike information criterion (WAIC) is the generalised AIC onto singular statistical models.

The WAIC is given by

$$WAIC = -2LLPD + 2p_{WAIC}, (3.64)$$

with the log pointwise predictive density (LLPD) given by

$$LLPD = \sum_{i=1}^{n} \log \left(\int p(y_i | \boldsymbol{\theta}) p_{post}(\boldsymbol{\theta}) \right) d\boldsymbol{\theta}.$$
 (3.65)

LLPD can be seen as the Bayesian analogue of $l\left(\widehat{\boldsymbol{\theta}}_{ML}\right)$ in the calculation of the AIC. The penalty term of the WAIC is also fully Bayesian and given by

$$p_{\text{WAIC}} = \sum_{i=1}^{n} \text{Var}_{\text{post}} \left(\log \left(p\left(y_{i} | \boldsymbol{\theta} \right) \right) \right), \tag{3.66}$$

where the term represents the variance of the individual terms in the LLPD over all data points [WO10; Yon18].

3.7.4 The Conditional Predictive Ordinate

The conditional predictive ordinate (CPO) is a Bayesian diagnostic that can be used to detect surprising observations. It is often used in the context of univariate sampling, the multivariate normal distribution and regression models.

The conditional predictive ordinate is given by

$$CPO = p(y_i|y_{-i}) \tag{3.67}$$

with y the data, y_{-i} the data without the i-th observation, and $p(\cdot|y_{-i})$ the predictive distribution of a new observation at y_{-i} . Low values of CPO are an indication that y_i is surprising given prior knowledge and the other observations [Pet90; Cox80].

Analysis of Geospatial Health

Data

4.1 Geographic Data

In spatial statistics, two fundamental types of geographic data exist, namely *vector data* and *raster data*. In the vector data model, the world is represented by points, lines and polygons with discrete, well-defined boundaries, which tends to result in high accuracy. Raster data, on the other hand, divides the surface into cells of uniform size, and raster datasets are used as the basis for background images in web mapping.

Determining which data type to use depends on the domain of the application. Vector data dominates in the social sciences because human settlements typically have discrete boundaries, while raster data are commonly used in many environmental sciences because they are based on remote sensing data. Naturally, there is also some overlap and both types can be used together or one form can be converted into the other [LNM19].

4.1.1 Vector Data

The geographic vector data model is based on points located within a *coordinate* reference system (CRS), in which points either represent self-standing features or form more complex geometric shapes, i.e. lines and polygons. Using this system, Trondheim can be represented by the coordinates (10.4, 63.4), meaning 10.4 degrees east of the prime meridian and 63.4 degrees north of the equator. It could also be written as (1157722.70, 9199010.75), which is the position of Trondheim using the Web Mercator projection, the de facto standard for web mapping applications. More will be said about CRS later, but for now it is sufficient to know that it is possible to display coordinates in various ways. An example of a CRS is shown in Figure 4.1.

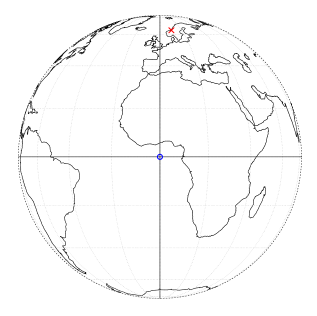


Fig. 4.1: A geographic CRS with an origin at 0° longitude and latitude. The red X denotes the location of Trondheim.

Different Types of Vector Data

As mentioned earlier, there are different types of vector data. There are 17 different geometry types in the standard *simple features*, but there are seven core types that can be used in most analysis software. These types are visualised in Figure 4.2.

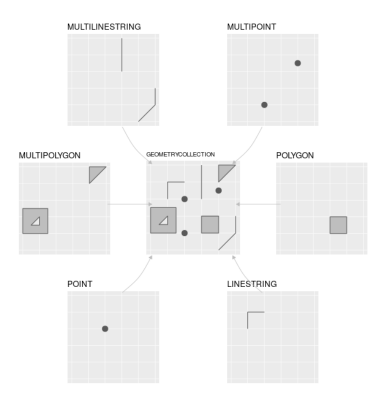


Fig. 4.2: The most commonly used simple feature types.

Simple Features was developed by the Open Geospatial Consortium and is an open, standardised, hierarchical data model that represents a wide range of geometry types. The use of this data model ensures that scientific work can be transferred to other institutions, e.g. when importing from and exporting to spatial databases [LNM19].

4.1.2 Raster Data

The geographic raster data model consists in most cases of a raster header and a matrix representing uniformly distributed cells/pixels. The raster header defines the CRS, the origin (starting point) and the extent. Since the number of columns and rows and the resolution of the cell size are stored in the extent, starting from the origin, it is easy to access and change each cell by its ID or by specifying the row and column number. In this type of representation, the coordinates of the four vertices of each cell are not explicitly stored, instead only the origin is stored. This speeds up data processing and makes it more efficient, but each raster layer can only contain a single value, which can be either numeric or categorical. Typically, raster maps are

used to depict continuous features such as elevation or temperature, but categorical variables, for example soil or land cover, as shown in Figure 4.3 [LNM19].

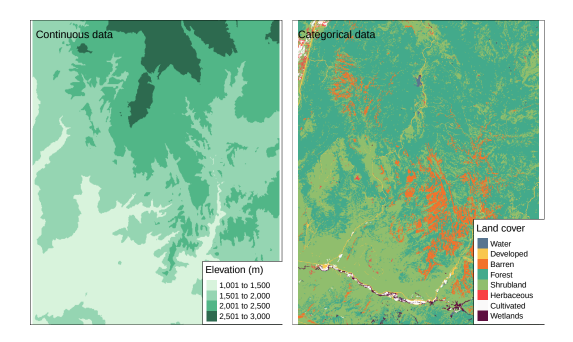


Fig. 4.3: An example of continuous and categorical raster data

Coordinate Reference Systems

A common denominator of vector and raster data are that both use the coordinate reference system (CRS), which defines how spatial elements relate to the surface of the Earth. The CRS can be either geographic or projected.

Geographic Coordinate Systems

Geographic coordinate systems use two values, *longitude* and *latitude*, to identify any location on Earth. Longitude is defined as the east-west location at an angular distance from the prime meridian plane, while latitude is the angular distance north or south of the equator. Consequently, distances in geographic CRS are not measured in metres.

The Earth's surface is typically represented in geographical coordinate systems by

a spherical or ellipsoidal surface. The former assumes that the Earth is a perfect sphere of a certain radius, which has the advantage of being a simplistic model, but is associated with inaccuracies owing to the fact that the Earth is not a sphere. Ellipsoidal models are defined by the equatorial radius and the polar radius, providing a better model since the equatorial radius is approximately 11.5 km longer than the polar radius.

The *datum* is a broader component of CRS that contains information about which ellipsoid to use and the exact relationship between Cartesian coordinates and the location on the Earth's surface. The notation *proj4string* is used to store these additional details. It allows for local variations of the Earth's surface, such as large mountain ranges, to be taken into account in local CRS. Datum can again be divided into two categories, *local* and *geocentric*, the difference being that in the local datum the ellipsoidal surface is shifted to match the surface at a particular location, whereas in the geocentric datum the centre of gravity of the Earth is the centre and the accuracy of the projections is not optimised for any particular location [LNM19].

Projected Coordinate Systems

Projected CRS are based on Cartesian coordinates on an implicitly flat surface and have an origin, x and y axes, and a linear unit of measurement, metres for instance. They are based on geographic CRS and rely on map projections to convert between the three-dimensional surface of the Earth and the east/north values (x and y) in a projected CRS.

This transition always entails some distortion, skewing some of the properties of the earth's surface, such as area, direction, distance and shape. Generally, the name of a projection is based on a property it preserves, e.g. equal area projection preserves area, equidistant projection preserves distance and conformal projection preserves local shape.

Again, subgroups exist in projection coordinate systems, *conic*, *cylindrical* and *planar* projections. In a conic projection, the earth's surface is projected onto a cone along one or two tangent lines. Along these lines the distortions are minimised and increase with the distance to the lines. The projection is therefore best suited for maps of mid-latitude areas. Cylindrical projections map the surface onto a cylinder. These types of projections can be created by touching the surface of the Earth along one or two tangent lines. They are often used to map the entire Earth. A planar projection projects data onto a flat surface that touches the globe at a point or along a tangent line, and is typically used in mapping polar projections [LNM19].

4.2 Spatial Point Processes

A stochastic process that describes the location of particular events/points that occur in a region is known as a point process. The number of points as well as the location of the points are random. An example of a point process would be the number of earthquakes and their locations.

4.2.1 Fundamentals of Point Processes

Let Z be a random, at most countable set of points in a space \mathbb{X} , for example \mathbb{R}^d . Ignoring measurability issues, Z can be thought of as a mapping $\omega \mapsto Z(\omega)$ from Ω into the set of countable subsets of \mathbb{X} , where $(\Omega, \mathcal{F}, \mathbb{P})$ defines an underlying probability space. Z can then be identified with the family of mappings

$$\omega \mapsto \eta(\omega, B) := \operatorname{card}(z(\omega) \cap B), \qquad B \subset \mathbb{X},$$
 (4.1)

which counts the number of points from Z in B. For any fixed $\omega \in \Omega$, $\eta(\omega, \cdot)$ is the counting measure supported by $Z(\omega)$ [CI80].

For a general definition of a point process, let $(\mathbb{X}, \mathcal{X})$ be a measurable space and let $N_{<\infty}(\mathbb{X}) \equiv N_{<\infty}$ be the space of all measures μ on \mathbb{X} such that $\mu(B) \in \mathbb{N}_0 := \mathbb{N} \cup \{0\} \, \forall B \in \mathcal{X}$. Let $N(\mathbb{X}) \equiv N$ be the space of all measures describable as a countable sum of measures from $N_{<\infty}$, for example the zero measure 0 which is equal to 0 on \mathcal{X} . In general, any sequence $(x_n)_{n=1}^k$ of elements of \mathbb{X} , where $k \in \overline{\mathbb{N}} := \mathbb{N} \cup \{\infty\}$ denotes the number of terms in the sequence, can be used to define a measure

$$\mu = \sum_{n=1}^{k} \delta_{x_n}.$$

$$\Rightarrow \mu(B) = \sum_{n=1}^{k} \mathbf{1}_B(x_n), \qquad B \in \mathcal{X}.$$
(4.2)

More generally, for any measurable $f: \mathbb{X} \to [0, \infty]$,

$$\int f d\mu = \sum_{n=1}^{k} f(x_n) \tag{4.3}$$

For k=0 in (4.2), μ is equal to the zero measure. The point set $\mathbf{x}=(x_1,...,x_n)^T$ is said to be not pairwise different and if $x_i=x_j$ with $i\neq j$, μ is said to have multiplicities. The multiplicity of x_i is equal to the number

card
$$\{j \le k : x_j = x_i\}$$
.

