# Bayesian Ridge Regression - Final Report

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# Introduction

# 1 Mathematical Model

This chapter focuses on the underlying theoretical model of Bayesian Ridge Regression in the context of the Gaussian Location-Scale Regression model considered in the lmls package. First, the distributional assumptions for the parameter vectors  $\boldsymbol{\beta}$  and  $\boldsymbol{\gamma}$  as well as the scalar quantities  $\tau^2$  and  $\xi^2$  are clearly stated. Based on these prior distributions, the full conditional distributions of each parameter given all of the remaining model components are derived. Throughout this rather theoretical exposition we will build connections from the derived equations to practical consequences that have to be considered in the code implementation discussed in sections 2.2.1 and 2.2.2.

### 1.1 Prior Distributions

Assuming conditional independence among the regression coefficients  $\beta$  and  $\gamma$  as well as flat priors for the intercept parameters

$$f(\beta_0) \propto const$$
 and  $f(\gamma_0) \propto const$ ,

first note that

$$f(\boldsymbol{\beta} \mid \tau^2) = f(\beta_0) f(\tilde{\boldsymbol{\beta}} \mid \tau^2) \propto f(\tilde{\boldsymbol{\beta}} \mid \tau^2) \quad \text{and} \quad f(\boldsymbol{\gamma} \mid \xi^2) = f(\gamma_0) f(\tilde{\boldsymbol{\gamma}} \mid \xi^2) \propto f(\tilde{\boldsymbol{\gamma}} \mid \xi^2),$$

where the notation  $\boldsymbol{\beta} = (\beta_0, ..., \beta_K) \in \mathbb{R}^{K+1}$ ,  $\tilde{\boldsymbol{\beta}} = (\beta_1, ..., \beta_K) \in \mathbb{R}^K$  and, analogously,  $\boldsymbol{\gamma} = (\gamma_0, ..., \gamma_J) \in \mathbb{R}^{J+1}$ ,  $\tilde{\boldsymbol{\gamma}} = (\gamma_1, ..., \gamma_J) \in \mathbb{R}^J$  is used.

Thus, the Prior distributions of the parameters in the Bayesian Ridge Regression model, which are responsible for the regularizing effect compared to models without penalty, are given by

- $\tilde{\boldsymbol{\beta}} \mid \tau^2 \sim \mathcal{N}\left(\mathbf{0}, \, \tau^2 \cdot \mathbf{I}_K\right)$ ,
- $\tilde{\gamma} \mid \xi^2 \sim \mathcal{N}\left(\mathbf{0}, \, \xi^2 \cdot \mathbf{I}_J\right)$ ,
- $\tau^2 \sim IG(a_{\tau}, b_{\tau})$ , with fixed hyperparameters  $a_{\tau}$  and  $b_{\tau}$ ,
- $\xi^2 \sim IG(a_{\xi}, b_{\xi})$ , with fixed hyperparameters  $a_{\xi}$  and  $b_{\xi}$ .

### 1.2 Full Posterior Distribution

The starting point for deriving the Full Conditional distributions, which will majorly impact the implementation of the Metropolis-Hastings sampling process, is always the Full Posterior distribution. As usual in the Bayesian literature, we write the Full Posterior  $f(\beta, \gamma, \tau^2, \xi^2 \mid \mathbf{y})$  in terms of the Likelihood function / observation model  $f(\mathbf{y} \mid \beta, \gamma, \tau^2, \xi^2)$  and the *joint* Prior distribution  $f(\beta, \gamma, \tau^2, \xi^2)$ , i.e.

$$f(\boldsymbol{\beta}, \boldsymbol{\gamma}, \tau^2, \boldsymbol{\xi}^2 \mid \mathbf{y}) \propto f(\mathbf{y} \mid \boldsymbol{\beta}, \boldsymbol{\gamma}, \tau^2, \boldsymbol{\xi}^2) \cdot f(\boldsymbol{\beta}, \boldsymbol{\gamma}, \tau^2, \boldsymbol{\xi}^2).$$

