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Does non-stationary spatial data always require 
non-stationary random fields?

Geir-Arne Fuglstad∗1, Daniel Simpson1, Finn Lindgren2, and Håvard Rue1

1Department of Mathematical Sciences, NTNU, Norway
2Department of Mathematical Sciences, University of Bath, UK

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Abstract

A stationary spatial model is an idealization and we expect that the true 
dependence structures of physical phenomena are spatially varying, but how should 
we handle this non-stationarity in practice? We study the challenges involved in 
applying a flexible non-stationary model to a dataset of annual precipitation in 
the conterminous US, where exploratory data analysis shows strong evidence of a 
non-stationary covariance structure.

The aim of this paper is to investigate the modelling pipeline once non-stationarity 
has been detected in spatial data. We show that there is a real danger of over-fitting 
the model and that careful modelling is necessary in order to properly account 
for varying second-order structure. In fact, the example shows that sometimes non-
stationary Gaussian random fields are not necessary to model non-stationary spatial 
data.

Keywords: Annual precipitation, Penalized maximum likelihood, Non-stationary 
Spatial modelling, Stochastic partial differential equations, Gaussian random fields, 
Gaussian Markov random fields

1 Introduction

There are, in principle, two sources of non-stationarity present in any dataset: the non-
stationarity in the mean and the non-stationarity in the covariance structure. Classical 
geostatistical models based on stationary Gaussian random fields (GRFs) ignore the 
latter, but include the former through covariates that capture important structure in the 
mean. The focus of non-stationary spatial modelling is non-stationarity in the covariance 
structure. However, it is impossible to separate the non-stationarity in the mean and 
the non-stationarity in the covariance structure based on a single realization, and even 
with multiple realizations it is challenging.

∗Corresponding author, fuglstad@math.ntnu.no
The Karhunen-Loève expansion states that under certain conditions a GRF can be decomposed into an infinite linear combination of orthogonal functions, which is weighted by independent Gaussian variables with decreasing variances. For a single realization these orthogonal functions will be confounded with the covariates in the mean, and the mean structure and the covariance structure cannot be separated. This can give apparent long range dependencies and global non-stationarity if spatially varying covariates are missing in the mean. Such spurious global non-stationarity and its impact on the local estimation of non-stationarity is an important topic in the paper.

A high degree of flexibility in the covariance structure combined with weak information about the covariance structure in the data makes overfitting by interpreting ostensible patterns in the data as non-stationarity a critical issue. However, the most important point from an applied viewpoint is the computational costs of running a more complex model versus the scientific gain. Non-stationarity in the mean is computationally cheap, whereas methods for non-stationarity in the covariance structure are much more expensive. This raises two important questions: How much do we gain by including non-stationarity in the covariance structure? What sort of non-stationarity, if any, is most appropriate for the problem at hand?

The computational cost of non-stationary models usually comes from a high number of highly dependent parameters that makes it expensive to run MCMC methods or likelihood optimizations, but the challenges with non-stationary models are not only computational. Directly specifying non-stationary covariance functions is difficult and we need other ways of constructing models. Additionally, we need to choose where to put the non-stationarity. Should we have non-stationarity in the range, the anisotropy, the marginal variance, the smoothness or the nugget effect? And how do we combine it all to a valid covariance structure?

### 1.1 Non-stationarity

Most of the early literature on non-stationary methods deals with data from environmental monitoring stations where multiple realizations are available. In this situation it is possible to calculate the empirical covariances between observed locations, possibly accounting for temporal dependence, and finding the required covariances through, for example, shrinkage towards a parametric model (Loader and Switzer 1989) or kernel smoothing (Oehlert, 1993). It is also possible to deal efficiently with a single realization with the moving window approach of Haas (Haas, 1990a,b, 1995), but this method does not give valid global covariance structures.

However, the most well-known method from this time period is the deformation method of Sampson and Guttorp (1992), in which an underlying stationary process is made non-stationary by applying a spatial deformation. The original formulation has been extended to the Bayesian framework (Damian et al. 2001, 2003; Schmidt and O’Hagan 2003), to a single realization (Anderes and Stein 2008), to covariates in the covariance structure (Schmidt et al. 2011) and to higher dimensional base spaces (Bornn et al. 2012).

Another major class of non-stationary methods is based on the process convolution
method developed by Higdon (Higdon, 1998; Higdon et al., 1999). In this method a spatially varying kernel is convolved with a white noise process to create a non-stationary covariance structure. Paciorek and Schervish (2006) looked at a specific case where it is possible to find a closed form expression for a Matérn-like covariance function and Neto et al. (2014) used a kernel that depends on wind direction and strength to control the covariance structure. The process convolution methods have also been extended to dynamic multivariate processes (Calder, 2007, 2008) and spatial multivariate processes (Kleiber and Nychka, 2012).

It is possible to take a different approach to non-stationarity, where instead of modelling infinite-dimensional Gaussian processes one uses a linear combination of basis functions and models the covariance matrix of the coefficients of the basis functions (Nychka et al., 2002, 2015). One such approach is the fixed rank kriging method (Cressie and Johannesson, 2008), which uses a linear combination of a small number of basis functions and estimates the covariance matrix for the coefficients of the linear combination. This approach leads to a continuously indexed spatial process with a non-stationary covariance structure. The predictive processes (Banerjee et al., 2008) corresponds to a specific choice of the basis functions and the covariance matrix, but does not give a very flexible type of non-stationarity. All such methods are variations of the same concept, but lead to different computational schemes with different computational properties. The dimension of the finite-dimensional basis is in all cases used to control the computational cost and the novelty of each method lies in how the basis elements are selected and connected to each other, and the computational methods used to exploit the structure.

An overview of the literature before around 2010 is given in Sampson (2010). This overview also includes less well-known methods such as the piece-wise Gaussian process of Kim et al. (2005), processes based on weighted linear combination of stationary processes (Fuentes, 2001, 2002a,b; Nott and Dunsmuir, 2002).

Recently, a new class of methods based on the SPDE-approach introduced by Lindgren et al. (2011) is emerging. This class of methods is based on a representation of the spatial field as a solution of a stochastic partial differential equation (SPDE) with spatially varying coefficients. The methodology is closely connected with Gaussian Markov random fields (GMRFs) (Rue and Held, 2005) and is able to handle more observations than is possible with the deformation method and the process convolution method. In a similar way as a spatial GMRF describes local behaviour for a discretely indexed process, an SPDE describes local behaviour for a continuously indexed process. This locality in the continuous description can be transferred to a GMRF approximation of the solution of the SPDE, and gives a GMRF with a spatial Markovian structure that can be exploited in computations.

This type of methodology has been applied to global ozone data (Bolin and Lindgren, 2011) and to annual precipitation in Norway with covariates in the covariance structure (Ingebrigtsen et al., 2014, 2015). Additionally, Sigrist et al. (2012) used similar type of modelling to handle a spatio-temporal process where wind direction and strength enters in the covariance structure.

Despite all the work that has been done in non-stationary spatial modelling, it is still
an open field where no model stands out as the clear choice. However, we believe that
modelling locally such as in the SPDE-based models is more attractive than modelling
globally such as in the deformation method and the process convolution method. There-
fore, we choose to use an extension of the model by [Fuglstad et al. 2015] that allows
for both a spatially varying correlation structure and a spatially varying marginal vari-
ance. This method is closely connected to the already well-known deformation method
of [Sampson and Guttorp 1992] and the Matérn-like process convolution of [Paciorek and
Schervish 2006], but is focused at the local behaviour and not the global behaviour.

In a similar way as in the model of [Paciorek and Schervish 2006] the global structure
is defined through the combination of ellipses at each location that describe anisotropy.
However, their model only combines the ellipses at two locations and does not account for
the local behaviour between locations. The new model incorporates the local anisotropy
everywhere into the covariance for each pair of locations and is not the same as the
model of [Paciorek and Schervish 2006]. The model works in a similar way as the
deformation method. However, instead of describing a global deformation, the ellipses
augment the local distances around each point and describe locally a change of distances
such that lengths are different in different directions, but does not, in general, lead to
a deformation of $\mathbb{R}^2$ to $\mathbb{R}^2$. Such local modelling tends to lead to a deformation in an
ambient space of dimension higher than 2. The interest of this paper is to study the
challenges and results of applying the method to a dataset of annual precipitation in the
conterminous US.

1.2 Annual precipitation in the conterminous US

This case study of non-stationarity will use the measurements of monthly total precipi-
tation at different measurement stations in the conterminous US for the years 1895–1997
that are available at [http://www.image.ucar.edu/GSP/Data/US.monthly.met/]. This
dataset was chosen because it is publicly available in a form that is easily downloaded
and loaded into software, and because the large spatial scale of the dataset and the
complexity of the physical process that generates weather makes it intuitively feels like
there must be non-stationarity in the dataset. The main focus of this study is to see
how much non-stationarity improves the spatial predictions compared to a stationary
model, but the approach could also be used to gain insight about the spatial structure
of climate in the conterminous US (Smith, 1996).

In total there are 11918 measurement stations in the dataset, but measurements are
only available at a subset of the stations each month and the rest of the stations have
in-filled data [Johns et al. 2003]. For each year, we aggregate the monthly data at those
stations which have measurements available at all months in that year and produce a
dataset of yearly total precipitation. This gives a different number of locations for each
year. We then take the logarithm of each observation to create the transformed data
that is used in this paper. Figure 1 shows the transformed data at the 7040 stations
available for 1981. The only covariate available in the dataset is the elevation at each
station, and no work was done to find other covariates from alternate sources.

If the focus were to model this data in the best possible way, it would, in general, be
good to look for more covariates or consider alternatives such as spatially heterogeneous coefficients (Fotheringham et al., 2002; Gelfand et al., 2003) before using a full non-stationary model. Non-stationarity can be achieved both through a flexible model for the mean structure and through a flexible model for the covariance structure. The difference between the two alternatives lies in the interpretation of the model, and with focus on computations and predictions, a simple second-order structure and a flexible first-order structure could be an attractive option. But it is our explicit intention to focus on non-stationarity in the second-order structure and we keep the model for the mean simple.

We will assume that the transformed data can be treated as Gaussian, which is a reasonable assumption because we are modelling annual precipitation data. However, it would not be a reasonable assumption, for example, for daily data, and it would be necessary to consider not only how to deal with non-stationarity, but also how to deal with the lack of Gaussianity. Bolin and Wallin (2013) compare the predictions made by a stationary Gaussian model, a stationary Gaussian model for transformed data and two stationary non-Gaussian models for monthly precipitation for two different months from the same dataset as in this paper. They apply the non-Gaussian model of Bolin (2014), but do not find clear evidence that one model perform better than the others. The approach of Bolin (2014) is based on the discretization of an SPDE in a similar way as the approach in this paper, but using a non-Gaussian noise process, and a possible extension of the non-stationary model in this paper would be to non-Gaussian data.