Any μ of the form (4.2) is interpreted as a counting measure with possible multiplicities, but in general it cannot be guaranteed that every $\mu \in N$ can be written in this particular form.

A point process η on $\mathbb X$ is called *proper point process* if random elements $X_1, X_2, ...$ exist $\in \mathbb X$ and a $\overline{\mathbb N}_0$ -valued random variable κ such that almost surely

$$\eta = \sum_{n=1}^{\kappa} \delta_{X_n}.\tag{4.4}$$

For $\kappa = 0$ this is the zero measure on \mathbb{X} .

This terminology is motivated by the intuition that a point process is a (random) set of points, rather than an integer measure. A proper point process fits this intuition better, since it can be interpreted as a countable set of points in \mathbb{X} [LP17, pp. 9–12].

4.2.2 Poisson Processes

Poisson processes are defined by the fact that the number of points in a given set follows a Poisson distribution. Furthermore, the numbers of points in disjoint sets are stochastically independent.

In application, Poisson processes are used in a wide range of fields, including biology, economics and image processing.

Let λ be an s-finite measure on \mathbb{X} . Let a *Poisson process* with intensity measure λ be defined as a point process η on \mathbb{X} with the following two properties:

1.
$$\forall B \in \mathcal{X} : \eta(B) \sim \text{Po}(\lambda(B); k) \ \forall k \in \mathbb{N}_0 \iff \mathbb{P}(\eta(B) = k)$$

2. $\forall m \in \mathbb{N}$ and all pairwise disjoint sets $B_1, ..., B_m \in \mathcal{X}$: the random variables $\eta(B_1), ..., \eta(B_m)$ are independent.

A point process satisfying the second of these conditions is called *completely independent*. If η is a Poisson process with intensity measure λ , then

$$\mathbb{E}\left[\eta\left(B\right)\right] = \lambda(B). \tag{4.5}$$

For the zero measure,

$$\mathbb{P}\left(\eta(\mathbb{X})=0\right)=1,$$

with $\lambda = 0$ [LP17, p. 19].

4.3 Modeling and Visualising Health Data

4.3.1 Areal Data

Areal or lattice data are the result of segmenting a fixed domain into a finite number of sub-regions where results are aggregated, e.g. the number of infections with a specific disease in districts or the number of overweight people in provinces. Often the aim of disease risk models is to assess the risk within the same areas for which data are available. This can be done with a simple measure such as the *standardised incidence ratio* (SIR) or by using a Bayesian hierarchical model, which allows information to be drawn from neighbouring areas and incorporates covariates, thereby smoothing and reducing extreme values.

A widely used model is the *Besag-York-Mollié* (BYM) [BYM91], which takes spatial correlation and the potential for observations in neighbouring areas to be more similar than those in distant regions into account. It includes a spatial random effect that smoothes the data according to a neighbourhood structure, and an unstructured exchangeable component that models uncorrelated noise. In settings where disease numbers are monitored over time, spatio-temporal models account for temporal correlations in addition to spatial correlation, while also accounting for spatio-temporal interactions [Mor19].

4.3.1.1 Spatial Neighbourhood Matrices

Spatial or proximity matrices are useful for exploratory analysis of area data. Let w_{ij} denote the (i,j) element of a *spatial neighbourhood matrix* \boldsymbol{W} . w_{ij} connects the two areas in some spatial way. The neighbourhood structure over the complete study region is defined by \boldsymbol{W} , and the elements of the matrix can be considered as weights. The closer j is to i, the more weight is associated with it. The simplest neighbourhood definition is given by the binary matrix

$$w_{ij} = \begin{cases} 1 & \text{if regions } i \text{ and } j \text{ share a border} \\ 0 & \text{else} \end{cases}$$
 (4.6)

Since a region cannot share a boundary with itself, $w_{ii} = 0$ [Mor19].

In Figure 4.4, the number of shared borders of each canton in Switzerland are mapped.

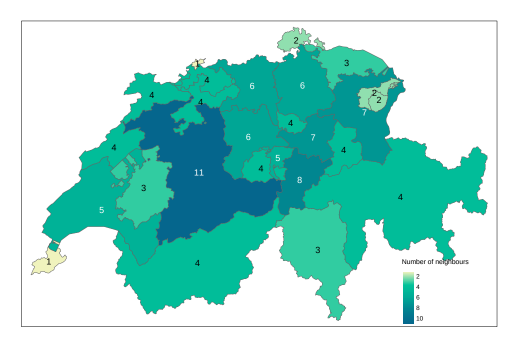


Fig. 4.4: The number of shared borders of cantons in Switzerland

4.3.1.2 Standardised Incidence Ratio

A basic measure of disease risk is the *standardised incidence ratio*, which yields an estimate in each of the areas that form a partition of the study region. It is defined as the ratio of observed counts to expected counts

$$SIR_i = \frac{Y_i}{E_i}. (4.7)$$

 E_i represents the sum of the expected number of cases of a given area i that behave according to the way the standard population behaves. It is calculated using indirect standardisation as

$$E_i = \sum_{j=1}^m r_j^{(s)} n_j^{(i)}, \tag{4.8}$$

with $r_j^{(s)}$ the rate in stratum j in the standard population and $n_j^{(i)}$ the population in stratum j of area i. If the stratum information is unavailable, the expected counts can be calculated as follows

$$E_i = r^{(s)} n^{(i)},$$

where $r^{(s)}$ denotes the rate in the standard population and $n^{(i)}$ is the population of area i. If the standardised incidence rate is greater than 1, area i has a higher risk than expected from the standard population, while for $SIR_i = 1$ the risk is the same

and for $SIR_i < 1$ it is lower than expected. The ratio is also called the standardised mortality ratio when applied to mortality data [Mor19].

4.3.1.3 Spatial Small Area Disease Risk Estimation

While SIRs may prove useful in some situations, in areas with low population sizes or rare diseases, expected counts may be low, making SIRs insufficiently reliable for reporting. It is therefore preferable to assess disease risk using models that allow information to be borrowed from neighbouring areas and incorporate information from covariates, thus smoothing or shrinking extreme values due to small sample sizes [Gel+10].

The observed counts Y_i in area i are typically modeled with a Poisson distribution with mean $E_i\theta_i$, where E_i is the expected counts and θ_i denotes the relative risk in area i. To account for extra Poisson reliability, the logarithm of the relative risk is expressed as the total of the intercept and the random effects. θ_i quantifies whether area i has a higher $(\theta_i>1)$ or lower $(\theta_i<1)$ risk than the average risk in the standard population. If the risk of an area i is half the average risk, then $\theta_i=0.5$. The general model for spatial data is formulated as follows:

$$Y_i \sim \text{Po}(E_i \theta_i), \quad i = 1, ..., n,$$
 (4.9)

$$\log\left(\theta_{i}\right) = \alpha + u_{i} + v_{i}.\tag{4.10}$$

The overall risk in the region of study is represented by α , u_i is a random effect specific to each area to model the spatial dependence between relative risks, and v_i is an unstructured exchangeable component that models uncorrelated noise, $v_i \sim \mathcal{N}\left(0,\sigma_v^2\right)$. Covariates are often included to measure risk factors and other random effects to deal with different sources of variability. For example,

$$\log\left(\theta_{i}\right) = \boldsymbol{d}_{i}\boldsymbol{\beta} + u_{i} + v_{i},$$

with $d_i = (1, d_{i1}, ..., d_{ip})$ a vector of the intercept and p covariates corresponding to the area i and $\boldsymbol{\beta} = (\beta_0, ..., \beta_p)^T$ the vector of coefficients. An increase in d_j (j = 1, ..., p) by one unit, leads to an increase in the relative risk by a factor of $\exp(\beta_i)$, provided that all other covariates remain constant.

In the Besag-York-Mollié (BYM) [BYM91] model, this spatial random effect u_i is assigned a conditional autoregressive (CAR) distribution that smooths the data

according to a given neighbourhood structure that defines two areas as neighbours if they share a common boundary. Specifically,

$$u_i | \boldsymbol{u}_{-i} \sim \mathcal{N}\left(\overline{u}_{\delta_i}, \frac{\sigma_u^2}{n_{\delta_i}}\right),$$
 (4.11)

where $\overline{u}_{\delta_i}^{-1} = n_{\delta_i}^{-1} \sum_{j \in \delta_i} u_j$, while δ_i and n_{δ_i} represent the set and the amount of neighbours of the area i, respectively. The unstructured component v_i is modeled as an independent and identically distributed (i.i.d.) normal variable with zero mean and variance σ_v^2 .

In 2017, Simpson et al. proposed BYM2 [Sim+17], a new parameterisation of the BYM model that yields interpretable parameters and facilitates the assignment of meaningful penalised complexity priors. It uses a scaled, spatially structured component u_* and an unstructured component v_* ,

$$\boldsymbol{b} = \frac{1}{\sqrt{\tau_b}} \left(\sqrt{1 - \phi} \boldsymbol{v_*} + \sqrt{\phi} \boldsymbol{u_*} \right). \tag{4.12}$$

The precision parameter $\tau_b > 0$ controls the marginal variance contribution of the weighted sum of u_* and v_* . The mixing parameter $0 \le \phi \le 1$ captures the proportion of the marginal variance explained by the structured effect u_* . Therefore, the BYM2 model is equal to a pure spatial model for $\phi = 1$ and equal to unstructured spatial noise for $\phi = 0$ [Rie+16]. To define the prior for the marginal accuracy τ_b , the following probability statement is used:

$$\mathbb{P}\left(\frac{1}{\sqrt{\tau_b}} > U\right) = \alpha$$

$$\iff \mathbb{P}\left(\phi < U\right) = \alpha \tag{4.13}$$

[Mor19].