The specific form of the Likelihood function is given by the Location-Scale Regression model that the lmls package is built upon:

$$y_i \mid \boldsymbol{\beta}, \boldsymbol{\gamma}, \tau^2, \xi^2 = y_i \mid \boldsymbol{\beta}, \boldsymbol{\gamma} \sim \mathcal{N}\left(\mathbf{x}_i^T \boldsymbol{\beta}, \exp\left(\mathbf{z}_i^T \boldsymbol{\gamma}\right)^2\right).$$

Taking the independence structure into account, the joint Prior distribution can be written as

$$f(\beta, \gamma, \tau^{2}, \xi^{2}) = f(\beta \mid \gamma, \tau^{2}, \xi^{2}) \cdot f(\gamma, \tau^{2}, \xi^{2})$$

$$= f(\beta \mid \gamma, \tau^{2}, \xi^{2}) \cdot f(\gamma \mid \tau^{2}, \xi^{2}) \cdot f(\tau^{2}, \xi^{2})$$

$$= f(\beta \mid \tau^{2}) \cdot f(\gamma \mid \xi^{2}) \cdot f(\tau^{2}) \cdot f(\xi^{2}).$$

Combining these results yields for the Full Posterior distribution the general form

$$f(\boldsymbol{\beta}, \boldsymbol{\gamma}, \tau^2, \xi^2 \mid \mathbf{y}) \propto f(\mathbf{y} \mid \boldsymbol{\beta}, \boldsymbol{\gamma}, \tau^2, \xi^2) \cdot f(\boldsymbol{\beta} \mid \tau^2) \cdot f(\boldsymbol{\gamma} \mid \xi^2) \cdot f(\tau^2) \cdot f(\xi^2)$$
$$\propto f(\mathbf{y} \mid \boldsymbol{\beta}, \boldsymbol{\gamma}) \cdot f(\tilde{\boldsymbol{\beta}} \mid \tau^2) \cdot f(\tilde{\boldsymbol{\gamma}} \mid \xi^2) \cdot f(\tau^2) \cdot f(\xi^2),$$

in which the corresponding densities for the observation model and the individual prior distributions can be inserted. These are given by

• 
$$f(\mathbf{y} \mid \boldsymbol{\beta}, \boldsymbol{\gamma}) = \prod_{i=1}^{n} \frac{1}{\sqrt{2\pi \exp(\mathbf{z}_{i}^{T} \boldsymbol{\gamma})^{2}}} \cdot \exp\left(-\frac{1}{2 \exp(\mathbf{z}_{i}^{T} \boldsymbol{\gamma})^{2}} \cdot (y_{i} - \mathbf{x}_{i}^{T} \boldsymbol{\beta})^{2}\right),$$

• 
$$f(\boldsymbol{\beta} \mid \tau^2) \propto \prod_{k=1}^K \frac{1}{\sqrt{2\pi\tau^2}} \cdot \exp\left(-\frac{1}{2\tau^2} \cdot \beta_k^2\right) = (2\pi)^{-\frac{K}{2}} \tau^{-K} \exp\left(-\frac{1}{2\tau^2} \cdot \tilde{\boldsymbol{\beta}}^T \tilde{\boldsymbol{\beta}}\right)$$

• 
$$f(\gamma \mid \xi^2) \propto \prod_{j=1}^J \frac{1}{\sqrt{2\pi\xi^2}} \cdot \exp\left(-\frac{1}{2\xi^2} \cdot \gamma_j^2\right) = (2\pi)^{-\frac{J}{2}} \xi^{-J} \exp\left(-\frac{1}{2\xi^2} \cdot \tilde{\gamma}^T \tilde{\gamma}\right)$$

• 
$$f(\tau^2) = \frac{b_{\tau}}{\Gamma(a_{\tau})} \left(\frac{1}{\tau^2}\right)^{a_{\tau}+1} \exp\left(-\frac{b_{\tau}}{\tau^2}\right)$$
,