The main motivation for focusing on the year 1981 is that Paciorek and Schervish (2006) previously studied the annual precipitation in the subregion of Colorado for this year. They did not see major improvements over a stationary model and our preliminary analysis showed that there was little non-stationarity left in the subregion after introducing a joint mean and elevation. However, Colorado constitutes a small part of the conterminous US, and as shown in Figure 2 there are large differences in the topography of the western and the eastern part of the conterminous US. A large proportion of the western part is mountainous whereas in the eastern part a large proportion is mostly flat. This varied topography is a strong indication that the process cannot possibly be stationary.

To substantiate our claims of non-stationarity we explore the difference in the covariance structure in the western and eastern part through variograms. The data from years 1971–1985 is selected and divided into two regions: the area west of 100°W and the area east of 100°W. For each year the variogram of each region is calculated. Figure 3 shows that there is no overlap between the variograms of the western region and the eastern region. There is significant variation within each region, but the overall appearance clearly indicates different covariance structures within the regions. Based on the evidence of non-stationarity seen in the variograms for the full region, we want to know if a non-stationary model will improve the predictions. It has been observed by several authors (Schmidt et al., 2011; Neto et al., 2014) and it has also been the experience of the authors that non-stationary models do not lead to much difference in the predicted values, and that the differences are found in the prediction variances. However, predictions
should always have associated error estimates and when we write improved predictions, we are interested in whether the predictive distributions, summarized by the predicted values and their associated prediction variances, better describe the observed values.

There are two cases of interest: a single realization and multiple realizations. In the former it is impossible to separate the non-stationarity in the mean and in the covariance structure, and the non-stationary model might be more accurately described as adaptive smoothing, but many spatial datasets are of this form and a non-stationary model might still perform better than a stationary model. We will investigate both of these cases and evaluate whether the non-stationary model improves predictions and whether the computational costs are worth it. It is clear that stationarity is not the truth, but that does not mean that it does not necessarily constitutes a sufficient model for predictions.
Figure 3: Estimated semi-variograms for the years 1971 to 1985 using the locations to the west of 100°W coloured in blue and marked with circles and locations to the east of 100°W coloured in red and marked with crosses.

1.3 Overview

The paper is divided into five sections. Section 2 describes how we model the data. We discuss what type of non-stationarity is present in the model and how it is specified, how we parametrize the non-stationarity and how we perform computations with the non-stationary model. Then in Section 3 a hierarchical model incorporating the non-stationary model is applied to annual precipitation in a single realization setting, and in Section 4 the data is studied from a multiple realizations perspective. The differences between the estimated covariance structures and the prediction scores for the different models are discussed. The paper ends with discussion and concluding remarks in Section 5.

2 Modelling the data

Before analyzing the data we need to introduce the model that will be used. Particularly, we need to say which types of non-stationarity will be present in the model and how this non-stationarity will be modelled. A good spatial model should provide a useful way to do both the theoretical modelling and the associated computations. We first discuss how non-stationarity is introduced and then how to parametrize the non-stationary.

2.1 Modelling the non-stationarity

It is difficult to specify a global covariance function when one only has intuition about local behaviour. Consider the situation in Figure 4. The left hand side and the right hand side have locally large “range” in the horizontal direction and somewhat shorter “range” in the vertical direction, and the middle area has locally much shorter “range” in the horizontal direction, but slightly longer in the vertical direction. We write “range”
Figure 4: Example of a correlation function caused by varying local behaviour. For each location marked with a black cross, the 0.9, 0.7, 0.5, 0.25 and 0.12 level contours of the correlation function are shown.

with quotation marks because the concept of a global range does not have a well-defined meaning in non-stationary modelling. Instead we will think of range as a local feature and use the word to mean what happens to dependency in a small region around each point. From the figure one can see that for the point in the middle, the chosen contours look more or less unaffected by the two other regions since they are fully contained in the middle region, but that for the point on the left hand side and the point on the right hand side, there is much skewness introduced by the transition into a different region.

It would be hard to specify a fitting global correlation function for this situation. However, if one instead starts with an isotropic process and then stretches the left hand side and the right hand side in the x-direction, the task is much easier. This is a flexible way to create interesting covariance structures and is the core of the deformation method [Sampson and Guttorp, 1992], but can be challenging since one has to create a valid global deformation. We present instead a model where the modelling can be done locally without worrying about the global structure. We let the local structure automatically specify a valid global structure. In this example one would only specify that locally the range is longer in the horizontal direction in the left hand side and the right hand side, and then let this implicitly define the global structure without directly modelling a global deformation.

In the SPDE-based approach the correlation between two spatial locations is determined implicitly by the behaviour between the spatial locations. If there are mountains, the model could specify that locally the distances are longer than they appear on the map and the correlation will decrease more quickly when crossing those areas, and if there are plains, the model could specify that distances are shorter than they appear on the map and the correlation will decrease more slowly in those areas. A major advantage of this approach is that the local specification naturally leads to a spatial GMRF
with good computational properties. It is possible to approximate the local continuous description with a local discrete description. The result is a spatial GMRF with a very sparse precision matrix.

The starting point for the non-stationary SPDE-based model is the stationary SPDE introduced in Lindgren et al. (2011),

\[(\kappa^2 - \nabla \cdot \nabla)u(\mathbf{s}) = \sigma \mathcal{W}(\mathbf{s}), \quad \mathbf{s} \in \mathbb{R}^2, \tag{1}\]

where \(\kappa > 0\) and \(\sigma > 0\) are constants, \(\nabla = \left( \frac{\partial}{\partial x}, \frac{\partial}{\partial y} \right)^T\) and \(\mathcal{W}\) is a standard Gaussian white noise process. The SPDE describes the GRF \(u\) as a smoothed version of the Gaussian white noise on the right hand side of the equation. Whittle (1954, 1963) showed that any stationary solution of this SPDE has the Matérn covariance function

\[r(\mathbf{s}_1, \mathbf{s}_2) = \frac{\sigma^2}{4\pi \kappa^2} (\kappa \|\mathbf{s}_2 - \mathbf{s}_1\|) K_1(\kappa \|\mathbf{s}_2 - \mathbf{s}_1\|), \tag{2}\]

where \(K_1\) is the modified Bessel function of second kind, order 1. This covariance function is a member of the commonly-used Matérn family of covariance functions, and one can see from Equation (2) that one can first use \(\kappa\) to select the range and then \(\sigma\) to achieve the desired marginal variance. In some methods for non-stationarity it is possible to spatially vary the smoothness, but this is not a feature that is available in the non-stationary model presented here. However, with the flexibility present in the rest of the non-stationarity it is not clear if the smoothness would be jointly identifiable.

The next step is to generate a GRF with an anisotropic Matérn covariance function. The cause of the isotropy in SPDE (1) is that the Laplacian, \(\Delta = \nabla \cdot \nabla\) is invariant to a change of coordinates that involves rotation and translation. To change this a \(2 \times 2\) matrix \(H > 0\) is introduced into the operator to give the SPDE

\[(\kappa^2 - \nabla \cdot H\nabla)u(\mathbf{s}) = \sigma \mathcal{W}(\mathbf{s}). \tag{3}\]

This choice is closely related to the change of coordinates \(\tilde{\mathbf{s}} = H^{1/2} \mathbf{s}\) (Fuglstad et al., 2015, Section 3) and gives the covariance function

\[r(\tilde{\mathbf{s}}_1, \tilde{\mathbf{s}}_2) = \frac{\sigma^2}{4\pi \kappa^2 \sqrt{\det(H)}} (\kappa \|H^{-1/2}(\tilde{\mathbf{s}}_2 - \tilde{\mathbf{s}}_1)\|) K_1(\kappa \|H^{-1/2}(\tilde{\mathbf{s}}_2 - \tilde{\mathbf{s}}_1)\|). \tag{4}\]

Compared to Equation (2), there is a change in the marginal variance and a directionality is introduced through a distance measure different than the standard Euclidean distance. Figure 5 shows how the eigenpairs of \(H\) and the value of \(\kappa\) act together to control range. One can see that the construction leads to elliptic iso-covariance curves. In what follows \(\sigma\) is assumed to be equal to 1 since the marginal variance can be controlled by varying \(\kappa^2\) and \(H\) together.

The final step is to construct a non-stationary GRF where the local behaviour at each location is governed by SPDE (3) with \(\sigma = 1\) and the values of \(\kappa^2\) and \(H\) varying over the domain. The intention is to create a GRF by chaining together processes with different local covariance structures. The SPDE becomes

\[(\kappa^2(\mathbf{s}) - \nabla \cdot H(\mathbf{s})\nabla)u(\mathbf{s}) = \mathcal{W}(\mathbf{s}). \tag{5}\]
Figure 5: Iso-correlation curve for the 0.6 level, where \((\lambda_1, \vec{v}_1)\) and \((\lambda_2, \vec{v}_2)\) are the eigenpairs of \(\mathbf{H}\).

For technical reasons concerned with the discretization of the SPDE, \(\kappa^2\) is required to be continuous and \(\mathbf{H}\) is required to be continuously differentiable. This does not present any problems and is easily achieved by using continuously differentiable basis functions for \(\kappa^2\) and \(\mathbf{H}\). The restricted form where \(\kappa^2\) is constant was investigated in Fuglstad et al. \(2015\), but this restricted form only allows for varying local anisotropy without control over the marginal variances. This extended model allows for spatially varying “range”, anisotropy and marginal variance.

The non-stationary covariance structure is fully described by SPDE \(\mathbf{5}\), but before the model can be used in practice the description must be brought into a form which is useful for computations. This can be done by discretizing the SPDE using a finite-dimensional basis expansion where the distribution of the coefficients is a sparse GMRF that possesses approximately the same covariance structure as \(u\). See \(\text{A}\) for more details. This kind of construction alleviates one of the largest problems with GMRFs, namely that they are hard to specify in a spatially coherent manner. The computational benefits of spatial GMRFs are well known, but a GMRF needs to be constructed through its conditional distributions and it is notoriously hard to do this for non-stationary models. But with this approach it is possible to model the problem with an SPDE and then do computations with the computational benefits of a spatial GMRF.

### 2.2 Parametrizing the non-stationarity

Before we can turn the theoretical and computational description of the non-stationary model into a statistical model, we need to describe the non-stationarity through parameters. This means both decomposing the model into parameters and connecting the
parameters together through a penalty.

The first step is to decompose the function $H$, which must give positive definite $2 \times 2$ matrices at each location, into simpler functions. One usual way to do this is to use two strictly positive functions $\lambda_1$ and $\lambda_2$ for the eigenvalues and a function $\phi$ for the angle between the $x$-axis and the eigenvector associated with $\lambda_1$. However, with a slight re-parametrization $H$ can be written as the sum of an isotropic effect, described by a constant times the identity matrix, plus an additional anisotropic effect, described by direction and magnitude.