DEFINE U AND ALPHA AND CONNECT IT TO PC PRIORS. ALSO ADD THE LEROUX AND BESAG MODEL

4.3.1.4 Spatio-Temporal Small Area Disease Risk Estimation

When disease counts are monitored over time, spatio-temporal models are useful as they take into account not only the spatial structure but also temporal correlations and spatio-temporal interactions [MLB08]. Let Y_{ij} be the counts observed in area i

and at time j, θ_{ij} be the relative risk, E_{ij} be the expected number of cases in area i and at time j, then

$$Y_{ij} \sim \text{Po}(E_{ij}\theta_{ij}), \qquad i = 1, ..., I, j = 1, ..., J.$$
 (4.14)

 $\log{(\theta_{ij})}$ is written as the sum of several components, including spatial and temporal structures, to consider that neighbouring areas and successive times may have similar risk. Spatio-temporal interactions can be included to account for the fact that temporal trends may differ from area to area but may be more alike in neighbouring areas.

Bernardinelli et al. [Ber+95], for example, propose a spatio-temporal model with parametric time trends that expresses the logarithm of relative risks as

$$\log(\theta_{ij}) = \alpha + u_i + v_i + (\beta + \delta_i) \times t_j. \tag{4.15}$$

The intercept is denoted by α , u_i+v_i is a random area effect, β represents a global linear trend effect and δ_i is an interaction between space and time which is the difference between β and the area-specific trend. For modeling u_i and δ_i , a CAR distribution is used and v_i is i.i.d.. This specification allows each of the areas to have its individual time trend, where the spatial intercept is given by $\alpha+u_i+v_i$ and the slope by $\beta+\delta_i$. δ_i is referred to as the differential trend of the i-th area and represents the amount by which the time trend of area i deviates from the overall time trend β . If $\delta_i \neq 0$, then area i has a time trend with a slope that is either steeper or less steep than the overall time trend β .

For models that do not demand linearity of the time trend, non-parametric models such as the one proposed by Knorr-Held [Kno00] can be used. This specific model incorporates spatial effects, temporal random effects and an interaction between space and time as follows:

$$\log(\theta_{ij}) = \alpha + u_i + v_i + \gamma_j + \phi_j + \delta_{ij}. \tag{4.16}$$

The intercept is again denoted by α , $u_i + v_i$ is a spatial random effect defined as before, i.e. u_i follows a CAR distribution and v_i is i.i.d.. $\gamma_j + \phi_j$ represents a temporal random effect and γ_j follows either a first order random walk in time (RW1)

$$\gamma_j | \gamma_{j-1} \sim \mathcal{N}\left(\gamma_{j-1}, \sigma_{\gamma}^2\right),$$
 (4.17)

or second order random walk in time (RW2)

$$\gamma_j | \gamma_{j-1}, \gamma_{j-2} \sim \mathcal{N}\left(2\gamma_{j-1} - \gamma_{j-2}, \sigma_{\gamma}^2\right).$$
 (4.18)

The unstructured temporal effect is given by $\phi_j \stackrel{i.i.d.}{\sim} \mathcal{N}\left(0,\sigma_\phi^2\right)$. The interaction between space and time, δ_{ij} , can be specified in a number of ways by combining the structure of the random effects that interact. The interactions proposed by Knorr-Held are those between the effects (u_i,γ_j) , (u_i,ϕ_j) , (v_i,γ_j) and (v_i,ϕ_j) [Kno00]. Using the last of these interactions leads to the assumption that there is no spatial or temporal structure on δ_{ij} . Thus, the interaction term can be modeled as $\delta_{ij} \sim \mathcal{N}\left(0,\sigma_\delta^2\right)$ [Mor19].

4.3.1.5 Issues With Areal Data

The analysis of spatially aggregated data is subject to the "misaligned data problem" (MIDP), which arises when the data to be analysed is at a different scale from that at which it was collected [BCG14]. This may be solely due to the fact that the aim is to obtain the spatial distribution of a variable at a new spatial level of aggregation, e.g. if predictions are to be made at the county level with data that was originally collected at the postcode level. Another objective may be to try to find an association between variables available at different spatial scales, e.g. determining whether the risk of an unfavourable outcome provided at the country level correlates with exposure to an environmental pollutant measured at different stations, taking into account the population at risk and other demographic information available at the postcode level.

The Modifiable Area Unit Problem (MAUP) [Ope84] describes a problem where the inference may differ when the same underlying data are grouped at a new spatial level of aggregation. It consists of two interrelated effects, the first of which is the scale/aggregation effect. It relates to the different conclusions obtained when the same data are grouped into larger and larger areas. The other effect is the grouping/zoning effect, which accounts for the variability in results due to alternative formations of the areas, resulting in differences in area shape given the same or similar scales.

Ecological studies are defined by their reliance on aggregated data [Rob09] and the inherent potential for ecological fallacies. This phenomenon occurs when estimated associations obtained from the analysis of variables measured at the aggregate level lead to conclusions that differ from analyses based on the same variables measured at the individual level. This can be considered a special case of MAUP and the resulting so-called ecological bias is composed of two effects similar to the aggregation and zoning effects in MAUP. Namely, the aggregation bias caused by the aggregation of individuals and the specification bias due to the different distribution of confounding variables that results from the aggregation [GY02; Mor19].

4.3.2 Geostatistical Data

Geostatistical data are measurements of one or more spatially continuous features collected at specific locations. They can be a disease risk measured by a survey in different villages, the level of a pollutant recorded at several monitoring stations, or the density of mosquitoes responsible for disease transmission measured by traps set at different locations [WG04]. Let $Z(s_1),...,Z(s_n)$ be the observations of a spatial variable Z at locations $s_1,...,s_n$. Geostatistical data are often assumed to be partial realisations of a random process

$$\left\{ Z\left(s\right):s\in D\subset\mathbb{R}^{2}\right\} ,\tag{4.19}$$

where D denotes a fixed subset of \mathbb{R}^2 and the spatial index \mathbf{s} varies continuously over D. For practical reasons, it is only possible to observe $Z(\cdot)$ at a finite set of locations. The inference of the characteristics, e.g. mean and variability of the process, of the spatial process is based on this partial realisation. Using these characteristics, it is possible to predict the process at unobserved locations and construct a spatially continuous surface of the variable of interest.

4.3.2.1 Stochastic Partial Differential Equation Approach

With geostatistical data, an underlying spatially continuous variable can often be assumed and modelled using a Gaussian random field. A spatial model can be fitted using the stochastic partial differential equation (SPDE) approach and the variable of interest can be predicted at new locations. A GRF with a Matérn covariance matrix can be written as a solution to the following continuous domain SPDE [Whi63]:

$$\left(\kappa^{2} - \Delta\right)^{\alpha/2} (\tau x(\mathbf{s})) = \mathcal{W}(\mathbf{s}). \tag{4.20}$$

The GRF is represented by x (s), where smoothness is controlled by α , while $\mathcal{W}(s)$ denotes a Gaussian spatial white noise process. $\kappa>0$ is a scale parameter and Δ denotes the Laplacian given by $\sum_{i=1}^d \frac{\partial^2}{\partial x_i^2}$, where d is the dimension of the spatial domain D.

The smoothness parameter ν of the Matérn covariance function is linked to the SPDE by

$$\nu = \alpha - \frac{d}{2}$$

while the marginal variance σ^2 is related to the SPDE by

$$\sigma^{2} = \frac{\Gamma(\nu)}{\Gamma(\alpha) (4\pi)^{d/2} \kappa^{2\nu} \tau^{2}}.$$

For d=2 and $\nu=0.5$ this corresponds to the exponential function.

The SPDE can be solved approximately using the *finite element* method, which partitions the spatial domain D into a set of non-intersecting triangles, resulting in a triangulated mesh with n vertices and n basis functions $\psi_k\left(\cdot\right)$. These functions are piecewise linear functions on each triangle, equal to 1 at vertex k and 0 otherwise. The continuously indexed Gaussian field x is thus represented as a discretely indexed Gaussian Markov random field by the finite basis functions defined on the triangulated mesh

$$x(\mathbf{s}) = \sum_{k=1}^{n} \psi_k(\mathbf{s}) x_k, \tag{4.21}$$

with n the number of vertices of the triangulate, $\psi_k\left(\cdot\right)$ the piecewise linear basis functions and $\{x_k\}$ zero-mean Gaussian distributed weights.

The joint distribution of the weight vector follows a Gaussian distribution, $\mathbf{x} = (x_1, ..., x_n) \sim \mathcal{N}\left(0, \mathbf{Q}^{-1}(\tau, \kappa)\right)$, which approximates the solution $x(\mathbf{s})$ of the SPDE in the mesh nodes, and the basis functions transform $x(\mathbf{s})$ from the mesh nodes to the other spatial locations of interest [LRL11].

Dataset Collection

As is often the case with statisticians, the construction of the dataset used to analyse a research question is an essential task and frequently involves the merging of multiple data sources to create a dataset. This was the case in this thesis and in the following chapter a brief overview of the data sources used, their pre-processing and how they were combined is given.

5.1 Covid-19 Data

5.1.1 Covid-19 Data for Norway

The Covid-19 data for Norway comes from a dataset made available to the public via the a repository on the website Github.com, created by the user thohan88. The repository contains a daily updated dataset that is the result of combining several data sources, which include the Institute of Public Health and the Norwegian Directorate of Health. According to the author of the repository, the project is "an open-source effort to make data about the Covid-19 situation in Norway available to the public in a timely and coherent manner" [Han20].

A few sample data points from this dataset are displayed in Table 5.1.

Tab. 5.1: An excerpt from the Covid-19 data for Norway. Does not contain all variables.

kommune_no	kommune_name	population	2020-03-26	2020-03-27
1103	Stavanger	143574	87	88
1507	Ålesund	66258	20	20
4601	Bergen	283929	231	248
5001	Trondheim	205163	113	136

5.1.2 Covid-19 Data for Germany

In Germany, the Robert Koch Institute publishes daily situation reports in which the number of new cases is published at NUTS 3 level. These reports are available as pdf files via the Institute's website. They can be downloaded and grouped via the R package covid19germany[Sch+21], as was done for this work.

A few sample data points from this dataset are displayed in Table 5.2.

The variable *CumNumberTestedIll* contains the cumulative number of people that have tested positive for Covid-19.

Tab. 5.2: An excerpt from the Covid-19 data for Germany. Does not contain all variables.

Landkreis	Date	CumNumberTestedIll	population
SK München	2020-01-29	1	1471508
SK München	2020-02-03	2	1471508
SK München	2020-02-11	3	1471508
LK Rosenheim	2020-02-29	1	260983
LK Rosenheim	2020-03-08	2	260983
LK Rosenheim	2020-03-10	6	260983

5.2 Demographic Data

As demographics tend to differ between different geographic units, the decision was made to include demographic variables in the analysis of the research question to see if the risk for infection may be higher when a certain characteristic is present in the population.