• 
$$f(\xi^2) = \frac{b_{\xi}}{\Gamma(a_{\xi})} \left(\frac{1}{\xi^2}\right)^{a_{\xi}+1} \exp\left(-\frac{b_{\xi}}{\xi^2}\right)$$

leading to the Full Posterior

$$f(\boldsymbol{\beta}, \boldsymbol{\gamma}, \tau^{2}, \xi^{2} \mid \mathbf{y}) \propto \prod_{i=1}^{n} \frac{1}{\exp(\mathbf{z}_{i}^{T} \boldsymbol{\gamma})} \cdot \exp\left(-\frac{1}{2 \exp(\mathbf{z}_{i}^{T} \boldsymbol{\gamma})^{2}} \cdot (y_{i} - \mathbf{x}_{i}^{T} \boldsymbol{\beta})^{2}\right)$$
$$\cdot \tau^{-K} \exp\left(-\frac{1}{2\tau^{2}} \tilde{\boldsymbol{\beta}}^{T} \tilde{\boldsymbol{\beta}}\right) \cdot \xi^{-J} \exp\left(-\frac{1}{2\xi^{2}} \tilde{\boldsymbol{\gamma}}^{T} \tilde{\boldsymbol{\gamma}}\right)$$
$$\cdot \left(\frac{1}{\tau^{2}}\right)^{a_{\tau}+1} \exp\left(-\frac{b_{\tau}}{\tau^{2}}\right) \cdot \left(\frac{1}{\xi^{2}}\right)^{a_{\xi}+1} \exp\left(-\frac{b_{\xi}}{\xi^{2}}\right).$$

### 1.3 Full Conditional Distributions

The Full Posterior distribution contains complete information about the statistical model. For our purposes, we are mostly interested in the Full Conditional Distribution of each model parameter. These can be obtained by simply neglecting all factors of the Full Posterior that do not depend on the parameter in consideration.

The Full Conditional distribution can then be recovered by the resulting density kernel, either by recognizing a known distribution or by adding a normalization constant (which, however, is not needed for Markov Chain Monte Carlo sampling).

# 1.3.1 Full Conditional of $\tau^2$ :

$$f(\tau^{2} \mid \cdot) \propto \tau^{-K} \cdot \exp\left(-\frac{1}{2\tau^{2}} \cdot \tilde{\boldsymbol{\beta}}^{T} \tilde{\boldsymbol{\beta}}\right) \cdot \left(\frac{1}{\tau^{2}}\right)^{a_{\tau}+1} \exp\left(-\frac{b_{\tau}}{\tau^{2}}\right)$$
$$\propto \left(\frac{1}{\tau^{2}}\right)^{a_{\tau}+\frac{K}{2}+1} \cdot \exp\left(-\frac{1}{\tau^{2}} \left(b_{\tau} + \frac{1}{2} \tilde{\boldsymbol{\beta}}^{T} \tilde{\boldsymbol{\beta}}\right)\right).$$

This is the kernel of an Inverse Gamma distribution parameterized by

$$\tau^2 \mid \cdot \sim IG(a_{\tau} + \frac{K}{2}, b_{\tau} + \frac{1}{2}\tilde{\boldsymbol{\beta}}^T\tilde{\boldsymbol{\beta}}).$$

### 1.3.2 Full Conditional of $\xi^2$ :

$$f(\xi^{2} \mid \cdot) \propto \xi^{-J} \cdot \exp\left(-\frac{1}{2\xi^{2}} \cdot \tilde{\gamma}^{T} \tilde{\gamma}\right) \cdot \left(\frac{1}{\xi^{2}}\right)^{a_{\xi}+1} \exp\left(-\frac{b_{\xi}}{\xi^{2}}\right)$$
$$\propto \left(\frac{1}{\xi^{2}}\right)^{a_{\xi}+\frac{J}{2}+1} \cdot \exp\left(-\frac{1}{\xi^{2}} \left(b_{\xi} + \frac{1}{2} \tilde{\gamma}^{T} \tilde{\gamma}\right)\right).$$

Thus, the Full Conditional of  $\xi^2$  follows an Inverse Gamma distribution as well:

$$\xi^2 \mid \cdot \sim IG(a_{\xi} + \frac{J}{2}, b_{\xi} + \frac{1}{2}\tilde{\gamma}^T\tilde{\gamma}).$$