Express $H$ through the scalar functions $\gamma, v_x$ and $v_y$ by

$$H(\vec{s}) = \gamma(\vec{s})I_2 + \begin{bmatrix} v_x(\vec{s}) & v_y(\vec{s}) \\ v_y(\vec{s}) & v_x(\vec{s}) \end{bmatrix},$$

where $\gamma$ is required to be strictly positive. The eigendecomposition of this matrix has eigenvalue $\lambda_1(\vec{s}) = \gamma(\vec{s}) + v_x(\vec{s})^2 + v_y(\vec{s})^2$ with eigenvector $\vec{v}_1(\vec{s}) = (v_x(\vec{s}), v_y(\vec{s}))$ and eigenvalue $\lambda_2(\vec{s}) = \gamma(\vec{s})$ with eigenvector $\vec{v}_2(\vec{s}) = (-v_y(\vec{s}), v_x(\vec{s}))$. From Figure 5 this means that for a stationary model, $\gamma$ affects the length of the shortest semi-axis of the iso-correlation curves and $\vec{v}$ specifies the direction of and how much larger the longest semi-axis is. The above decomposition through $\gamma, v_x$ and $v_y$ is general and is valid for every symmetric positive-definite $2 \times 2$ matrix.

Since we want flexible covariance structures, some representation of the functions $\kappa^2$, $\gamma$, $v_x$ and $v_y$ is needed. To ensure positivity of $\kappa^2$ and $\gamma$, they are first transformed into $\log(\kappa^2)$ and $\log(\gamma)$. Each of these functions will be expanded in a basis, and requires a penalty that imposes regularity and makes sure the function is not allowed to vary too much. The choice was made to give $\log(\kappa^2)$, $\log(\gamma)$, $v_x$ and $v_y$ spline-like penalties. The steps that follow are the same for each function. Therefore, they are only shown for $\log(\kappa^2)$.

The function $\log(\kappa^2)$ is given a penalty according to the distribution generated from the SPDE

$$-\Delta \log(\kappa^2(\vec{s})) = W_{\kappa}(\vec{s})/\sqrt{\tau_\kappa}, \quad \vec{s} \in \mathcal{D},$$

where $\tau_\kappa > 0$ is the parameter controlling the penalty, with the Neumann boundary condition of zero derivatives at the edges. This extra requirement is used to restrict the resulting distribution so it is only invariant to the addition of a constant function, and the penalty parameter is used to control how much $\log(\kappa^2)$ can vary from a constant function. The penalty defined through SPDE (6) is in this paper called a two-dimensional second-order random walk due to its similarity to a one-dimensional second-order random walk (Lindgren and Rue, 2008).

The first step of making the above penalty applicable for the computational model is to expand $\log(\kappa^2)$ in a basis through a linear combination of basis functions,

$$\log(\kappa^2(\vec{s})) = \sum_{i=1}^{k} \sum_{j=1}^{l} \alpha_{ij} f_{ij}(\vec{s}),$$

where $\{\alpha_{ij}\}$ are the parameters and $\{f_{ij}\}$ are real-valued basis functions. For convenience, the basis is chosen in such a way that all basis functions satisfy the boundary
conditions specified in SPDE (6). If this is done, one immediately satisfies the boundary condition. The remaining tasks are then to decide which basis functions to use and what the resulting penalties on the parameters are.

Due to a desire to make \( H \) continuously differentiable and a desire to have “local” basis functions, the basis functions are chosen to be based on 2-dimensional, second-order B-splines (piecewise-quadratic functions). The basis is constructed as a tensor product of two 1-dimensional B-spline bases constrained to satisfy the boundary condition.

The penalty is based on the distribution defined by SPDE (6), so the final step is to determine a Gaussian distribution for the parameters such that the distribution of \( \log(\kappa^2) \) is close to a solution of SPDE (6). The approach taken is based on a least-squares formulation of the solution and is described in B. Let \( \vec{\alpha} \) be the \( \{\alpha_{ij}\} \) parameters stacked row-wise, then the result is that \( \vec{\alpha} \) should be given a zero-mean Gaussian distribution with precision matrix \( \tau_{\kappa} Q_{RW2} \). This matrix has rank \( (kl - 1) \), due to the Neumann boundary conditions, and the distribution is invariant to the addition of a vector of only the same values, but for convenience the penalty will still be written as \( \vec{\alpha} \sim N_{kl}(\vec{0}, Q_{RW2}/\tau_{\kappa}) \).

2.3 Hierarchical model

Observations \( y_1, y_2, \ldots, y_N \) are made at locations \( \vec{s}_1, \vec{s}_2, \ldots, \vec{s}_N \). The observed value at each location is assumed to be the sum of a fixed effect due to covariates, a spatial “smooth” effect and a random effect. The covariates at location \( \vec{s}_i \) are described by the \( p \)-dimensional row vector \( \vec{x}(\vec{s}_i)^T \) and the spatial field is denoted by \( u \). This gives the observation equation

\[
y_i = \vec{x}(\vec{s}_i)^T \vec{\beta} + u(\vec{s}_i) + \epsilon_i,
\]

where \( \vec{\beta} \) is a \( p \)-variate random vector for the coefficients of the covariates and \( \epsilon_i \sim \mathcal{N}(0, 1/\tau_{\text{noise}}) \) is the random effect for observation \( i \), for \( i = 1, 2, \ldots, N \).

The \( u \) is modelled and parametrized as described in the previous sections and the GMRF approximation is used for computations. In this GMRF approximation the domain is divided into a regular grid consisting of rectangular cells and each element of the GMRF approximation describes the average value on one of these cells. So \( u(\vec{s}_i) \) is replaced with the approximation \( \vec{e}(\vec{s}_i)^T \vec{u} \), where \( \vec{e}(\vec{s}_i)^T \) is the \( mn \)-dimensional row vector selecting the element of \( \vec{u} \) which corresponds to the cell which contains location \( \vec{s}_i \). In total, this gives

\[
\vec{y} = \vec{X}\vec{\beta} + \vec{E}\vec{u} + \vec{\epsilon},
\]

where \( \vec{y} = (y_1, y_2, \ldots, y_N) \), the matrix \( \vec{X} \) has \( \vec{x}(\vec{s}_1)^T, \ldots, \vec{x}(\vec{s}_N)^T \) as rows and the matrix \( \vec{E} \) has \( \vec{e}(\vec{s}_1)^T, \ldots, \vec{e}(\vec{s}_N)^T \) as rows. In this equation the spatial effect is approximated with a discrete model, but the covariate has not been gridded and is at a higher resolution than the grid.

The model for the observations can also be written in the form

\[
\vec{y} | \vec{\beta}, \vec{u}, \log(\tau_{\text{noise}}) \sim \mathcal{N}_N(\vec{X}\vec{\beta} + \vec{E}\vec{u}, \vec{I}_N / \tau_{\text{noise}}).
\]
The parameter $\tau_{\text{noise}}$ acts as the precision of a joint effect from measurement noise and small scale spatial variation (Diggle and Ribeiro, 2007). We make the underlying model for the $p$-dimensional random variable $\vec{\beta}$ proper by introducing a weak Gaussian penalty,

$$\vec{\beta} \sim \mathcal{N}_p(\vec{0}, \frac{1}{\tau\beta} \mathbf{I}_p).$$

The penalty can be made stronger, but we do not believe it will have a strong effect on the estimates for this dataset with only an intercept and one covariate.

To describe the full hierarchical model, we introduce symbols to denote the parameters that control the spatial field $u$. Denote the parameters that control $\log(\kappa_2)$, $\log(\gamma)$, $v_x$ and $v_y$ by $\vec{\alpha}_1$, $\vec{\alpha}_2$, $\vec{\alpha}_3$ and $\vec{\alpha}_4$, respectively. Further, denote the corresponding penalty parameters for each function by $\tau_1$, $\tau_2$, $\tau_3$ and $\tau_4$. With this notation the full model becomes

Stage 1: $\vec{y} | \vec{\beta}, \vec{u}, \log(\tau_{\text{noise}}) \sim \mathcal{N}_N(\mathbf{X}\vec{\beta} + \mathbf{E}\vec{u}, \frac{1}{\tau_{\text{noise}}} \mathbf{I}_N)$

Stage 2: $\vec{u} | \vec{\alpha}_1, \vec{\alpha}_2, \vec{\alpha}_3, \vec{\alpha}_4 \sim \mathcal{N}_{nm}(\vec{0}, \mathbf{Q}^{-1}), \vec{\beta} \sim \mathcal{N}_p(\vec{0}, \frac{1}{\tau\beta} \mathbf{I}_p)$

Stage 3: $\vec{\alpha}_i | \tau_i \sim \mathcal{N}_{kl}(\vec{0}, \frac{1}{\tau_i} \mathbf{I}_p)$ for $i = 1, 2, 3, 4$,

where $\tau_1$, $\tau_2$, $\tau_3$, $\tau_4$ and $\tau\beta$ are penalty parameters that must be pre-selected.

An important model choice when constructing the GMRF approximation of the spatial process is the selection of the resolution of the approximation. The approximation does not allow for variation of the spatial field within a grid cell and the spatial resolution must be chosen high enough to capture variations on the scale at which observations were made. The variation at sub-grid scale cannot be captured by the approximation and will be captured by the nugget effect.

### 2.4 Penalized likelihood and inference

The two things of main interest to us in this case study are the covariance parameters $\vec{\theta} = (\vec{\alpha}_1, \vec{\alpha}_2, \vec{\alpha}_3, \vec{\alpha}_4, \log(\tau_{\text{noise}}))$ and the predictive distributions for unmeasured locations. To estimate the covariance parameters, we need the integrated likelihood where the latent field consisting of the coefficients of the fixed effects and the spatial effect are integrated out. This integration can be done explicitly because the spatial field by construction is Gaussian and the parameters of the fixed effects are Gaussian due to the choice of a Gaussian penalty.