5.2.1 Demographic Data for Norway

The demographic data collected for Norway comes from Statistisk Sentralbyrå and is made available to the public through their online database, StatBank[@Sen16]. The first characteristic collected was the age of the population in a given municipality. For each age, starting at 0 and ending at 105, the number of people of that age is known

Next, unemployment data were collected for a given municipality. For each municipality, the percentage of all people out of work is known, as well as the percentage of all immigrants out of work.

Other data collected include data related to the number of workers in a particular industry, as well as immigration data. Since there is discussion about whether workers from certain industries, in this case construction, contribute to the spread of Covid-19, the decision was made to collect this type of data. For each community, the number of workers across all industries is known, as well as the number of workers in the construction industry. Workers, in this case, are individuals employed in a given municipality who are between the ages of 20 and 66. It is also known how many people work full-time and how many work part-time.

Finally, for immigration data, it is known how many immigrants live in a given municipality and how many Norwegians were born to immigrant parents. These figures are known in terms of the percentage of the population in 2020.

5.2.2 Demographic Data for Germany

The demographic data collected for Germany comes from the federal and state statistical offices and is made available to the public through their online database, Regionaldatenbank Deutschland [@Bun20].

The first characteristic collected was unemployment data at the NUTS 3 level. For each municipality, the number of unemployed people as well as the number of unemployed foreigners was collected.

Next, data related to the European elections in 2019 were collected. In each municipality, it is known how many people voted in total, how many people voted for the six largest parties, and how many votes the remaining parties received combined. Data was also collected in relation to people seeking protection, welfare recipients and in relation to asylum seeker benefits. It is known how many people sought protection in Germany, how many received social welfare and how many received asylum seeker benefits.

Finally, trade tax, income tax, and payroll tax data were collected for each municipality.

5.3 Shapefiles

In addition to numeric variables, the dataset also contains a geographic variable containing the geographic boundaries of a given municipality or city/district.

5.3.1 Shapefiles for Norway

The data for the Norwegian shapefiles comes from Geonorge [@Geo21] and is downloaded from a Github repository, as the data there was in a cleaner state [@Smi20]. In addition to the geographic shape, the dataset also includes a variable that contains the ID of each municipality.

5.3.2 Shapefiles for Germany

The data for the German shapefiles comes from Esri Germany [@Deu20].

5.4 OpenStreetMap Data

OpenStreetMap (OSM) is a free project that collects, structures and stores freely usable geodata in a database for use by anyone (Open Data). This data is available under a free license, the Open Database License. The core of the project is therefore an openly accessible database of all contributed geoinformation [@Ope17].

In R, the OpenStreetMap API can be queried using the R package osmdata [Pad+17]. To download all locations of a given type in a given region, a shape or bounding box must be specified along with a key and optionally a value. These key-value pairs are used to specify the type of location, for example, the "amenity" key is used for all facilities used by visitors and residents. If you use the "biergarten" value together with the "amenity" key, the locations of all beer gardens in a given geographic region will be downloaded.

OpenStreetMap users have the option to map a location as either POINT, POLYGON, MULTIPOLYGON, LINESTRING, or MULTILINESTRING. Conventionally, the first three are used. Therefore, only sites mapped as one of these were used for this work. If a location was mapped as either POLYGON or MULTIPOLYGON, the centroid of the location was calculated.

A complete list of all key-value pairs used for this work can be found in the Appendix.

5.5 Data Wrangling

The final step before analyzing the research question at hand is to combine all of these data sources into one dataset. This section will show how this was achieved.

5.5.1 Data Wrangling for Norway

The initial step in creating the final dataset was to convert the data from a wide format, as seen in Table 5.1, to a long format. This was done using the function melt() from the R package reshape2 [Wic07]. The long version of the dataset is shown in Table 5.3.

Tab. 5.3: An excerpt from the long version of the Norwegian Covid-19 data. Does not contain all variables.

kommune_no	kommune_name	population	date	value
1507	Ålesund	66258	2020-03-26	20
5001	Trondheim	205163	2020-03-26	113
1507	Ålesund	66258	2020-03-27	20
5001	Trondheim	205163	2020-03-27	136

Next, the demographic data for Norway was loaded and processed. Since the age data contains the number of people of a certain age, the median age was calculated for each region based on how many people of each age group live in each region. The other demographic variables are left unchanged. To combine the demographic

graphic data with the Covid-19 data, the municipality IDs were extracted using the str_extract() function from the stringr [Wic19] R package using the regular expression [0-9]{4}. Next, all demographic datasets and the Covid-19 dataset were merged using the merge() function.

Using the st_intersects() function from the sf [Peb18] R package, the number of points of interest downloaded via OpenStreetMap was calculated for each municipality. Since the shapefiles contain the ID for each community, these data were then merged with the data containing the demographic and Covid-19 data.

For each numeric variable, e.g. the number of schools or the number of employees, this number was scaled.

If there were missing values in the covariates, these values were imputed using the median of the respective variable.

Finally, seven new variables were created:

- 1. expected_count, which is the expected number of cases in each municipality
- 2. sir, which is the standardised incidence ratio in each municipality
- 3. idarea 1, which is a unique ID given to each municipality
- 4. higher_education, which counts the number of universities and colleges in a given area.
- 5. sex, which gives the proportion of females living in a given area.
- 6. pop dens, i.e. the number of people per square kilometre in a given area.
- 7. urb_dens, i.e. the number of residential buildings per square kilometre in a given area.

The final dataset contains the variables shown in Table 5.4.

Tab. 5.4: The variables contained in the final dataset.

Variable Name	Explanation	Scale	
kommune_no	The municipality ID	None	
kommune_name	The municipality name	None	
population	Population in a municipality	None	
date	The date of the data used	None	
value	The number of infected people	None	
median_age	The median age	None	
unemp_tot	The proportion of		
unemp_tot	unemployed people	scaled	
unemp_immg	The proportion of	scaled	
unemp_mmg	unemployed immigrants	scarca	
workers ft	The number of	scaled	
WOIRCIS_IL	full-time workers		
workers pt	The number of	scaled	
workers_pt	part-time workers	scarca	
construction ft	The number of full-time	scaled	
construction_n	construction workers	bearea	
construction pt	The number of part-time	scaled	
construction_pt	construction workers		
immigrants_total	The proportion of immigrants	scaled	
marketplace	The number of marketplaces	scaled	
entertainment	The number of entertainment venues	scaled	
sport	The number of sports amenities	scaled	
clinic	The number of clinics	scaled	
hairdresser	The number of hairdresser	scaled	
shops	The number of shops	scaled	
place_of_worship	The number of places of worship	scaled	
retail	The number of retail stores	scaled	
nursing_home	The number of nursing homes	scaled	
restaurant	The number of restaurants	scaled	
aerodrome	The number of aerodromes	scaled	
office	The number of offices	scaled	
platform	The number of public	scaled	
pidtioiiii	transport platforms	scarcu	
kindergarten	The number of kindergartens	scaled	
schools	The number of schools	scaled	
bakeries	The number of bakeries	scaled	
residential	The number of residential buildings	None	
higher education	The number of colleges	scaled	
ingiler_education	and universities		
expected_count	The expected number of infections	None	
sir	The standardised incidence ratio	None	
idarea_1	A unique ID	None	
area	The area in km ²	None	
pop_dens	People per km ²	scaled	
-			
urb_dens	Residential buildings per km ² The proportion of females	scaled scaled	

5.5.2 Data Wrangling for Germany

The data processing procedure for Germany is identical to that for Norway. First, all demographic variables were loaded and left unchanged before being merged with the Covid-19 data prior to calculating the spatial intersections between the points of interest and the NUTS-3 areas. After merging all the data, the scaled numbers were calculated for the numeric variables. For the variables containing the number of people who voted for a particular political party, the relative percentage of votes the party received was calculated. Again, missing values were imputed using the median. Finally, the same seven new variables were created. The final dataset contains the variables shown in Table 5.5.

Tab. 5.5: The variables contained in the final dataset.

Variable Name	Explanation	Scale
municipality_id	The municipality ID	None
municipality	The municipality name	None
population	Population in a municipality	None
date	The date of the data used	None
value	The number of infected people	None
trade_tax	The trade tax in Euros	scaled
income_tax	The income tax in Euros	scaled
income_total	The income and payroll tax in Euros	scaled
asyl_benefits	The number of people receiving asylum seeker benefits	scaled
welfare_recipients	The number of welfare recipients	scaled
unemployed_total	The number of unemployed people	scaled
unemployed foreigners	The number of unemployed foreigners	scaled
protection seekers	The number of protection seekers	scaled
Union	Percentage of vote for Union	scaled
SPD	Percentage of vote for SPD	scaled
Gruene	Percentage of vote for Gruene	scaled
FDP	Percentage of vote for FDP	scaled
die linke	Percentage of vote for die Linke	scaled
afd	Percentage of vote voted for AfD	scaled
marketplace	The number of marketplaces	scaled
entertainment	The number of entertainment venues	scaled
sport	The number of sports amenities	scaled
clinic	The number of clinics	scaled
hairdresser	The number of hairdresser	scaled
shops	The number of shops	scaled
place_of_worship	The number of places of worship	scaled
retail	The number of retail stores	scaled
nursing_home	The number of nursing homes	scaled
restaurant	The number of restaurants	scaled
aerodrome	The number of aerodromes	scaled
office	The number of offices	scaled
	The number of public	
platform	transport platforms	scaled
kindergarten	The number of kindergartens	scaled
schools	The number of schools	scaled
bakeries	The number of bakeries	scaled
residential	The number of residential buildings	None
higher_education	The number of colleges and universities	scaled
expected count	The expected number of infections	None
sir	The standardised incidence ratio	None
idarea_1	A unique ID	None
area	The area in km ²	None
pop dens	People per km ²	scaled
urb dens	Residential buildings per km ²	scaled
	The proportion of females	Jearca

Data Analysis

In this chapter the models calculated for each country are reviewed. First, a look at the standardised incidence rate for each country is taken, before spatial models, spatio-temporal models and finally predictive models are discussed.

6.1 Standardised Incidence Ratio

This section takes a brief look at the standardised incidence ratio for the countries of interest.

6.1.1 Standardised Incidence Ratio for Germany

When looking at the standardised incidence ratio for Germany, it is noticeable that the actual number of infections in the eastern parts of Germany, especially in Saxony, is considerably higher than the expected number of infections. Furthermore, parts of Bavaria have an increased standardised incidence ratio compared to the rest of Germany, excluding Saxony. This could be due to the fact that the regions share a border with the Czech Republic, a country that is substantially more affected by Covid-19 than Germany. The northern parts of Germany show the lowest SIR which is possibly due to the fact that this region is sparsely populated. For more information, see Figure 6.1.