### 1.3.3 Full Conditional of $\beta$ :

Here, the derivation is more involved. In order to keep the calculations structured, we introduce the following notation:

$$\mathbf{w}_i := \frac{\mathbf{x}_i}{\exp\left(\mathbf{z}_i^T \boldsymbol{\gamma}\right)} \in \mathbb{R}^{K+1}, \qquad \mathbf{W} := \begin{pmatrix} \mathbf{w}_1^T \\ \vdots \\ \mathbf{w}_n^T \end{pmatrix} \in \mathbb{R}^{n \times (K+1)}$$

and

$$u_i := \frac{y_i}{\exp\left(\mathbf{z}_i^T \boldsymbol{\gamma}\right)} \in \mathbb{R}, \quad \mathbf{u} := \begin{pmatrix} u_1 \\ \vdots \\ u_n \end{pmatrix} \in \mathbb{R}^n,$$

yielding  $\sum_{i=1}^{n} u_i \mathbf{w}_i = \mathbf{W}^T \mathbf{u}$  and  $\sum_{i=1}^{n} \mathbf{w}_i \mathbf{w}_i^T = \mathbf{W}^T \mathbf{W}$ .

Therefore the Full Conditional distribution of  $\beta$  can be written as

$$f(\beta \mid \cdot) \propto \exp\left(-\frac{1}{2} \cdot \left[\frac{1}{\tau^{2}} \tilde{\boldsymbol{\beta}}^{T} \tilde{\boldsymbol{\beta}} + \sum_{i=1}^{n} \frac{1}{\exp\left(\mathbf{z}_{i}^{T} \boldsymbol{\gamma}\right)^{2}} \left(y_{i} - \mathbf{x}_{i}^{T} \boldsymbol{\beta}\right)^{2}\right]\right)$$

$$= \exp\left(-\frac{1}{2} \cdot \left[\frac{1}{\tau^{2}} \tilde{\boldsymbol{\beta}}^{T} \tilde{\boldsymbol{\beta}} + \sum_{i=1}^{n} \left(\frac{y_{i}^{2}}{\exp\left(\mathbf{z}_{i}^{T} \boldsymbol{\gamma}\right)^{2}} - \frac{2y_{i} \mathbf{x}_{i}^{T}}{\exp\left(\mathbf{z}_{i}^{T} \boldsymbol{\gamma}\right)^{2}} \boldsymbol{\beta} + \boldsymbol{\beta}^{T} \frac{\mathbf{x}_{i}}{\exp\left(\mathbf{z}_{i}^{T} \boldsymbol{\gamma}\right)} \frac{\mathbf{x}_{i}^{T}}{\exp\left(\mathbf{z}_{i}^{T} \boldsymbol{\gamma}\right)} \boldsymbol{\beta}\right)\right]\right)$$

$$\propto \exp\left(-\frac{1}{2} \cdot \left[\frac{1}{\tau^{2}} \tilde{\boldsymbol{\beta}}^{T} \tilde{\boldsymbol{\beta}} - 2 \cdot \sum_{i=1}^{n} u_{i} \mathbf{w}_{i}^{T} \boldsymbol{\beta} + \sum_{i=1}^{n} \boldsymbol{\beta}^{T} \mathbf{w}_{i} \mathbf{w}_{i}^{T} \boldsymbol{\beta}\right]\right)$$

$$= \exp\left(-\frac{1}{2} \cdot \left[\boldsymbol{\beta}^{T} \left(\sum_{i=1}^{n} \mathbf{w}_{i} \mathbf{w}_{i}^{T} + \frac{1}{\tau^{2}} \begin{pmatrix} 0 & 0 \\ 0 & \mathbf{I}_{K} \end{pmatrix} \right) \boldsymbol{\beta} - 2 \cdot \sum_{i=1}^{n} \boldsymbol{\beta}^{T} u_{i} \mathbf{w}_{i}\right]\right)$$

$$= \exp\left(-\frac{1}{2} \cdot \left[\boldsymbol{\beta}^{T} \left(\mathbf{W}^{T} \mathbf{W} + \frac{1}{\tau^{2}} \begin{pmatrix} 0 & 0 \\ 0 & \mathbf{I}_{K} \end{pmatrix} \right) \boldsymbol{\beta} - 2 \cdot \boldsymbol{\beta}^{T} \mathbf{W}^{T} \mathbf{u}\right]\right).$$