First, collect the fixed effect and the spatial effect in $\vec{z} = (\vec{u}^T, \vec{\beta}^T)$. The model given the value of $\vec{\theta}$ can then be written as

$$\vec{z} | \vec{\theta} \sim \mathcal{N}_{mn+p}(\vec{0}, \mathbf{Q}_z^{-1})$$

and

$$\vec{y} | \vec{z}, \vec{\theta} \sim \mathcal{N}_N(\mathbf{S}\vec{z}, \frac{1}{\tau_{\text{noise}}} \mathbf{I}_N)$$,

where

$$\mathbf{S} = \begin{bmatrix} \mathbf{E} & \mathbf{X} \end{bmatrix} \quad \text{and} \quad \mathbf{Q}_z = \begin{bmatrix} \mathbf{Q} & \mathbf{0} \\ \mathbf{0} & \frac{1}{\tau\beta} \mathbf{I}_p \end{bmatrix}.$$
We then use the fact that both these distributions are Gaussian to integrate out \( \mathbf{z} \) from the likelihood, as shown in C. This gives the full penalized log-likelihood

\[
\log(\pi(\tilde{\theta} | \tilde{y})) = \text{Const} - \frac{1}{2} \sum_{i=1}^{4} \mathbf{y}_i^T Q_{\text{RW2}} \mathbf{y}_i \cdot \tau_i + \frac{1}{2} \log(\det(Q_z)) + \frac{N}{2} \log(\tau_{\text{noise}}) + \\
- \frac{1}{2} \log(\det(Q_C)) - \frac{1}{2} \mathbf{y}_C^T Q_C \mathbf{y}_C \cdot \tau_{\text{noise}} - \frac{\tau_{\text{noise}}}{2} (\tilde{y} - S \tilde{\mu}_C)^T (\tilde{y} - S \tilde{\mu}_C),
\]

where \( Q_C = Q_z + S^T S \cdot \tau_{\text{noise}} \) and \( \tilde{\mu}_C = Q_C^{-1} S^T \tilde{y} \cdot \tau_{\text{noise}}. \)

The first step of the inference scheme is to estimate the covariance parameters \( \tilde{\theta} \) with the value \( \hat{\tilde{\theta}} \) that maximizes Equation (8). This value is then used to calculate predictions and prediction standard deviations at new locations \( \tilde{y}^* \) by using the predictive distribution \( \tilde{y}^* | \hat{\tilde{\theta}}, \tilde{y} \). However, the penalty parameters that control the penalty of the covariance parameters are difficult to estimate. The profile likelihoods are hard to calculate and there is not enough information on such a low stage of the hierarchical model to estimate them together with the covariance parameters. Thus they have to be pre-selected, based on intuition about how much the covariance structure should be allowed to vary, or chosen with a cross-validation procedure based on a scoring rule for the predictions.

During implementation of the inference scheme it became apparent that an analytic expression for the gradient was needed for the optimization to converge. Its form is given in D and its value can be computed for less cost than a finite difference approximation of the gradient for the number of parameters used in the application in this paper. The calculations require the use of techniques for calculating only parts of the inverse of a sparse precision matrix (Rue and Held, 2010).

3 Non-stationarity in a single realization

3.1 Adaptive smoothing framework

We begin by considering the common situation in spatial statistics where only a single realization is available. In this situation it is theoretically impossible to separate non-stationarity in the mean and in the covariance structure, and the non-stationary model is better described as adaptive smoothing. The non-stationary model allows the degree of smoothing to vary over space, and areas with long range will have high smoothing and areas with short range will have low smoothing. In practice, parts of the non-stationarity in the mean structure will be captured in the covariance structure, but this is not necessarily a problem and might lead to better predictions. The main interest is finding out whether the complex non-stationary model improves predictions at unobserved locations and whether the computational costs are worth it.

We select the year 1981 which has 7040 measurement stations and want to predict the annual precipitation in the entire conterminous US with associated prediction standard deviations. Two covariates are used in the mean structure: an intercept and a linear effect of elevation. This means that the design matrix, \( X \), in Equation (7) has two
Table 1: Estimated values of the parameters and associated approximate standard deviations for the stationary model.

| Parameter          | Estimate | Standard deviation |
|--------------------|----------|--------------------|
| log($\kappa^2$)   | -1.75    | 0.15               |
| log($\gamma$)     | -0.272   | 0.042              |
| $v_x$              | 0.477    | 0.053              |
| $v_y$              | -0.313   | 0.057              |
| log($\tau_{\text{noise}}$) | 4.266 | 0.030 |

columns. The first column contains only ones, and corresponds to the intercept, and the second column contains elevations measured in kilometres. There should be strong information about the two covariates and a weak penalty is applied to the coefficients of the fixed effects, $\mathbf{\beta} \sim \mathcal{N}_2(\mathbf{0}, \mathbf{I}_2 \cdot 10^4)$.

3.2 Stationary model

The spatial effect is constructed on a rectangular domain with longitudes from $130.15^\circ W$ to $60.85^\circ W$ and latitudes from $21.65^\circ N$ to $51.35^\circ N$. This is larger than the actual size of the conterminous US as can be seen in Figure 1 and is chosen to reduce boundary effects. The domain is discretized into a $400 \times 200$ grid and the parameters $\log(\kappa^2)$, $\log(\gamma)$, $v_x$, $v_y$ and $\log(\tau_{\text{noise}})$ are estimated. In this case the second order random walk penalty is not used as no basis (except a constant) is needed for the functions. The estimated values with associated approximate standard deviations are shown in Table 1. The approximate standard deviations are calculated from the observed information matrix.

From Section 2.1 one can see that the estimated model implies a covariance function approximately equal to the Matérn covariance function

$$r(\bar{s}_1, \bar{s}_2) = \hat{\sigma}^2 \left| \left( \frac{\mathbf{H}}{\hat{\kappa}^2} \right)^{-1/2} (\bar{s}_2 - \bar{s}_1) \right| K_1 \left( \left| \left( \frac{\mathbf{H}}{\hat{\kappa}^2} \right)^{-1/2} (\bar{s}_2 - \bar{s}_1) \right| \right),$$

where $\hat{\sigma}^2 = 0.505$ and

$$\frac{\mathbf{H}}{\hat{\kappa}^2} = \begin{bmatrix} 5.71 & -0.86 \\ -0.86 & 4.96 \end{bmatrix},$$

together with a nugget effect with precision $\hat{\tau}_{\text{noise}} = 71.2$. Figure 6 shows contours of the estimated covariance function with respect to a chosen location. One can see that the model gives high dependence within a typical-sized state, whereas there is little dependence between the centres of different typically-sized states.

Next, the parameter values are used together with the observed logarithms of annual precipitations to predict the logarithm of annual precipitation at the centre of each cell in the discretization. The elevation covariate for each location is selected from bilinear interpolation from the closest points in the high resolution elevation data set GLOBE (Hastings et al., 1999). The predictions and prediction standard deviations are shown in Figures 7(a) and 7(c). Since there only are observations within the conterminous US and this is the area of interest, the locations outside are coloured white.
Figure 6: The 0.95, 0.70, 0.50, 0.36, 0.26, 0.19, 0.14 and 0.1 level correlation contours of the estimated covariance function for the stationary model.

Figure 7: Predicted values and prediction standard deviations for the stationary model and the non-stationary model.
3.3 Non-stationary model

The parameters $\tau_1$, $\tau_2$, $\tau_3$ and $\tau_4$, that appear in the penalty for the functions $\log(\kappa^2)$, $\log(\gamma)$, $\nu_x$ and $\nu_y$, respectively, have to be chosen before the rest of the inference is started. The parameters are chosen with 5-fold cross-validation based on the log-predictive density. The data is randomly divided into five parts and in turn one part is used as test data and the other four parts are used as training data. For each choice of $\tau_1$, $\tau_2$, $\tau_3$ and $\tau_4$ the cross-validation error is calculated by

$$CV(\tau_1, \tau_2, \tau_3, \tau_4) = -\frac{1}{5} \sum_{i=1}^{5} \log(\pi(y_i^*|\tilde{\theta}_i)),$$

where $y_i^*$ is the test data and $\tilde{\theta}_i$ is the estimated covariance parameters based on the training data $\tilde{y}_i$ using the selected $\tau$-values. The cross validation is done over $\log(\tau_i) \in \{2, 4, 6, 8\}$ for $i = 1, 2, 3, 4$. We selected four values for each parameter to have a balance between the need to test strong and weak penalties and to make the problem computationally feasible. Controlling the penalty on non-stationarity is important, but appropriate penalty values are not easily deduced from the model. Therefore, different values were tested to determine values of $\tau_i$ that corresponds to a weak penalty and a strong penalty and then four points were chosen linearly on log-scale since $\tau_i$ acts as a scale parameter. We use the same domain size as for the stationary model, but reduce the grid size to $200 \times 100$ with $8 \times 4$ basis functions for each function. The choice that gave the smallest cross-validation error was $\log(\tau_1) = 2$, $\log(\tau_2) = 4$, $\log(\tau_3) = 2$ and $\log(\tau_4) = 8$.

After the penalty parameters are selected, the grid size is increased to $400 \times 200$ and each of the four functions in the SPDE is given a $16 \times 8$ basis functions. Together with the precision parameter of the random effect this gives a total of 513 parameters. These parameters are estimated together based on the integrated likelihood. Note that there are not 513 “free” parameters as they are connected together in four different penalties enforcing slowly changing functions. This means that an increase in the number of parameters increases the resolutions of the functions, but not directly the degrees of freedom in the model.

The nugget effect is estimated to have a precision of $\hat{\tau}_{\text{noise}} = 107.4$. The estimates of $\kappa^2$ and $H$ are not shown since the exact values themselves are not interesting. We calculate instead the marginal standard deviations for all locations and 0.7 level correlation contours for selected locations in Figure 8(a) and Figure 8(b) respectively. From these figures one can see that the estimated covariance structure is different from the estimated covariance structure for the stationary model shown in Figure 6. In the non-stationary model we have a much longer range in the eastern part and a much short range in the mountainous areas in the west.

The estimated covariance structure implies strong smoothing in the eastern region and weak smoothing in the western region. This must be understood to say something about both how well the covariates describe the data at different locations and the underlying non-stationarity in the covariance structure of the physical phenomenon. In
Figure 8: Estimated covariance structure of the spatial field. (a) Marginal standard deviations (b) Contours of 0.7 correlation for selected locations marked with red crosses.

In this case there is a good fit for the elevation covariate in the mountainous areas in the western part, but it offers less information in the eastern part. From Figure 1 one can see that at around longitude 97° W there is an increase in precipitation which cannot be explained by elevation, and thus is not captured by the covariates. This jump must therefore be explained by the covariance structure, and in this case it is explained by having the covariates fit well in the western region and explaining the high values in the eastern region as being caused, randomly, by a spatial process with a long range.

In the same way as in Section 3.2 the logarithm of annual precipitation is predicted at the centre of each cell in the discretization. This gives predictions for 400 × 200 regularly distributed locations, where the value of the elevation covariate at each location is selected with bilinear interpolation from the closest points in the GLOBE [Hastings et al. 1999] dataset. The prediction and prediction standard deviations are shown in Figures 7(b) and 7(d). As for the stationary model, the values outside the conterminous US are coloured white. One can see that the overall look of the predictions is similar to the predictions from the stationary model, but that the prediction standard deviations differ. The prediction standard deviations vary strongly over the spatial domain because of the extreme differences in spatial range for the estimated non-stationary model.