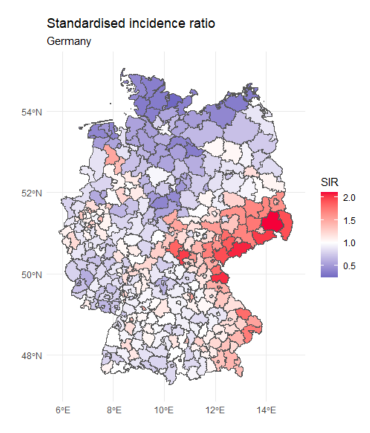


Fig. 6.1: The standardised incidence ratio for Germany based on the data of the 24th of March 2021

6.1.2 Standardised Incidence Ratio for Norway

Looking at the standardised incidence rate for Norway, a standardised incidence rate of less than 1 can be seen for most municipalities north of Trondheim. In the southern parts of Norway there are several municipalities with a rate above 1, for example the standardised incidence rate around the capital Oslo is around 2. However, the two small municipalities, Hyllestad and Ulvik, have the highest standardised incidence rate in Norway. In Hyllestad, 95 of 1328 people have been infected with Covid-19 so far, while in Ulvik, 134 of 1080 people have been infected so far.

The SIR in Hyllestad is around 4.5, following an outbreak in a shipyard in autumn 2020 [Kor20], while Ulvik has a ratio of around 8, following an outbreak of the UK variant of Covid-19. According to the head of the municipality, Hans Petter Thorbjørnsen, the infections are thought to have spread through children [NTB21]. See Figure 6.2 for more information.

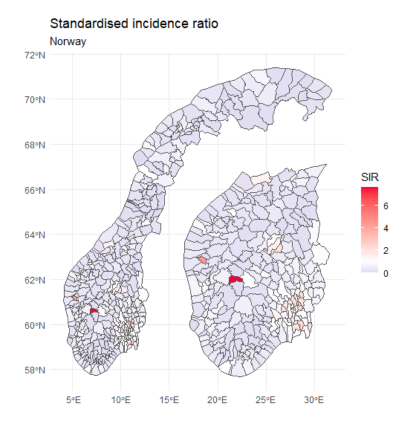


Fig. 6.2: The standardised incidence ratio for Norway based on the data of the 24th of March 2021

Because the high numbers from two small municipalities complicate the interpretation of Figure 6.2, Figure 6.3 shows the SIR on a log10 scale. It is now clearer that the standardized incidence ratio is below 1 in most parts of Norway, but that there is a higher risk in the region around Oslo.

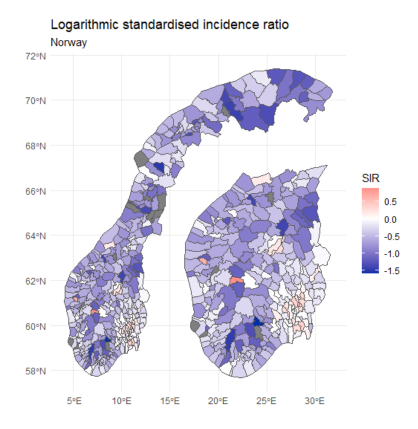


Fig. 6.3: The log10 standardised incidence ratio for Norway based on the data of the 24th of March 2021

6.2 Spatial Models

After looking at the standardised incidence rates for the countries of interest, the next step is to take a closer look at the current figures for the respective countries. Spatial models are used to try to extract the factors that cause some populations to be at higher risk than other populations. Three different types of models are used for each country:

- 1. The Besarg-Yollie-Mollie Model
- 2. Besags Proper Spatial Model
- 3. The Leroux-Model

All of these models were computed using the INLA [BGR15] R package. To specify each type of model, the code shown in Listing 7.1 can be used. Three measures are used to compare the models, the DIC, the WAIC and the CPO. For all countries, the models were computed with

- 1. only the demographic variables as covariates
- 2. only the infrastructural variables as covariates
- 3. both, demographic and infrastructural variables, as covariates
 - 3.1 Without variable selection
 - 3.2 With variable selection

For each model type, different values for the penalised prior were tried. As this resulted in a large number of models, only the model with the best performance for each model class is examined in more detail.

The models were compared using the mean absolute error. For this, 20% of the observations were removed from the training and used for testing instead. The predicted number of infections for these municipalities was then compared to the actual numbers.

Finally, due to the amount of covariates, forwards and backwards stepwise variable selection was performed with the intention of obtaining a model that fits the data well and at the same time is relatively easy to interpret. This can be done with the R package INLAutils [Red+17], as shown in Listing 7.3. Backwards as well forwards variable selection was performed.

A list of all calculated models along with their performance measures is provided in the appendix.

6.2.1 Model Selection

Before the models are computed, however, the distribution that fits the number of cases must first be found. For this, the function descdist() from the fitdistrplus R package is used. The Cullen and Frey graph can be used to give an initial idea of which distributions fit the data, in this case the number of infections, reasonably well depending on the kurtosis and the square of the skewness.

The plots for Germany and Norway can be seen in Figure 6.4 and Figure 6.5.

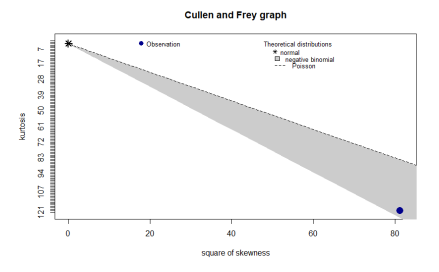


Fig. 6.4: The Cullen and Frey graph for Germany

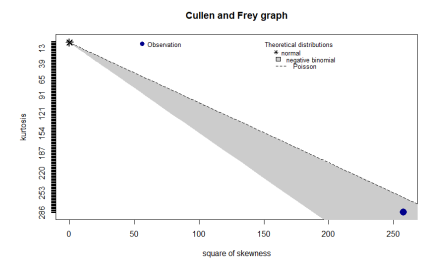


Fig. 6.5: The Cullen and Frey graph for Norway

Next, a negative binomial distribution, a normal distribution, and a Poisson distribution are fitted to the data using the maximum likelihood method. The negative binomial fits for both countries can be seen in Figure 6.6 and Figure 6.7. The fits for the normal and Poisson distribution for both countries, are shown in the Appendix in Figure 7.1, Figure 7.2, Figure 7.3 and Figure 7.4.

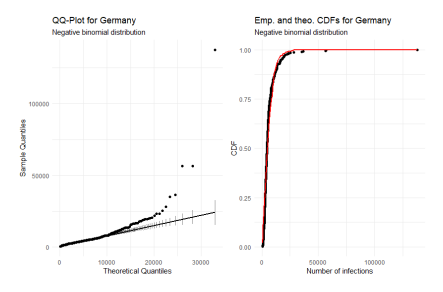


Fig. 6.6: A negative binomial fit to the number of cases in German municipalities

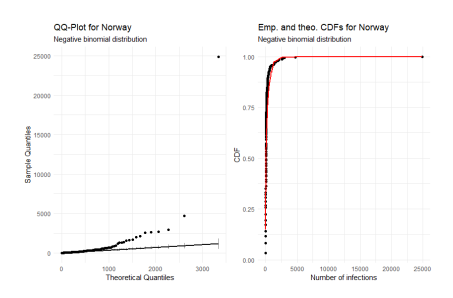


Fig. 6.7: A negative binomial fit to the number of cases in Norwegian municipalities

Lastly, the AIC was calculated for fitting a normal distribution to the data, a Poisson distribution to the data and a negative binomial distribution to the data. The values can be seen in Table 6.1. Afterwards, the negative binomial distribution was chosen

as the distribution of the target variable in both cases.

Tab. 6.1: The AIC for different distributions for Germany and Norway

Country	Distribution	AIC
Germany	Normal	8360
Germany	Poisson	2148100
Germany	Negative Binomial	7731
Norway	Normal	6166
Norway	Poisson	366181
Norway	Negative Binomial	4086

The poor fit for the Poisson distribution can be explained by looking at the range of the number of confirmed cases in a given municipality. For Germany, this number ranges from 508 to 137634 (as of March 18, 2021), while for Norway, the number ranges from 0 to 24905 (as of March 20, 2021). This results in a mean and standard deviation for Germany of 6617 and 9014, respectively. For Norway, the values for these metrics are 236 and 1389.

6.2.2 Spatial Models for Germany

First, a look is taken at the spatial models calculated for Germany. These models are based on data from 24 March 2021, when 2.695.037 people in Germany were confirmed infected with Covid-19. The five municipalities with the most infections are shown in Table 6.2.

Tab. 6.2: The municipalities with the most infections as of March 24th 2021.

Municipality	Population	Number of infections
SK Berlin	3644826	141639
SK Hamburg	1841179	58661
SK Munich	1471508	57690
SK Cologne	1085664	37592
Region Hannover	1157624	36241

6.2.2.1 Demographic Models

The model with the best performance based on the demographic variables was a Leroux model computed using the formula in Listing 7.6. The variables used for this

model were the percentages of the vote for the six biggest German political parties. The performance measures of this model and the best-performing BYM2 and Besag proper models are shown in Table 6.3.

Tab. 6.3: The performance measures for the best performing demographic model of each type.

Model	DIC	WAIC	СРО	MAE
Besag	4916	4919	-2719	193968
BYM2	4682	4670	-2726	193922
Leroux	4728	4753	-3005	191965

The summary of the fixed effects is shown in Table 6.4. To compute the posterior mean of the coefficients, the code shown in Listing 7.4 can be used.

To obtain a credibility interval of the fixed effects on the transformed scale, the code in Listing 7.5 can be used.

Tab. 6.4: The fixed effects for the Leroux model. Values are rounded.

Variable	Mean	exp(mean _p)	exp(q0025p)	exp(q0975 _p)
(Intercept)	-0.07468	0.9282	0.8976	0.9594
afd	0.1230	1.134	0.9820	1.302
FDP	-0.01211	0.9881	0.9515	1.026
Gruene	-0.1363	0.8750	0.7555	1.008
SPD	-0.1702	0.8443	0.7749	0.9179
Union	-0.1714	0.8453	0.7175	0.9889
die_linke	-0.2159	0.8067	0.7353	0.8830

The posterior mean of the exponentiated intercept implies a -7.2% risk rate across Germany with a credibility interval ranging from -10.2% to 4.1%.