Comparing this representation with the kernel of a multivariate normal distribution leads to the conclusion

$$oldsymbol{eta} \mid \cdot \sim \mathcal{N}_{K+1} \left(oldsymbol{\mu}_{eta}, oldsymbol{\Sigma}_{eta}
ight)$$

with the parameters

$$\mathbf{\Sigma}_{eta} = \left(\mathbf{W}^T\mathbf{W} + rac{1}{ au^2} \left(egin{array}{cc} 0 & 0 \ 0 & \mathbf{I}_K \end{array}
ight)
ight)^{-1} \qquad ext{and} \qquad oldsymbol{\mu}_{eta} = oldsymbol{\Sigma}_{eta} \mathbf{W}^T \mathbf{u}.$$

### 1.3.4 Full Conditional of $\gamma$ :

Using the notation

$$\mathbf{z}_i \in \mathbb{R}^{J+1}, \qquad \mathbf{Z} = egin{pmatrix} \mathbf{z}_1^T \ dots \ \mathbf{z}_n^T \end{pmatrix} \in \mathbb{R}^{n \times (J+1)} \qquad ext{and} \qquad oldsymbol{\gamma} \in \mathbb{R}^{J+1},$$

the Full Conditional distribution of  $\gamma$  is given by

$$f(\boldsymbol{\gamma} \mid \cdot) \propto \exp\left(-\frac{1}{2} \cdot \left[\frac{1}{\xi^2} \tilde{\boldsymbol{\gamma}}^T \tilde{\boldsymbol{\gamma}} + \sum_{i=1}^n \left(\frac{1}{\exp\left(\mathbf{z}_i^T \boldsymbol{\gamma}\right)^2} \left(y_i - \mathbf{x}_i^T \boldsymbol{\beta}\right)^2 + 2 \cdot \mathbf{z}_i^T \boldsymbol{\gamma}\right)\right]\right)$$
$$= \exp\left(-\frac{1}{2} \cdot \left[\frac{1}{\xi^2} \tilde{\boldsymbol{\gamma}}^T \tilde{\boldsymbol{\gamma}} + 2 \cdot \mathbf{1}_n^T \mathbf{Z} \boldsymbol{\gamma} + \sum_{i=1}^n \left(\frac{y_i - \mathbf{x}_i^T \boldsymbol{\beta}}{\exp\left(\mathbf{z}_i^T \boldsymbol{\gamma}\right)}\right)^2\right]\right).$$

In contrast to the Full Conditionals for  $\beta$ ,  $\tau^2$  and  $\xi^2$ , this kernel cannot be assigned to a known distribution. Thus, for sampling from the Full Posterior distribution, it is not feasible to use a Gibbs Sampler in its purest form. More specifically, we will include a Metropolis Hastings step for sampling the  $\gamma$  parameter vector.

Although this 'inconvenience' is not required for  $\beta$  (since we can use independent samples from a multivariate normal distribution), we will briefly explore and analyze the convergence properties of the modified version of sampling both  $\gamma$  and  $\beta$  via the Metropolis-Hastings procedure.

# 2 The asp21bridge Package

This chapter introduces numerous facets of the asp21bridge package.

Section 2.1 explains how to use and combine the functions that are contained in the package most efficiently.

Section 2.2 focuses on the implementation of the Markov Chain Monte Carlo Sampler with Ridge Penalty, links the source code to the mathematical model from chapter 1 and explains how the main function mcmc\_ridge() as well as the helper functions implementing the Metropolis-Hastings algorithm are structured.

Finally, some additional components of the package development process and ideas, that the package is based upon, are discussed in section 2.3.

### 2.1 User Guide

This section aims to provide guidance for new users of the asp21bridge package. Although all exported functions are fully documented such that function arguments and brief examples can be looked up at the corresponding help page, the following tutorial extends the documentation by illustrating a typical workflow of simulation via the penalized MCMC Sampler, extracting meaningful statistical quantities from the samples and visually analyzing the results.

The asp21bridge package inherits all functions from the lmls package and exports 9 additional functions, which can be grouped into three categories:

- Sampling: The whole sampling process is covered by the very flexible and robust mcmc\_ridge() function that is explained in detail in section 2.2.1.
- Graphical Analysis: The trace\_plot(), density\_plot() and acf\_plot() functions provide the three most common visualizations of a *single* chain's development over time, its distribution and the autocorrelation between the samples. Since all of these are *diagnostic* tools, they are combined in the high-level diagnostic\_plots() function, which simply collects all three plots in a grid and will be used more often than the separate building blocks.
  - For visualizing *multiple* chains together, the mult\_plot() function can be used. Several arguments for customizing the graphical output exist. In the most basic form, trace plots of all selected chains are displayed in the upper panel while the corresponding density plots are integrated into the lower panel.
- Statistical Analysis: Here, the summary\_complete() function represents the main tool and provides a more thorough statistical summary than the generic summary() function. In addition, the burnin() and thinning() functions are convenient in the context of Markov Chains and in particular for extracting more meaningful features of the samples from the posterior distributions.