3.4 Evaluation of predictions

The predictions of the stationary model and the non-stationary model are compared with the continuous rank probability score (CRPS) [Gneiting et al. 2005] and the logarithmic scoring rule. CRPS is defined for a univariate distribution as

\[ \text{crps}(F, y) = \int_{-\infty}^{\infty} (F(y) - 1(y \leq t))^2 \, dt, \]

where \( F \) is the distribution function of interest, \( y \) is an observation and \( 1 \) is the indicator function. This gives a measure of how well a single observation fits a distribution. The
total score is calculated as the average CRPS for the test data,

\[ \text{CRPS} = \frac{1}{N} \sum_{i=1}^{N} \text{crps}(F_k, y_k), \]

where \( \{y_k\} \) is the test data and \( \{F_k\} \) are the corresponding marginal predictive distributions given the estimated covariance parameters and the training data. The logarithmic scoring rule is based on the joint predictive distribution of the test data \( \tilde{y}^* \) given the estimated covariance parameters \( \tilde{\theta} \) and the training data \( \tilde{y} \),

\[ \text{LogScore} = -\log \pi(\tilde{y}^*|\tilde{\theta}, \tilde{y}). \]

The comparison of the models is done using holdout sets where each holdout set consists of 20% of the locations chosen randomly. The remaining 80% of the locations are used to estimate the parameters and to predict the values at the locations in the holdout set. This procedure is repeated 20 times. For each repetition the CRPS, the logarithmic score and the root mean square error (RMSE) are calculated. From Figure 9 one can see that measured by both log-predictive score and CRPS the non-stationary model gives better predictions, but that the RMSE does not show any improvement.

However, the RMSE is based only on the point predictions and does not incorporate the prediction variances. The log-predictive score and the CRPS are more interesting since they say something about how well the predictive distributions fit. The difference in log-predictive score is large and indicates that the non-stationary model is better, but the difference in CRPS is small and indicates only a small improvement. The likely cause for this is that the log-predictive score evaluates the joint predictive distributions and there are differences which are not showing in the univariate predictive distributions.

The full cross-validation procedure for selecting the penalty parameters is expensive and takes weeks and must be evaluated against the potential gain in any application. The results show that the choice of scoring rule has a strong influence on the conclusion of whether the non-stationary model was worth it. The CRPS does not show evidence that all the extra computation time was worth it, but according to the log-predictive score there is a large improvement.

### 3.5 Criticism

The log-predictive score and CRPS are better for the non-stationary model for each hold-out set, but the covariance structure shown in Figure 8 is troubling. The range was estimated long and the marginal variances were estimated high in the eastern part because this was the “best” way to explain the changes observed, but we do not truly believe the estimates. The long estimated range means that most of the eastern part is highly correlated and the high marginal variance means that next year there might be a large change in the level in the eastern part. Whereas the low marginal variance in the west means that there will be far less changes in the spatial field there the next year. This is clearly wrong since the data for different years do not show huge changes, which
Figure 9: Scatter plots of prediction scores from the stationary and the non-stationary model. 20% of the locations are randomly chosen to be held out and the remaining 80% are used to estimate parameters and predict the 20% held out data. This was repeated 20 times. For values below the line the non-stationary model is better, and conversely for values above the line.

are compatible with the estimated standard deviations of the spatial field, in the level of precipitation between years in the eastern region.

It is well-known that the range and the marginal variance of the stationary Matérn model are not identifiable from a fixed-size observation window \cite{Zhang2004}, and the situation is not likely to improve for a complex model with spatially varying marginal variances and covariance structure, but what we are seeing is the result of forcing the model to include mean structure in the covariance structure. Based on data from multiple years it is clear that the difference in level between the western and eastern region is actually caused by a change in the mean. Further, the short range in the west is also problematic because it means that few of surrounding data points are being used to predict values in this part of the domain. This could mean that the spatial effect is weak in this region, but the estimated covariance structure gives evidence that we need to investigate the cause more thoroughly.

This makes an important point regarding the worth of the non-stationary model. Whether we have improved the CRPS and the log-predictive score is not the only question worth asking. We have gained understanding about issues in the estimated covariance structure that we need to investigate to understand where the non-stationarity is coming from and whether it is correctly captured in the model. In this case we have gained something more than an improvement in prediction scores. We have identified two potential issues with the model: the wrongly specified mean, which we knew about, and the weak spatial effect in the western region, which we need investigate.

4 Non-stationarity in multiple realizations

4.1 Non-stationary modelling framework

If we use multiple realizations, the non-stationarity in the mean and the non-stationarity in the covariance structure are separable if the spatio-temporal process can be assumed
to be stationary in time. Modelling the mean structure and the covariance structure separately goes beyond adaptive smoothing and is a situation where the term non-stationary modelling is accurate. The goal in this section is to separate out the non-stationarity in the mean and to investigate the two issues we discovered in the analysis of a single year in the previous section: over-smoothing in the eastern region and under-smoothing in the western region.

We repeat the analysis using data from the years 1971–1985, and we want to see how much the predictions improve and how the estimated non-stationary changes with a better model for the mean. Ideally, one could fit a full spatio-temporal model to these years, but since the focus is on the spatial non-stationarity we will assume that the 15 years are independent realizations of the same spatial process. Since we are using precipitation data aggregated to yearly data, the temporal dependence is weak and this is a reasonable simplification.

### 4.2 De-trending

The first step in the analysis is to de-trend the dataset. Each year has a different number of observations and some observations are at different locations, which means that there will be different missing locations for each year. The de-trending is done with a simple model that assumes that each year is an independent realization of a stationary spatial field and is observed with measurement noises with the same variance. The model is estimated based on the observations, and the values at locations of interest at each year is filled in based on the posterior marginal conditional means. Then we take the average of the fitted values over the 15 year period as an estimate of the true mean.

The simple model is fitted using the R package INLA, which is based on the INLA method of [Rue et al., 2009]. The model used is

$$y(\mathbf{s}_i, t) = \mu + x(\mathbf{s}_i)\beta + u_t(\mathbf{s}_i) + a_t + \epsilon_{i,t}, \quad i = 1, 2, \ldots, N_t \quad t = 1971, 1972, \ldots, 1985,$$

where $\mu$ is the joint mean for all observations, $x(\mathbf{s}_i)$ is the elevation at location $\mathbf{s}_i$ and $\beta$ is the associated coefficient for the covariate, $u_t$ for $t = 1971, 1972, \ldots, 1985$ are independent realizations of the spatial effect for each year, $a_t$ is an AR(1) process supposed to capture temporal changes in the joint mean between years, and $\epsilon_{i,t}$ are independent Gaussian measurement errors. The spatial effect is approximately Matérn with smoothness parameter $\nu = 1$. The model is estimated and used to predict the values at all locations of interest in all 15 years. The estimate of the true mean $\hat{\mu}(\mathbf{s})$, at location $\mathbf{s}_i$ is found by taking the average over the estimated value at each year.

In the rest of the section we focus on the residuals $y(\mathbf{s}_i, t) - \hat{\mu}(\mathbf{s}_i)$. This means that the estimate of the mean is assumed to be without uncertainty. The intention is to remove most of the non-stationarity in the mean and then evaluate whether there is remaining non-stationarity in the covariance structure of the de-trended data that benefits from being modelled with a non-stationary model. However, if the mean structure is not estimated well with the de-trending procedure, there could be significant non-stationarity left in the mean structure. Any such residual structure will make the estimated covariance structure biased, but the de-trended data for 1981 shown in Figure 10 shows much
less clear evidence for a spatially varying mean than the original data in Figure 1. For example, the de-trending has removed the obvious shift in the level of the precipitation between the western and eastern sides.

4.3 Fitting the non-stationary model

We fit a stationary model (STAT1) and a non-stationary model (NSTAT1) as in Section 3, but without covariates and with the assumption that there are 15 independent replications of the residuals. Each year has observations at potentially different locations, but this does not pose any problems in the SPDE-based model since the entire field is modelled explicitly through the values on each cell in the discretization. The observations are mapped to statements about the values on the grid cells in each year and the inference proceeds in a similar way as for the adaptive smoothing application that used only the year 1981.

The penalty parameters $\tau_1$, $\tau_2$, $\tau_3$ and $\tau_4$ should be changed, but with 15 realizations the cross-validation becomes far more computationally expensive. Therefore, we performed an exploratory analysis where the fits for low, medium and high smoothing were compared, and we decided to use $\log(\tau_1) = 10$, $\log(\tau_2) = 10$, $\log(\tau_3) = 10$ and $\log(\tau_4) = 10$. This might not lead to the highest possible decrease in the prediction scores, but at this point the main interest lies in the qualitative changes in the estimated structure. And, it would, potentially, be a waste of time to put in the required effort before we are certain that there are not major components missing in the model.

The parameters were estimated in the same way as in Section 3, and the maximum penalized likelihood estimates for non-stationarity were used to give the predictions shown in Figure 11. The figure shows both the predictions and the prediction standard deviations for STAT1 and NSTAT1. There are several interesting features in these plots. First, the predicted values are similar for the two models and the main difference is found in the prediction standard deviations. Second, the prediction standard deviations for the western region is troubling for NSTAT1. The range appears to be too short and the
Figure 11: Prediction for de-trended data for year 1981 based on the 15 year period 1971–1985. (a) shows the prediction for STAT1, (b) shows the prediction for NSTAT1, (c) shows the prediction standard deviations for STAT1 and (d) shows the prediction standard deviations for NSTAT1.

spatial effect appears to be close to independent measurement noise in this area. This is not consistent with Figure 10, which appears to have a spatial effect in this region as well.

The problem can be seen clearly when looking at the estimated covariance structure shown in Figure 12. The correlation structure in the eastern part looks regular after de-trending the data, but the correlation structure in the western region is almost degenerating to independent noise. This is a problem from a computational perspective, since the discretization of the SPDE requires that the range is not too small compared to the size of the grid cells, and from a modelling perspective, since the parameters are supposed to describe a slowly changing spatial dependence structure. In the case that the spatial range is that low, the SPDE models requires a high resolution to properly capture the dependence between neighbouring grid cells in the discretization, but if the range is that low, a spatial effect might not be needed. Furthermore, Figure 12(a) shows that the variance of the spatial field is higher in the western region. This indicates that the nugget effect in the western region needs to be different from the nugget effect in the eastern region.

The fits of STAT1 and NSTAT1 are compared with the log-predictive score, the CRPS and the RMSE. The scores are calculated by randomly dividing the data in each year in five parts and then holding out the first part from each year and do the entire fitting and prediction of this data using only the remaining part of the data. Then holding out the second part of the data in each year and so on, for a total of 5 values. This process was then repeated three more times for a total of 20 values of the scores. Scatter plots comparing the scores for the two models are shown in Figure 13.
Figure 12: (a) Estimated marginal standard deviations and (b) estimated 0.7 level contour curves for the correlation functions with respect to the locations marked with red crosses for the spatial effect in NSTAT1.