This summary shows that especially people who come from regions where the AfD gets a higher share of the vote have a higher risk of contracting Covid-19. In Figure 6.1, regions in eastern Germany often had a higher SIR. These are also the regions where the AfD tends to perform better in elections. The connection between the two can be attributed to the fact that the AfD is a right-wing party that openly criticises the measures taken to prevent the spread of the coronavirus. As a result, a large proportion of people who vote for the AfD do not take the measures seriously either and refuse to keep a safe distance or wear a mask in public spaces.

A one standard deviation increase for the AfD leads to a risk increase of about 13.4%. The only other party with a posterior mean close to 1 is the liberal FDP, which is

known for its capitalist views. However, it is still below 1. An increase of 1 standard deviation leads to a risk reduction of 1.8%.

As for the rest of the parties, they are either the leading party in Germany (Union) or left-wing oriented (SPD, the Greens and the Left). People who vote for left-wing parties may respect the Corona rules more and show more consideration for other people. As this was the overall best-performing spatial model for Germany, a look is taken at the posterior means $\zeta = \exp(\xi)$ of the area-specific risks. Since the uncertainty associated with them can also be mapped, the posterior probability $\mathbb{P}(\zeta_i > 1|y)$ is visualised as well. In Figure 6.8, it can be seen that the relative risk is above 1 for large parts of Germany, except for most of northern Germany and some parts in the south of the country. The posterior probability is highest for parts of North Rhine-Westphalia, home to Germany's most populous metropolitan area, the Ruhr area, and for Saxony and Thuringia, both political strongholds of the AfD. The higher numbers in Bavaria can be explained by the large common borders with Austria and the Czech Republic, two countries that were more affected by the pandemic. Many people cross these borders because of tourism and many guest workers come from the Czech Republic to work on agricultural land in Bavaria, which resulted in several Covid-19 outbreak events.

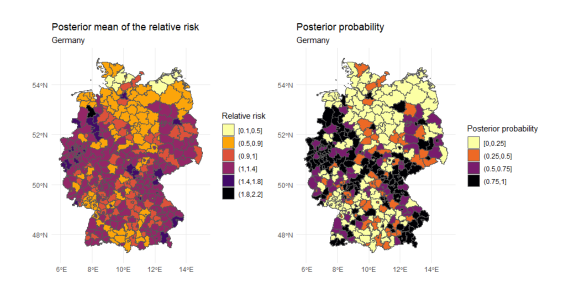


Fig. 6.8: Posterior mean of the area-specific risk and the posterior probability.

6.2.2.2 Infrastructure Models

When it comes to the models based on the infrastructural variables, a BYM2 model performed the best. It was computed using formula in Listing 7.7.

The performance measures of this model and the best-performing Besag proper and Leroux model are shown in Table 6.5

Tab. 6.5: The performance measures for the best performing infrastructure model of each type.

Model	DIC	WAIC	СРО	MAE
Besag	4749	4745	-2799	194009
BYM2	4635	4620	-2758	193472
Leroux	4785	4823	-3444	1947560

The values of the covariates are shown in Table 6.6.

Tab. 6.6: The fixed effects for the BYM2 model. Values are rounded.

Variable	Mean	exp(mean _p)	exp(q0025p)	exp(q0975 _p)
(Intercept)	-0.07373	0.9289	0.9159	0.9421
shops	0.3860	1.479	1.197	1.808
hairdresser	0.1123	1.122	0.9655	1.296
marketplace	0.04187	1.043	0.9846	1.104
schools	0.02157	1.023	0.9199	1.135
nursing_home	0.01399	1.014	0.9879	1.042
aerodrome	0.01273	1.013	0.9648	1.063
bakeries	0.008426	1.011	0.8684	1.171
sport	0.007894	1.009	0.9374	1.084
office	0.005687	1.006	0.9533	1.061
retail	-0.0028888	0.9977	0.9317	1.067
place_of_worship	-0.005710	0.9945	0.9562	1.034
higher_education	-0.03179	0.9689	0.9324	1.006
clinic	-0.03258	0.9683	0.9154	1.023
entertainment	-0.03454	0.9676	0.8656	1.078
platform	-0.05867	0.9434	0.8913	0.9976
restaurant	-0.1990	0.8213	0.7206	0.9316
kindergarten	-0.2268	0.7993	0.6871	0.9243

Some findings from these results are that shops, defined as supermarkets, convenience stores and drugstores, hairdressers, marketplaces, schools, nursing homes and bakeries, among others, increase the risk of infection. The two highest are shops and hairdressers, where a 1 standard deviation increase leads to a 47.9% increase in

risk in the case of shops and a 12.2% increase in risk for hairdressers. It should not be surprising that shops in particular increase the risk of infection, as people tend to congregate there and people living in areas where a supermarket is nearby may prefer to go shopping more often to buy fresh produce rather than just once a week when a supermarket is further away.

In Germany, there was a big discussion about whether hairdressers should be allowed to stay open during lockdowns, as they were forced to close and could only work when there was no lockdown. Seeing how much hairdressers increase the risk of infection, it might have been a wise decision not to allow them to continue working during a closure.

Finally, the number of schools in a given community also increases the risk of infection. An increase of 1 standard deviation increases the risk by about 2.3%.

There are also effects that have a posterior mean of less than 1, suggesting that they reduce the risk of infection. However, this is not necessarily the case, as it is probably safer for a person to stay at home than to go to a restaurant, for example. It is interesting to note, however, that the two lowest posterior means were observed for restaurants and kindergartens. In the case of restaurants, this could be due to the fact that during most of the period restaurants were either only allowed to serve take-away food or had very strict hygiene rules. The low value could indicate that these measures worked as intended.

For the kindergartens, the reason could be that they were also closed most of the time and the children therefore stayed at home with their families and did not interact with other children.

6.2.2.3 Combined Models

Finally, models are considered that include both the infrastructural and demographic covariates. Due to the amount of variables, all models run are based on variable selection.

The best performing model was again a BYM2 model, run with the formula shown in Listing 7.8.

The performance measures of the computed models are shown in Table 6.7.

Tab. 6.7: The performance measures for the best performing demographic + infrastructure model of each type.

Model	DIC	WAIC	СРО	MAE
Besag	4755	4721	-2756	198553
BYM2	4753	4774	-2776	198461
Leroux	4847	4818	-2684	204102

The fixed effects are shown in Table 6.8.

Tab. 6.8: The fixed effects for the BYM2 model. Values are rounded.

Variable	Mean	exp(mean _p)	exp(q0025 _p)	exp(q0975 _p)
(Intercept)	-0.0716	0.9309	0.9168	0.9450
afd	0.2528	1.288	1.222	1.357
pop_dens	0.09614	1.101	1.055	1.148
welfare_recipients	0.07515	1.079	1.004	1.157
sport	0.05575	1.058	0.9945	1.124
schools	0.03926	1.041	0.9485	1.140
nursing_home	0.008492	1.009	0.9856	1.032
office	-0.007482	0.9928	0.9510	1.036
log(trade_tax)	-0.007660	0.9924	0.9695	1.016
place_of_worship	-0.01141	0.9888	0.9557	1.023
platform	-0.02052	0.9800	0.9322	1.029
kindergarten	-0.02174	0.9799	0.8812	1.087
FDP	-0.03154	0.9691	0.9340	1.005
SPD	-0.03197	0.9688	0.9288	1.010
entertainment	-0.03963	0.9619	0.8891	1.039
bakeries	-0.06149	0.9419	0.8412	1.051
_die_linke	-0.08336	0.9204	0.8697	0.9734

According to this model, the three biggest driving factors are the percentage of votes the AfD receives, the population density and the number of welfare recipients in a municipality. The influence of the AfD has already been discussed in section 6.2.2.1, only in this case a 1 standard deviation increase leads to a risk increase of 28.8%. Regarding population density, it can be said that if it increases by 1 standard deviation, the risk of infection increases by 10.1%. The more people are concentrated per square kilometre, the easier it is for a virus to spread because the distances between people become smaller. Other factors that positively influence the risk of infection are the number of sports facilities, the number of schools and the number of nursing homes, although the risk increase for the latter is only just above 0. Sports facilities are again a place where many people come together, and here, depending

on the type of sport, there can also be more contact, which in turn favours the spread of a virus. The last effect worth mentioning is the logarithmic trade tax. If a region receives more revenue through trade tax, the risk of infection in that region decreases. Therefore, people in communities where there is less money are at higher risk.

6.2.3 Spatial Models for Norway

Next, the same types of models are evaluated for Norway. These models are based on data from 24 March 2021, when 87537 people in Norway were confirmed infected with Covid-19. The five municipalities with the most infections are shown in Table 6.9.

Tab. 6.9: The municipalities with the most infections as of March 24th 2021.

Population	Number of infections
693494	26151
283929	4738
101386	3149
127731	2834
85983	2779
	693494 283929 101386 127731

6.2.3.1 Demographic Models

As with Germany, the best demographic model was again a Leroux model calculated using the formula shown in Listing 7.9.

The performance measures for all three models are shown in Table 6.10. It is noticeable that the mean error in Norway is significantly smaller, suggesting that the models for Germany have greatly overestimated or underestimated the number of infections.

Tab. 6.10: The performance measures for the best performing demographic model of each type.

Model	DIC	WAIC	СРО	MAE
Besag	2396	2403	-4369	10149
BYM2	2224	2202	-6793	10002
Leroux	2225	2192	-6810	8460

All other performance measures for these models are also lower than for the models computed for Germany.

The effects of the covariates are shown in Table 6.11.

Tab. 6.11: The fixed effects for the Leroux model. Values are rounded.

Variable	Mean	exp(mean _p)	exp(q0025p)	exp(q0975 _p)
(Intercept)	-1.040	0.3540	0.3173	0.3929
unemp_total	0.3371	1.406	1.192	1.645
unemp_immg	0.03737	1.042	0.8788	1.226

The posterior mean of the section implies a risk rate of -64.6%, which does not say much except for the fact that there is a very low risk of getting Covid-19 in Norway. According to this model, the risk depends on only two factors, namely total unemployment and immigrant unemployment. A 1 standard deviation increase in total unemployment leads to a 40.6% increase in risk and a 1 standard deviation increase in immigrant unemployment leads to a 4.2% increase in risk. A possible explanation for this could be that the unemployed spend less time at home and instead move around more and thus have more contact with people, but this should be taken with a grain of salt.

6.2.3.2 Infrastructure Models

For the models based on the infrastructural variables, the best predictive performance was observed for a Besag model. The formula is shown in Listing 7.10. The performance measure for the models are shown in Table 6.12.

Tab. 6.12: The performance measures for the best performing infrastructure model of each type.

Model	DIC	WAIC	СРО	MAE
Besag	2326	2335	-8405	11694
BYM2	2218	2187	-6532	11866
Leroux	2230	2167	-7128	12943

The effect of the covariates are shown in Table 6.13.

Tab. 6.13: The fixed effects for the Besag model. Values are rounded.

Variable	Mean	exp(mean _p)	exp(q0025 _p)	exp(q0975 _p)
(Intercept)	-1.061	0.3518	0.2437	0.4965
schools	0.3253	1.407	0.9864	1.966
kindergarten	0.2418	1.285	0.9825	1.663
pop_dens	0.1531	1.171	0.9733	1.406
place_of_worship	0.03138	1.041	0.8053	1.335
restaurant	0.01374	1.044	0.6384	1.626
nursing_home	0.001030	1.002	0.9258	1.085
office	-0.07424	0.9263	0.7991	1.084
shops	-0.5186	0.6451	0.2793	1.303

According to this model, schools, kindergartens, places of worship and nursing homes are infrastructural features that increase the risk of infection. In addition, higher population density also increases the risk of infection. Schools and kindergartens are the two biggest driving factors, as a 1 standard deviation increase in schools leads to a 40.7% increase in risk and a 1 standard deviation increase in kindergartens leads to a 28.5% increase in risk. The risk of schools and kindergartens is reflected in the Corona outbreak that occurred in Tromsø in early March 2021, where several infections have been linked to school classes and kindergartens [Kom20]. A 1 standard deviation increase in population density leads to a 17.1% increase in risk, and for places of worship and nursing homes, a 1 standard deviation increase leads to a 4.1% and 0.2% increase in risk, respectively. Finally, for restaurants, a 1 standard deviation increase leads to a 4.4% increase in risk. Why restaurants do not have a large impact on the risk of infection has already been discussed in Section 6.2.2.2

For offices and shops, on the other hand, there is no increased risk of infection. For offices, one reason why they do not increase the risk could be that more and more people work from home and therefore there is less contact between people at office locations and thus no increased risk of infection. Finally, shops do not increase the risk of infection here, whereas in Germany they were one of the driving factors, as can be seen in Table 6.6. Why this is so is difficult to say. Because of the lower infection numbers, there is of course less risk when people congregate in supermarkets, which may be an explanation, even if these are not driving factors of the pandemic in Norway. It could also mean that Norwegians are more likely to be careful to keep enough distance from each other.

Looking at the posterior mean relative risk, it can be seen that the region in and around the capital Oslo is most at risk, as the posterior mean of the relative risk can be as high as 6, with a posterior probability between 0.75 and 1. In addition, some

municipalities around Bergen and Ulvik are also more at risk, with a relative risk of up to 3.4 and a posterior probability between 0.75 and 1.

The risk seems to be lowest in the parts north of Trondheim and south of Bodø, where the relative risk is mostly below 1. For many municipalities in Troms og Finnmark, the northernmost part of Norway, the posterior mean is also below 1, but for a few regions it is above 1 with posterior probabilities between 0.5 and 0.75 and 0.75-1.

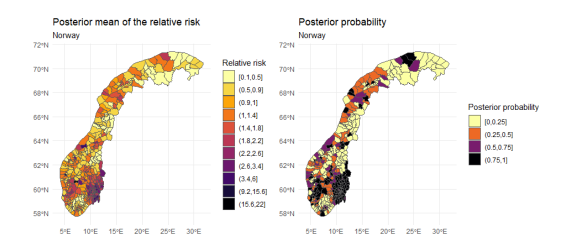
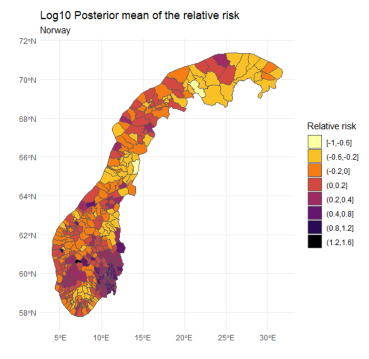


Fig. 6.9: Posterior mean of the area-specific risk and the posterior probability.

For better interpretability, the logarithmic posterior mean is shown in Figure 6.10.



 $\textbf{Fig. 6.10:} \ \ Logarithmic \ posterior \ mean \ of \ the \ area-specific \ risk.$

6.2.3.3 Combined Models

For the models containing variables related to the demographic of Norway and the infrastructure, again a Besag model performed the best. Its formula can be seen in Listing 7.11.

The performance measures for the models are shown in Table 6.14

Tab. 6.14: The performance measures for the best performing demographic + infrastructure model of each type.

Model	DIC	WAIC	CPO	MAE
Besag	2352	2355	-6939	13036
BYM2	2215	2184	-6406	13932
Leroux	2229	2193	-7677	14841

The fixed effects are shown in Table 6.15.

Tab. 6.15: The fixed effects for the Besag model. Values are rounded.

Variable	Mean	exp(mean _p)	exp(q0025p)	exp(q0975 _p)
(Intercept)	-1.068	0.3484	0.2478	0.4736
construction_pt_work	0.1873	1.234	0.7946	1.834
clinic	0.1855	1.266	0.6488	2.241
pop_dens	0.1717	1.194	0.9703	1.455
unemp_tot	0.1559	1.170	1.057	1.292
schools	0.0978	1.117	0.8074	1.507
sex	0.04402	1.046	0.9426	1.158
restaurant	0.03072	1.079	0.5717	1.855
higher_education	-0.05942	0.9443	0.8295	1.069
median_age	-0.1147	0.8929	0.8015	0.9910
workers_ft_work	-0.5200	0.6403	0.2784	1.259

Some of these factors have already been discussed in the previous sections, so they will not be discussed in detail again.

The number of clinics, the number of part-time construction workers, the population density, total unemployment and the number of schools are the five biggest drivers of relative risk, as a 1 standard deviation increase in any of these factors leads to an increase in relative risk of 26.6%, 23.4%, 19.4%, 17.0% and 11.7% respectively. That clinics increase the relative risk could be due to the fact that when there are more clinics in a community, more people who have symptoms go to a clinic and get tested, which naturally leads to more detected cases. In Norway, there has been some discussion about whether the number of foreign construction workers coming from countries with higher infection rates, such as Poland, could contribute to the spread of Covid-19. Since most of these are classified as part-time workers, this model suggests that this may be the case.

The other two factors that increase the risk are the proportion of women in a community and the number of restaurants. For the proportion of women, a 1 standard deviation increase leads to a 4.6% increase in risk and for restaurants, a 1 standard deviation increase leads to a 0.8% increase in risk.

The median age lowers the risk of infection, as an increase of 1 standard deviation lowers the relative risk by 9.8%. This in turn means that people living in communities with a lower median age have a higher risk than people living in "older" communities. This could be due to the fact that younger people tend to be more mobile and have more contact with other people as a result of this increased movement, which in turn helps the virus spread.

- 6.3 Spatio-Temporal Models
- 6.3.1 Spatio-Temporal Models for Germany
- 6.3.2 Spatio-Temporal Models for Norway
 - 6.4 Predictive Models

- 6.4.1 Predictive Models for Germany
- 6.4.2 Predictive Models for Norway

Appendix

7.1 Probability Distributions

7.1.1 The Normal Distribution

The normal distribution is an important type of continuous probability distribution in stochastics. The special significance of the normal distribution is based, among other things, on the central limit theorem, according to which distributions that result from the additive combination of a large number of independent influences are approximately normally distributed under weak conditions.

The density is given by

$$f(x|\mu,\sigma) = \frac{1}{\sigma\sqrt{2\pi}} \exp\left(-\frac{1}{2}\left(\frac{x-\mu}{\sigma}\right)^2\right). \tag{7.1}$$

The first two moments of the distribution are given by

$$\mathbb{E}\left[X\right] = \mu \tag{7.2}$$

$$Var[X] = \sigma^2. (7.3)$$

The graph of this density function has a "bell-shaped form" and is symmetrical with μ as the centre of symmetry [Fah+16, pp. 83–85].

7.1.2 The Poisson Distribution

The Poisson distribution is a discrete probability distribution that can be used to model the number of events that occur independently of each other at a constant mean rate in a fixed time interval or spatial area.

The density is given by

$$f(k) = \mathbb{P}(X = k) = \begin{cases} \frac{\lambda^k}{k!} \exp(-\lambda) & \text{for } x \in \{0, 1, ...\} \\ 0 & \text{else} \end{cases}$$
 (7.4)

with λ representing the expected value of X.

The first two moments of the distribution are given by

$$\mathbb{E}\left[X\right] = \lambda \tag{7.5}$$

$$Var[X] = \lambda. (7.6)$$

For $\lambda \ge 10$ the distribution becomes approximately symmetrical and can thus be approximated by a normal distribution [Fah+16, p. 243].

7.1.3 The Negative Binomial Distribution

The negative binomial distribution is a univariate probability distribution that belongs to the discrete probability distributions. It models the number of trials required to achieve a given number of successes in a Bernoulli process.

The density is given by

$$f(k,r,p) = \mathbb{P}(X=k) = {k+r-1 \choose r-1} (1-p)^k p^r,$$
 (7.7)

with r the number of successes, k the number of failures, and p the probability of success.

The first two moments of the distribution are given by

$$\mathbb{E}\left[X\right] = \frac{pr}{1-p} \tag{7.8}$$

$$Var[X] = \frac{pr}{(1-p)^2}. (7.9)$$

For large values of r, the negative binomial distribution can be approximated by a normal distribution [Hal41].