Figure 13: Scatter plots of (a) Log-predictive score, (b) CRPS and (c) Root mean square error for STAT1 and NSTAT1. The estimates were calculated with hold-out sets where 20% of the locations were held-out from each year as described in Section 4.3. NSTAT1 has a lower log-predictive score and CRPS than STAT1, but the RMSE is higher. The conclusions based on the log-predictive score and the CRPS is the same as for the single realization analysis in Section 3.4. However, the consistently higher RMSE values indicate that there is a problem with the model. The problem lies in the western region where the range is too low, which leads to worse point estimates because the spatial dependence is not exploited. The flexible non-stationary model is able to detect that a higher variance is required for the nugget effect in the western region, but is not able to achieve this in the correct way. Even with all the freedom available in the model it is impossible to have spatial dependence and different nugget effects because we have put the non-stationarity in the wrong components of the model. We need to treat the nugget effects in the western and eastern regions separately.

4.4 Removing the under-smoothing in the western part

The results in Section 4.3 indicate that the nugget effect is different in the western and the eastern part of the conterminous US. Therefore, we fit a stationary model (STAT2)
and a non-stationary model (NSTAT2) with separate nugget effects for locations with longitudes lower than 100°W and for locations with longitudes higher than or equal to 100°W. The placement of the frontier at 100°W is motivated by the change from mountainous regions to plains seen in Figure 2 and the change from low to high range seen in Figure 12(b), but we do not believe it would be particularly sensitive to the exact placement as long as it is in the area of transition from mountainous regions to plains. Except for this change, the models are unchanged, and we use the same penalties $\tau_1$, $\tau_2$, $\tau_3$ and $\tau_4$ for the non-stationarity structure. The intention is to see how much the predictions and the estimated dependence structure change with different nugget effects, but the same penalties.

The predictions and prediction standard deviations are shown in Figure 14. The prediction standard deviations for NSTAT2 do not have the strange artifacts in the western region that are present in Figure 11 for NSTAT1, but one can notice that there is a sharp change in prediction standard deviations at longitude 100°W. This is by construction due to the use of different nugget effects for the two parts of the conterminous US. STAT2 has an estimated standard deviation for the nugget effect of 0.17 in the western part and of 0.083 in the eastern part and for NSTAT2 the estimated standard deviation for the nugget effect is 0.16 in the western part and is 0.083 in the eastern part.

The estimated spatial dependence structure of NSTAT2 is shown in Figure 15. The clearest change from the dependence structure of NSTAT1 shown in Figure 12 is that the non-stationarity in the correlation structure is mostly gone. The appearance is much more reasonable than for NSTAT1 since the entire dependence structure is changing...
slowly and there are no areas with unreasonably large or small ranges. Some non-stationarity still remains in the marginal standard deviations, but together these plots indicate that the simple model STAT2, which does not use a complex non-stationary spatial field, should fit these data well.

We compare the predictions of STAT2 and NSTAT2 by the RMSE, the CRPS and the log-predictive score. The results are given in Figure 16. NSTAT2 performs better according to all of the scores. The scatter plots of the scores show that NSTAT2 performs better for all the hold-out sets, but that the differences in scores are small.

4.5 Discussion of models

The prediction scores for STAT1, NSTAT1, STAT2 and NSTAT2 are shown in Figure 17. The figure shows that the model performing the best according to all scores is NSTAT2, but is the extra computation time worth the effort in this case? The much simpler model STAT2 is performing almost as good as NSTAT2 and requires only one extra parameter. The cost of including one extra parameter is far less than the cost of introducing the
Figure 17: Comparison of STAT1 (blue), NSTAT1 (red), STAT2 (green) and NSTAT2 (black) based on (a) Log-predictive score, (b) CRPS and (c) RMSE. The estimates were calculated with hold-out sets where 20% of the locations were held-out from each year as described in Section 4.3.

flexible non-stationary model. Additionally, one can see that even though the expensive flexible model makes NSTAT1 consistently better than STAT1 in the log-predictive score and the CRPS, STAT2 makes an even greater improvement from STAT1 for the cost of only a single parameter.

The predictions and prediction standard deviations for STAT2 and NSTAT2 in Figure 14 are showing less extreme differences than the predictions and prediction standard deviations for STAT1 and NSTAT1 shown in Figure 11 but there are still some differences in the prediction standard deviations. Some further gain is possible by selecting the penalty parameters controlling the non-stationarity more carefully. We saw some improvement by trying different penalty parameters, but no major changes that would change the conclusion. When we take computation time into account, STAT2 appears to be the better choice. There is some gain with the flexible non-stationary model in NSTAT2, but it comes at a high computational cost.

The physical cause of the difference in the nugget effect between the western region and the eastern region is not known, but it is unlikely to be caused only by differences in the measurement equipment. It is more likely that it is caused by differences in the small-scale behaviour of the process generating the weather in the two different regions that is not captured by the model, but it has not been our intention to find the physical explanation. The intention has been to demonstrate how such a phenomenon can affect the estimation of general flexible models for non-stationarity and the need to carefully evaluate the fitted covariance structures.

5 Discussion

The question of whether we need non-stationary spatial models or not, is a deeper question than it might seem initially. The first step of the analysis should be to decide whether it is likely that non-stationarity is present in the data or not, and in this context simple data exploration, such as variograms, and formal tests (Fuentes, 2005; Jun and Genton, 2012; Bowman and Crujeiras, 2013) are useful tools. The second step is to
decide which non-stationary model we want to use and it can be tempting to look for complex models that allow for spatial fields that have large amounts of flexibility in the covariance structure. We then apply these models with the hope that the high degree of flexibility means that we will be able to capture any non-stationarity present in the data, but the analysis of the annual precipitation data shows that blindly applying such a model might not capture the non-stationarity in the correct and best way.

The case study clearly indicates the need to go beyond simply determining whether or not non-stationarity is present in the data. We need to determine what type of non-stationarity that is present in the data. A flexible model will try to adapt to the non-stationarity, but if the flexibility is available in the wrong parts of the model, the model might have to do suboptimal things to improve the predictive distributions. For example, the model might imitate a spatially varying nugget effect by decreasing the range and varying the marginal variances. This adaptation gives severe undersmoothing, but simply expanding the model with a smoothly varying nugget effect would make the model difficult to identify together with the rest of the flexibility. Therefore, we should determine what is causing the non-stationarity we are seeing before deciding which non-stationary model to use.

The first and most obvious source of non-stationarity in a dataset is the mean structure, and not accounting for this source of non-stationarity will confound the non-stationarity in the mean structure with the non-stationarity in the covariance structure. For example, unmeasured covariates can lead to the apparent long range dependence and global non-stationarity that we observed in the analysis of a single realization. The method presented in this paper is aimed at modelling local non-stationarity and is not appropriate for modelling this type of global non-stationarity. We handle this apparent structure in the covariances by de-trending the data, but it is also possible to model jointly the mean structure and the covariance structure. A simple example of the latter would be to combine the SPDE models with a small number of global basis functions to form a hybrid of fixed-rank kriging and the SPDE models, where the SPDE models captures the short range dependence and local non-stationarity, and the basis functions capture the long range dependence and global non-stationarity. Whichever approach is taken, the paper demonstrates the need to remove the global non-stationarity before modelling the local non-stationarity.

After we have removed the global non-stationarity induced by the mean structure we can model the remaining local non-stationarity, for which the Markovian structure of the SPDE models offers a computationally efficient modelling tool. In the SPDE models we construct a consistent global covariance structure by tying together the local behaviour specified by the SPDE at each location, and the covariance between any two locations will be a combination of the local behaviour at all locations in the model. We believe that this approach is a good way to model local non-stationarity that provides a more flexible, more computationally efficient and easier to parametrize approach than the deformation method, while still having a geometric interpretation of varying the local distance measures.

But modelling local non-stationarity requires information on the small-scale direc-
tional behaviour of the observations, and we would be hesitant to estimate flexible non-stationary models for sparser datasets. Methods such as the deformation method is routinely applied to much sparser datasets, but there is no way around the fact that for patches where we do not have observations we have no idea how the covariances behave. For sparse data it is possible to imagine multiple covariance structures that could give rise to the observed empirical covariances and the unobserved structure must be filled by the model based on the assumptions and restrictions that we have put into the model. This can, potentially, lead to highly model dependent estimates since in non-stationary modelling the missing covariances do not directly affect the observations, and it is important to not allow too much freedom in the covariance structure compared to the sparseness of the data, and to realize that the features seen in the estimated covariance structure will depend on the sparseness of the data.

In an analogous way as for other finite-dimensional methods, there is a confounding of the nugget effect and the resolution chosen for the finite-dimensional approximation. For predictive processes there exists a solution (Finley et al., 2009), but for the SPDE models it is an active field of research. In a GRF model the nugget effect is a combination of the small-scale behaviour and the measurement error, where small-scale behaviour is behaviour below the scale which the data can inform about. The sparser the data is, the more small-scale variation will be confounded with the nugget effect, but for the SPDE models the interpretation of the nugget effect is also tied to the discretization and is a combination of measurement error, small-scale variation and sub-grid variation. The approximation cannot capture variation within the grid cells and these variations increase the nugget variance and decrease the process variances, but this is only a worry when interpreting these parameters. If the precipitation data were sparser, the confounding between small-scale variation and the nugget effect would make it difficult to detect different nugget effects in the western region and the eastern region, and the approach might lead to a different conclusion about the nugget effect.

In each of the three cases studied, the flexible non-stationary model performs better according to the log-predictive score and the CRPS, but when we target directly the non-stationarity in the nugget effect, we can apply a much simpler model just using two nugget effects. Does this mean that the flexible non-stationary model was not useful? No, we were able to use the flexible non-stationary model to estimate a covariance structure that could be used to help determine possible sources of the non-stationarity. We could then include these sources directly and fit a simpler model performing almost equally well, and we could make the same changes to the flexible non-stationary model and fit it again to become confident that there were no other major uncaptured sources of non-stationarity. The idea that the nugget might be the source of heterogeneity is not new (Zimmerman, [1993]), but the case study demonstrates the dangers of putting the heterogeneity in the wrong components in the model.

If there were knowledge available about what was physically generating the non-stationarity, it would be possible to make simpler models where we reduce the flexibility and control the covariance structure by covariates. The use of two nugget effects is an extreme case of this, but covariates in the covariance structure has been a recent direction
of research within all the major families of approaches such as the deformation method, the process convolution method and the SPDE-based method [Schmidt et al., 2011; Neto et al., 2014; Ingebrigtsen et al., 2014]. However, even if we intend to use covariates, the more general non-stationary models could be used to gain intuition about which covariates should be selected and what type of non-stationarity they should control.