7.2 Distribution Fits

7.2.1 Distribution Fits for Germany

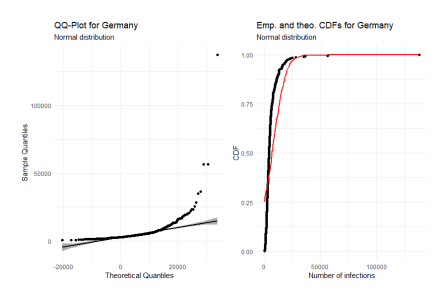


Fig. 7.1: A normal fit to the number of cases in German municipalities

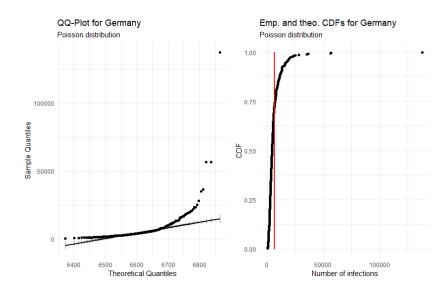


Fig. 7.2: A Poisson fit to the number of cases in German municipalities

7.2.2 Distribution Fits for Norway

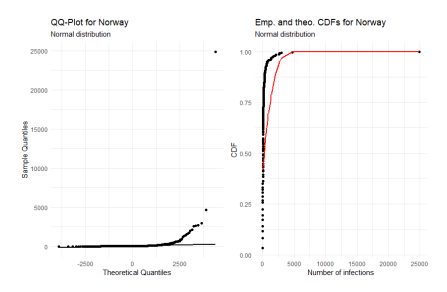


Fig. 7.3: A normal fit to the number of cases in Norwegian municipalities

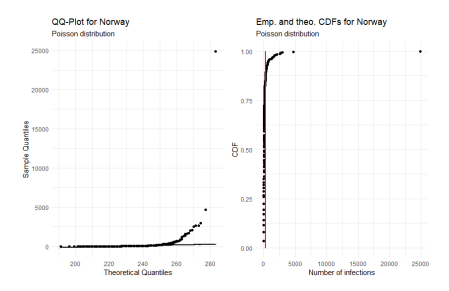


Fig. 7.4: A Poisson fit to the number of cases in Norwegian municipalities

7.3 Code examples

7.3.1 Specifying the Different Types of Models

```
# set the seed
    set.seed (420)
 3
    # draw a sample
    test <- sample(
 5
         seq_len(nrow(newest_numbers)),
 6
         size = floor(0.2 * nrow(newest_numbers))
 7
    # get the number of infections for the test data
    test_value <- newest_numbers$value[test]</pre>
10
    # set the number of infections to NA in the train data
11
    newest_numbers$value[test] <- NA</pre>
12
    # define the link function
13
    link <- rep(NA, nrow(newest_numbers))</pre>
    link[which(is.na(newest_numbers$value))] <- 1</pre>
15
    # define the penalised prior
16
    prior_1 <- list(</pre>
17
     prec = list(
18
        prior = "pc.prec",
19
         param = c(1, 0.01)
20
21
    )
22
    # create the neighbordhood matrix
    nb <- poly2nb(newest_numbers)</pre>
    # save the matrix
25
    nb2INLA("maps/map.adj", nb)
26
    g <- inla.read.graph(filename = "maps/map.adj")</pre>
27
    # define the C matrix for the Leroux model
28
    Q <- Diagonal(x = sapply(nb, length))
    for (i in 2:nrow(newest_numbers)) {
30
      Q[i - 1, i] <- -1
31
      Q[i, i-1] < --1
32
33
34
    C <- Diagonal(x = 1, n = nrow(newest_numbers)) - Q</pre>
35
    # define the formula for besags proper model
    formula_besag <- value ~
37
      pop_dens + urb_dens + sex +
38
      f(idarea_1, model = "besagproper", graph = g, hyper = prior_1)
     # define the formula for bym2 model
40
    formula_bym2 <- value <-</pre>
41
      pop_dens + urb_dens + sex +
42
      f(
43
         idarea_1, model = "bym2", graph = g,
```

```
44
        scale.model = TRUE, hyper = prior_1
45
46
    # define the formula for leroux model
47
    formula_leroux <- value ~</pre>
48
      pop_dens + urb_dens + sex +
49
      f(idarea_1, model = "generic1", Cmatrix = C, hyper = prior_1)
50
    # compute the models
51
    res_besag <- inla(
52
      formula_besag,
53
      family = "nbinomial",
54
      data = newest_numbers,
55
      E = expected_count,
56
      control.predictor = list(
57
        compute = TRUE,
58
        link = link
59
      ),
60
      Ntrials = newest_numbers$population,
      control.compute = list(dic = TRUE, waic = TRUE, cpo = TRUE)
61
62
    )
63
    res_bym2 <- inla(
64
      formula_bym2,
65
      family = "nbinomial",
66
      data = newest_numbers,
67
      E = expected_count,
      control.predictor = list(
68
        compute = TRUE,
70
        link = link
71
      ),
72
      Ntrials = newest_numbers$population,
      control.compute = list(dic = TRUE, waic = TRUE, cpo = TRUE)
73
74
    )
75
    res_leroux <- inla(
76
      formula_leroux,
77
      family = "nbinomial",
78
      data = newest_numbers,
79
      E = expected_count,
80
      control.predictor = list(
81
        compute = TRUE,
82
        link = link
83
      ),
84
      Ntrials = newest_numbers$population,
      control.compute = list(dic = TRUE, waic = TRUE, cpo = TRUE)
86
```

Listing 7.1: Specifying different models in INLA.

7.3.2 Making Predictions for the Test Data

```
# create a vector to save the predictions to
    predicted <- c()</pre>
   # now make the predictions
   for(i in seq_len(nrow(newest_numbers))) {
5
     predicted[i] <- inla.emarginal(</pre>
        function(x) x * newest_numbers$population[i],
7
        res_bym2$marginals.fitted.values[[i]]
8
      )
9
    }
10
    \# calculate the MAE for the test data
11  mean(abs(predicted[test] - test))
```

Listing 7.2: The code for making predictions in INLA.

7.3.3 Variable Selection using INLA

```
# define the stack
    stack_all <- inla.stack(</pre>
      data = list(value = newest_numbers$value),
      A = list(1),
      effects = list(
5
6
        data.frame(
7
          Intercept = 1,
          newest_numbers[, c(2:15, 20:35, 43:46)]
9
10
      )
    )
11
12
    # run backwards varaible selection
    result_all_backwards <- INLAstep(
      fam1 = "nbinomial",
14
15
     newest_numbers,
     in_stack = stack_all,
      invariant = "Intercept",
17
18
      direction = "backwards",
      include = c(2:15, 20:35, 43:46),
19
20
      y = "value",
21
     y2 = "value",
22
      powerl = 1,
23
      inter = 1,
24
      thresh = 2,
25
      num.threads = 7
26
27
    # run forwards variable selection
28 result_all_forwards <- INLAstep(
```

```
29
      fam1 = "nbinomial",
30
      newest_numbers,
31
      in_stack = stack_all,
      invariant = "Intercept",
33
      direction = "forwards",
34
      include = c(2:15, 20:35, 43:46),
      y = "value",
36
      y2 = "value",
37
      powerl = 1,
      inter = 1,
39
      thresh = 2,
      num.threads = 7
40
41
```

Listing 7.3: The code for variable selection in INLA.

7.3.4 Calculating the Posterior Mean

```
inla.emarginal(exp, model_leroux$marginals.fixed$trade_tax)

# calculate the increase in risk for an increase

# by 2 in the trade tax

inla.emarginal(
    exp,

model_leroux$marginals.fixed$trade_tax

) ^ 2
```

Listing 7.4: Calculating the posterior mean of a coefficient.

7.3.5 Calculating a Credibility Interval

```
inla.qmarginal(
c(0.025,0.975),
inla.tmarginal(
exp,
model_leroux$marginals.fixed$trade_tax
)
)
```

Listing 7.5: Extracting the credibility interval for a coefficient

7.3.6 Best Spatial Models For Germany

7.3.6.1 Best Spatial Model using Demographic Variables

```
1
   prior_2 <- list(</pre>
2
     prec = list(
3
       prior = "pc.prec",
4
       param = c(0.5 / 0.31, 0.01)
5
     )
6
   )
7
   formula <- value ~
     Union + SPD + Gruene + FDP +
9
     die_linke + afd +
   f(idarea_1, model = "generic1", Cmatrix = C, hyper = prior_2)
```

Listing 7.6: The code for the demographic model.

7.3.6.2 Best Spatial Model using Infrastructural Variables

```
1
    prior_1 <- list(</pre>
2
      prec = list(
3
        prior = "pc.prec",
4
        param = c(1, 0.01)
5
6
7
    formula <- value ~
8
      marketplace + entertainment + sport + clinic +
9
      hairdresser + shops + place_of_worship + retail + nursing_home +
10
     restaurant + aerodrome + office + platform + schools +
11
     higher_education + kindergarten + bakeries +
12
13
        idarea_1, model = "bym2", graph = G,
14
        scale.model = TRUE, hyper = prior_1
15
```

Listing 7.7: The code for the infrastructure model.

7.3.6.3 Best Spatial Model using Both Types of Variables

```
1
   prior_1 <- list(</pre>
2
     prec = list(
3
       prior = "pc.prec",
4
       param = c(1, 0.01)
5
     )
6
7
   formula <- value ~
8
     schools + afd + die_linke + pop_dens +
9
     place_of_worship + entertainment + bakeries + SPD +
   platform + sport + nursing_home + welfare_recipients +
```

```
FDP + kindergarten + log(trade_tax) + office +

f(
    idarea_1, model = "bym2", graph = g,
    scale.model = TRUE, hyper = prior_1
)
```

Listing 7.8: The code for the demographic + infrastructure model.

7.3.7 Best Spatial Models For Norway

7.3.7.1 Best Spatial Model using Demographic Variables

```
prior_1 <- list(</pre>
 1
 2
      prec = list(
 3
         prior = "pc.prec",
 4
         param = c(1, 0.01)
 5
      )
 6
 7
    formula <- value ~
8
      unemp_immg + unemp_tot
9
10
        idarea_1, model = "generic1",
11
         Cmatrix = C, hyper = prior_1
12
      )
```

Listing 7.9: The code for the demographic model.

7.3.7.2 Best Spatial Model using Infrastructural Variables

```
1
    prior_1 <- list(</pre>
2
      prec = list(
 3
        prior = "pc.prec",
 4
        param = c(1, 0.01)
5
6
 7
    formula <- value ~
8
      schools + shops + place_of_worship + office +
9
      schools + nursing_home + kindergarten + restaurant +
10
11
        idarea_1, model = "besagproper",
         graph = g, hyper = prior_1
13
```

Listing 7.10: The code for the infrastructural model.

7.3.7.3 Best Spatial Model using Both Types of Variables

```
prior_1 <- list(</pre>
2
     prec = list(
3
       prior = "pc.prec",
        param = c(0.5 / 0.31, 0.01)
4
5
6
    )
7
   formula <- value ~
8
     schools + unemp_tot + restaurant + sex +
9
     median_age + pop_dens + construction_pt_work +
10
      workers_ft_work + higher_education + clinic +
11
     f(
12
        idarea_1, model = "besagproper",
13
        graph = g, hyper = prior_2
14
   )
```

Listing 7.11: The code for the demographic + infrastructure model.

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