The comparison of the different models shows that the scoring rule used to evaluate the predictions has a large influence on the conclusion. The use of a non-stationary model instead of a stationary model mainly affects the prediction variances and not the predicted values. Therefore, the largest improvements are seen in the log-predictive score and the CRPS, and not the RMSE that only evaluates point predictions. However, consistently higher RMSE values for the flexible non-stationary model compared to the simple stationary, as observed when fitting the models using a single nugget effect to de-trended data, is useful to detect problems with the model such as undersmoothing.

The spatial structure of the annual precipitation is treated in detail in the case study, but the temporal structure is not given the same amount of focus. The treatment of the yearly data as independent realizations makes an explicit assumption of independence between years and an implicit assumption of stationarity in time. The assumption of stationarity is necessary to break the fundamental non-separability between the mean structure and the covariance structure, and without this assumption one would only have a single realization of the space-time process. If the dataset cannot be assumed to fulfill these conditions, it is necessary to make a decision about how to distribute the observed behaviour into mean structure and covariance structure through a spatio-temporal model. An interesting point for future work is whether parts of the patterns in the estimated covariance structures are caused by different spatial non-stationarity in different years, and to which degree increasing the spatial resolution would give additional information and to which degree the complexity of the non-stationarity varies with time-scale.

One of the major reasons not to use general non-stationary models unless they are absolutely needed is that they are computationally expensive. The covariate-based approach is less expensive, but requires assumptions about how the non-stationarity varies. Another approach would be to estimate the model locally in different parts of the domain and then try to piece everything together for predictions, but looking for the most efficient way to estimate the model is not the goal of this paper and the more complex one makes the model, the more computationally expensive it will be. The point we are trying to make is that in applications, time might in many cases be better spent on considering how to put the non-stationarity into the model than on developing more complex flexible models and ways to compute them.

Non-stationarity in the covariance structure of spatial models is needed even after the non-stationarity in the mean has been removed, but we need to think carefully about how we handle the non-stationarity. We need to go beyond determining whether there is non-stationarity or not, and determine what type of non-stationarity is present and if possible target this non-stationarity directly instead of using a general flexible model. But in this context the estimated covariance structure from a general flexible model can
in some cases be a useful tool to determine how to do this.

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A Computational details for the model

Let the SPDE

\[(\kappa^2(s) - \nabla \cdot H(s) \nabla) u(s) = \mathcal{W}(s), \quad s \in \mathbb{R}^2, \quad (9)\]

where \(\mathcal{W}(\cdot)\) is standard Gaussian white noise and \(\nabla = (\frac{\partial}{\partial x}, \frac{\partial}{\partial y})\), describe the desired covariance structure. The first thing to notice is that the operator in front of \(u\) only contains multiplications with functions and first order and second order derivatives. All of these operations involve only the local properties of \(u\) at each location. This means that if \(u\) is discretized using a finite-dimensional local basis expansion, the corresponding discretized operators (matrices) should only involve variables close to each other. This can be exploited to create a sparse GMRF which possesses approximately the same covariance structure as \(u\). The arguments above are not applicable for all smoothnesses, but we are constructing a model where the smoothness is fixed to 1 and the range is allowed to vary spatially (See discussion in Fuglstad et al. (2015, p. 5)). A detailed description of the basis function expansion, the choice of mesh, and the theoretical properties of the methods described in this section in Lindgren et al. (2011); Simpson et al. (2012, 2011).

The first step in creating the GMRF is to restrict SPDE (9) to a bounded domain,

\[(\kappa^2(s) - \nabla \cdot H(s) \nabla) u(s) = \mathcal{W}(s), \quad s \in D = [A_1, B_1] \times [A_2, B_2] \subset \mathbb{R}^2, \quad (10)\]

where \(B_1 > A_1\) and \(B_2 > A_2\). This restriction necessitates a boundary condition to make the distribution useful and proper. For technical reasons the boundary condition chosen is zero flux across the boundaries, i.e. at each point of the boundary the flux \(H(s) \nabla u(s)\), where \(\mathbf{n}\) is the normal vector of the boundary at that point, is zero. The derivation of a discretized version of this SPDE on a grid is involved, but for periodic boundary conditions the derivation can be found in the supplementary material to Fuglstad et al. (2015). The boundary conditions in this problem involve only a slight change in that derivation.

For a regular \(m \times n\) grid of \(D\), the end result is the matrix equation

\[
\mathbf{A}(\kappa^2, H)\mathbf{\bar{u}} = \frac{1}{\sqrt{V}}\mathbf{\bar{z}},
\]

where \(V\) is the area of each cell in the grid, \(\mathbf{\bar{u}}\) corresponds to the values of \(u\) on the cells in the regular grid stacked column-wise, \(\mathbf{\bar{z}} \sim \mathcal{N}_{mn}(\mathbf{0}, \mathbf{I}_{mn})\) and \(\mathbf{A}(\kappa^2, H)\) is a discretized version of \((\kappa^2 - \nabla \cdot H \nabla)\). This matrix equation leads to the multivariate Gaussian distribution

\[
\mathbf{\bar{u}} \sim \mathcal{N}_{mn}([\bar{0}, Q(\kappa^2, H)^{-1}]),
\]

where \(Q(\kappa^2, H) = \mathbf{A}(\kappa^2, H)^T \mathbf{A}(\kappa^2, H)V\). The precision matrix \(Q\) is proper and has up to 25 non-zero elements in each row, corresponding to the point itself, its eight closest neighbours and the eight closest neighbours of each of the eight closest neighbours. Since the approximation is constructed from an SPDE, it behaves consistently over different resolution and converges to a continuously indexed model for small resolutions. Changing the resolution changes which features can be represented by the model, but does not induce large changes to the covariance structure.
B Derivation of the second-order random walk prior

Each function, $f$, is a priori modelled as a Gaussian process described by the SPDE

$$-\Delta f(\vec{s}) = \frac{1}{\sqrt{\tau}} W(\vec{s}), \quad \vec{s} \in D = [A_1, B_1] \times [A_2, B_2],$$

(11)

where $A_1 < B_1$, $A_2 < B_2$ and $\tau > 0$, $W$ is standard Gaussian white noise and $\Delta = \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2}$, with the Neumann boundary condition of zero normal derivatives at the edges. In practice this is approximated by representing $f$ as a linear combination of basis elements $\{f_{ij}\}$ weighted by Gaussian distributed weights $\{\alpha_{ij}\}$,

$$f(\vec{s}) = \sum_{i=1}^{K} \sum_{j=1}^{L} \alpha_{ij} f_{ij}(\vec{s}).$$

The basis functions are constructed from separate bases $\{g_i\}$ and $\{h_j\}$ for the $x$-coordinate and the $y$-coordinate, respectively,

$$f_{ij}(\vec{s}) = g_i(x) h_j(y).$$

(12)

For convenience each basis function is assumed to fulfil the boundary condition of zero normal derivative at the edges.

Let $\vec{\alpha} = \text{vec}(\{\alpha_{ij}\}_{ij})$, then the task is to find the best Gaussian distribution for $\vec{\alpha}$. Where “best” is used in the sense of making the resulting distribution for $f$ “close” to a solution of SPDE (11). This is done by a least-squares approach where the vector created from doing inner products of the left hand side with $-\Delta f_{kl}$ must be equal in distribution to the vector created from doing the same to the right hand side,

$$\text{vec} (\langle -\Delta f, -\Delta f_{kl} \rangle_D) \overset{d}{=} \text{vec} (\langle W, -\Delta f_{kl} \rangle_D).$$

(13)

First, calculate the inner product that is needed

$$\langle -\Delta g_i h_j, -\Delta g_k h_l \rangle_D = \langle \Delta g_i h_j, \Delta g_k h_l \rangle_D$$

$$= \left\langle \left( \frac{\partial^2}{\partial x^2} g_i \right) h_j + g_i \frac{\partial^2}{\partial y^2} h_j, \left( \frac{\partial^2}{\partial x^2} g_k \right) h_l + g_k \frac{\partial^2}{\partial y^2} h_l \right\rangle_D.$$  

The bilinearity of the inner product can be used to expand the expression in a sum of four innerproducts. Each of these inner products can then be written as a product of two inner products. Due to lack of space this is not done explicitly, but one of these terms is, for example,

$$\left\langle \left( \frac{\partial^2}{\partial x^2} g_i \right) h_j, \left( \frac{\partial^2}{\partial x^2} g_k \right) h_l \right\rangle_D = \left\langle \frac{\partial^2}{\partial x^2} g_i, \frac{\partial^2}{\partial x^2} g_k \right\rangle_{[A_1, B_1]} \langle h_j, h_l \rangle_{[A_2, B_2]}.$$  

By inserting Equation (12) into Equation (13) and using the above derivations together with integration by parts one can see that the left hand side becomes

$$\text{vec} (\langle -\Delta f, -\Delta f_{kl} \rangle_D) = C\vec{\alpha},$$
\[ C = G_2 \otimes H_0 + 2G_1 \otimes H_1 + G_0 \otimes H_2 \text{ with } \\
G_n = \left[ \frac{\partial^n}{\partial x^n} g_i, \frac{\partial^n}{\partial x^n} g_j \right]_{[A_1,B_1]} \]

and

\[ H_n = \left[ \frac{\partial^n}{\partial y^n} h_i, \frac{\partial^n}{\partial y^n} h_j \right]_{[A_2,B_2]} \].

The right hand side is a Gaussian random vector where the covariance between the position corresponding to \( \alpha_{ij} \) and the position corresponding to \( \alpha_{kl} \) is given by

\[ \langle -\Delta f_{ij}, -\Delta f_{kl} \rangle \mathcal{D}. \]

Thus the covariance matrix of the right hand side must be \( C \) and Equation (13) can be written in matrix form as

\[ C\tilde{\alpha} = C^{1/2}z, \]

where \( \tilde{z} \sim \mathcal{N}_{KL}(\tilde{0}, I_{KL}) \). This means that \( \tilde{\alpha} \) should be given the precision matrix \( Q = C \). Note that \( C \) might be singular due to invariance to some linear combination of the basis elements.

### C Conditional distributions

From the hierarchical model

- Stage 1: \( \tilde{y}|\tilde{z},\tilde{\theta} \sim \mathcal{N}(S\tilde{z}, I_N/\tau_{\text{noise}}) \)
- Stage 2: \( z|\tilde{\theta} \sim \mathcal{N}_{mn+p}(\tilde{0}, Q_c^{-1}) \),

the posterior distribution \( \pi(\tilde{\theta}|\tilde{y}) \) can be derived explicitly. There are three steps involved.

#### C.1 Step 1

Calculate the distribution \( \pi(\tilde{z}|\tilde{\theta}, \tilde{y}) \) up to a constant,

\[ \pi(\tilde{z}|\tilde{\theta}, \tilde{y}) \propto \pi(\tilde{z}|\tilde{\theta})\pi(\tilde{y}|\tilde{\theta}, \tilde{z}) \]

\[ \propto \exp \left( -\frac{1}{2}(\tilde{z} - \tilde{0})^T Q_z (\tilde{z} - \tilde{0}) - \frac{1}{2}(\tilde{y} - S\tilde{z})^T I_N \cdot \tau_{\text{noise}} (\tilde{y} - S\tilde{z}) \right) \]

\[ \propto \exp \left( -\frac{1}{2}(\tilde{z}^T (Q_z + \tau_{\text{noise}} S^T S) \tilde{z} - 2\tilde{z}^T S^T \tilde{y} \cdot \tau_{\text{noise}}) \right) \]

\[ \propto \exp \left( -\frac{1}{2}(\tilde{z} - \mu_C)^T Q_C (\tilde{z} - \mu_C) \right) , \]

where \( Q_C = Q_z + S^T S \cdot \tau_{\text{noise}} \) and \( \mu_C = Q_c^{-1} S^T \tilde{y} \cdot \tau_{\text{noise}} \). This is recognised as a Gaussian distribution

\[ \tilde{z}|\tilde{\theta}, \tilde{y} \sim \mathcal{N}_N(\mu_C, Q_c^{-1}). \]
C.2 Step 2

Integrate out $\vec{z}$ from the joint distribution of $\vec{z}$, $\vec{\theta}$ and $\vec{y}$ via the Bayesian rule,

$$
\pi(\vec{\theta}, \vec{y}) = \frac{\pi(\vec{\theta}, \vec{z}, \vec{y})}{\pi(\vec{z}|\vec{\theta}, \vec{y})} = \frac{\pi(\vec{\theta})\pi(\vec{z}|\vec{\theta})\pi(\vec{y}|\vec{z}, \vec{\theta})}{\pi(\vec{z}|\vec{\theta}, \vec{y})}.
$$

The left hand side of the expression does not depend on the value of $\vec{z}$, therefore the right hand side may be evaluated at any desired value of $\vec{z}$. Evaluating at $\vec{z} = \vec{\mu}_C$ gives

$$
\pi(\vec{\theta}, \vec{y}) \propto \frac{\pi(\vec{\theta})\pi(\vec{z} = \vec{\mu}_C)\pi(\vec{y}|\vec{z} = \vec{\mu}_C, \vec{\theta})}{\pi(\vec{z} = \vec{\mu}_C|\vec{\theta}, \vec{y})} \times \exp \left( -\frac{1}{2} \vec{\mu}_C^T Q_z \vec{\mu}_z \right) \times \exp \left( -\frac{1}{2} (\vec{y} - S \vec{\mu}_C)^T I_N \cdot \tau_{\text{noise}} (\vec{y} - S \vec{\mu}_C) \right) \times \exp \left( +\frac{1}{2} (\vec{\mu}_C - \vec{\mu}_C)^T Q_C (\vec{\mu}_C - \vec{\mu}_C) \right).
$$

C.3 Step 3

Condition on $\vec{y}$ to get the desired conditional distribution,

$$
\log(\pi(\vec{\theta}|\vec{y})) = \text{Const} + \log(\pi(\vec{\theta})) + \frac{1}{2} \log(\det(\mathbf{Q}_z)) + \frac{N}{2} \log(\tau_{\text{noise}}) + \frac{1}{2} \log(\det(\mathbf{Q}_C)) - \frac{1}{2} \vec{\mu}_C^T Q_z \vec{\mu}_z - \frac{\tau_{\text{noise}}}{2} (\vec{y} - S \vec{\mu}_C)^T (\vec{y} - S \vec{\mu}_C). \quad (14)
$$

D Analytic expression for the gradient

This appendix shows the derivation of the derivative of the log-likelihood. Choose the evaluation point $\vec{z} = 0$ in C.2 to find

$$
\log(\pi(\vec{\theta}, \tau_{\text{noise}}|\vec{y})) = \text{Const} + \log(\pi(\vec{\theta}, \tau_{\text{noise}})) + \frac{1}{2} \log(\det(\mathbf{Q}_z)) + \frac{N}{2} \log(\tau_{\text{noise}}) + \frac{1}{2} \log(\det(\mathbf{Q}_C)) - \frac{\tau_{\text{noise}}}{2} (\vec{y} - S \vec{\mu}_C)^T (\vec{y} - S \vec{\mu}_C).
$$

This is just a rewritten form of Equation (14) which is more convenient for the calculation of the gradient, and which separates the $\tau_{\text{noise}}$ parameter from the rest of the covariance parameters. First some preliminary results are presented, then the derivatives are calculated with respect to $\theta_i$ and lastly the derivatives are calculated with respect to $\log(\tau_{\text{noise}})$. 

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Begin with simple preliminary formulas for the derivatives of the conditional precision matrix with respect to each of the parameters,

\[
\frac{\partial}{\partial \theta_i} Q_C = \frac{\partial}{\partial \theta_i} (Q + S^T S \cdot \tau_{\text{noise}}) = \frac{\partial}{\partial \theta_i} Q
\]

and

\[
\frac{\partial}{\partial \log(\tau_{\text{noise}})} Q_C = \frac{\partial}{\partial \log(\tau_{\text{noise}})} (Q + S^T S \cdot \tau_{\text{noise}}) = S^T S \cdot \tau_{\text{noise}}.
\]

### D.1 Derivative with respect to \( \theta_i \)

First the derivatives of the log-determinants can be handled by an explicit formula (Petersen and Pedersen, 2012)

\[
\frac{\partial}{\partial \theta_i} (\log(\det(Q)) - \log(\det(Q_C))) = \text{Tr}(Q^{-1} \frac{\partial}{\partial \theta_i} Q) - \text{Tr}(Q_C^{-1} \frac{\partial}{\partial \theta_i} Q_C)
\]

\[
= \text{Tr} \left[ (Q^{-1} - Q_C^{-1}) \frac{\partial}{\partial \theta_i} Q \right].
\]

Then the derivative of the quadratic forms are calculated

\[
\frac{\partial}{\partial \theta_i} \left( -\frac{1}{2} \bar{y}^T \tau_{\text{noise}} \bar{y} + \frac{1}{2} \bar{\mu}_C Q_C \bar{\mu}_C \right) = 0 + \frac{\partial}{\partial \theta_i} \left( \frac{1}{2} \bar{y}^T \tau_{\text{noise}} S Q_C^{-1} S^T \tau_{\text{noise}} \bar{y} \right)
\]

\[
= \frac{1}{2} \bar{\mu}_C^T \left( \frac{\partial}{\partial \theta_i} Q \right) Q_C^{-1} S^T \tau_{\text{noise}} \bar{y}
\]

\[
= -\frac{1}{2} \bar{\mu}_C \left( \frac{\partial}{\partial \theta_i} Q \right) \bar{\mu}_C.
\]

Combining these gives

\[
\frac{\partial}{\partial \theta_i} \log(\pi(\bar{\theta}, \tau_{\text{noise}} | \bar{y})) = \frac{\partial}{\partial \theta_i} \log(\pi(\bar{\theta}, \tau_{\text{noise}})) + \frac{1}{2} \text{Tr} \left[ (Q^{-1} - Q_C^{-1}) \frac{\partial}{\partial \theta_i} Q \right] - \frac{1}{2} \bar{\mu}_C \left( \frac{\partial}{\partial \theta_i} Q \right) \bar{\mu}_C
\]

### D.2 Derivative with respect to \( \log(\tau_{\text{noise}}) \)

First calculate the derivative of the log-determinants

\[
\frac{\partial}{\partial \log(\tau_{\text{noise}})} (N \log(\tau_{\text{noise}}) - \log(\det(Q_C))) = N - \text{Tr} \left( Q_C^{-1} \frac{\partial}{\partial \log(\tau_{\text{noise}})} Q_C \right)
\]

\[
= N - \text{Tr} \left( Q_C^{-1} S^T S \cdot \tau_{\text{noise}} \right).
\]
Then the derivative of the quadratic forms

$$\frac{\partial}{\partial \log(\tau_{\text{noise}})} \left( -\frac{1}{2} y^T \overrightarrow{\tau}_{\text{noise}} + \frac{1}{2} \overrightarrow{\mu}_C Q C \overrightarrow{\mu}_C \right) = -\frac{1}{2} y^T \overrightarrow{\tau}_{\text{noise}} + \frac{\partial}{\partial \log(\tau_n)} \frac{1}{2} y^T \overrightarrow{\tau}_{\text{noise}} S Q C^{-1} S^T \overrightarrow{\tau}_{\text{noise}} y +$$

$$= -\frac{1}{2} y^T \overrightarrow{\tau}_{\text{noise}} + y^T \overrightarrow{\tau}_{\text{noise}} S Q C^{-1} S \left( \frac{\partial \overrightarrow{\tau}_{\text{noise}}}{\partial \log(\tau_{\text{noise}})} \right) \overrightarrow{y} +$$

$$-\frac{1}{2} y^T \overrightarrow{\tau}_{\text{noise}} S Q C^{-1} \left( \frac{\partial}{\partial \log(\tau_{\text{noise}})} Q C \right) Q C^{-1} S^T \overrightarrow{\tau}_{\text{noise}} y$$

$$= -\frac{1}{2} y^T \overrightarrow{\tau}_{\text{noise}} + \vec{\mu}_C S^T \overrightarrow{y} \cdot \tau_{\text{noise}} - \frac{1}{2} \vec{\mu}_C S^T S \vec{\mu}_C \cdot \tau_{\text{noise}}$$

$$= -\frac{1}{2} (\overrightarrow{y} - A \vec{\mu}_C)^T (\overrightarrow{y} - A \vec{\mu}_C) \cdot \tau_{\text{noise}}.$$

Together these expressions give

$$\frac{\partial \log(\pi(\overrightarrow{\theta}, \tau_{\text{noise}} | \overrightarrow{y}))}{\partial \log(\tau_{\text{noise}})} = \frac{\partial}{\partial \log(\tau_{\text{noise}})} \log(\pi(\overrightarrow{\theta}, \tau_{\text{noise}})) + \frac{N}{2} - \frac{1}{2} \text{Tr} \left[ Q C^{-1} S^T S \cdot \tau_{\text{noise}} \right] +$$

$$\frac{1}{2} (\overrightarrow{y} - A \vec{\mu}_C)^T (\overrightarrow{y} - A \vec{\mu}_C) \cdot \tau_{\text{noise}}.$$

### D.3 Implementation

The derivative $\frac{\partial}{\partial \theta} Q_c$ can be calculated quickly since it is a simple functions of $\theta$. The trace of the inverse of a matrix $A$ times the derivative of a matrix $B$ only requires the values of the inverse of $A$ for non-zero elements of $B$. In the above case the two matrices have the same type of non-zero structure, but it can happen that specific elements in the non-zero structure are zero for one of the matrices. This way of calculating the inverse only at a subset of the locations can be handled as described in Rue and Held (2010).

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