### 2.1.1 Sampling with the mcmc\_ridge() function

As explained in section 2.2.1, there are two valid input types for simulating with the mcmc\_ridge() function. Most commonly, the sampling procedure will be built upon an existing model that was initialized by the lmls() function. In this case, only the name of the model object is required, although many further arguments can be specified such as the number of simulations nsim which is set to 1000 by default.

The mcmc\_ridge() function can, however, be used completely independent from the underlying lmls package. Therefore, the outcome vector  $\mathbf{y}$  as well as the design matrices  $\mathbf{X}$  and  $\mathbf{Z}$ , corresponding to the  $\boldsymbol{\beta}$  and  $\boldsymbol{\gamma}$  coefficient vectors respectively (as explained in chapter 1), must be manually specified.

In order to illustrate both options, we construct two separate models:

• The first model uses the built-in toy\_data, a data set which is specifically designed for introductory tutorials, documentation and unit tests. It consists of a column y representing a vector of observed values and the explanatory variables x1, x2, z1 and z2.

The data is simulated according to the correctly specified location-scale regression model from chapter 1, where all explanatory variables predict the mean of  $\mathbf{y}$  and only the latter two model the variance. This example will be used for model evaluation, since the true data generating values  $\boldsymbol{\beta} = \begin{pmatrix} 0 & -2 & -1 & 1 & 2 \end{pmatrix}^T$  and  $\boldsymbol{\gamma} = \begin{pmatrix} 0 & -1 & 1 \end{pmatrix}^T$  are known.

For the simulations, we use the model object that is created by the lmls() function as input and use most of the mcmc\_ridge() default settings, except for the number of simulations, which we increase to 10000.

• The second model uses the more realistic abdom data set from the lmls package.

In this case, we have to provide the data input as well as starting values for  $\beta$  and  $\gamma$  explicitly. Note that the dimension of beta\_start and gamma\_start have to match the number of columns in X and Z in the model *input*. If necessary, the mcmc\_ridge() functions adds intercept columns to both design matrices, such that the dimension of the coefficient vectors might have increased (as in the case illustrated here) in the model *output*.

This example differs from the first one with regards to the conclusions that can be drawn: Since the true data generating values of  $\beta = \begin{pmatrix} \beta_0 & \beta_1 \end{pmatrix}^T$  and  $\gamma = \begin{pmatrix} \gamma_0 & \gamma_1 \end{pmatrix}^T$  are *not* known, we have to rely more heavily on the model estimates. The number of simulations is again increased to 10000 such that statistically valid estimates of posterior characteristics are possible, even if preceding 'burnin' and 'thinning' steps might be required.

Note, that we first standardize the covariates in this case, which is strongly recommended when working with penalized methods.

```
library(asp21bridge)
set.seed(1234)

toy_fit <- lmls(
    location = y ~ x1 + x2 + z1 + z2, scale = ~ z1 + z2,
    data = toy_data, light = FALSE
) %>%
    mcmc_ridge(num_sim = 10000)

standardize <- function(x) (x - mean(x)) / sd(x)
y <- abdom$y
X <- as.matrix(standardize(abdom$x))
Z <- X

abdom_fit <- mcmc_ridge(
    y = y, X = X, Z = Z, beta_start = 1, gamma_start = 1,
    num_sim = 10000
)</pre>
```

The different forms of data input cause different structures in the resulting model objects: toy\_fit inherits the model structure as well as the S3 Class lmls from the lmls() function and adds a list entry mcmc\_ridge containing the sampling results.

In contrast, abdom\_fit only contains matrices filled with the simulations and the acceptance probability of the Metropolis-Hastings step for sampling  $\gamma$ :

```
str(abdom_fit, max.level = 1)
## List of 2
## $ sampling_matrices:List of 4
## $ acceptance_rate : num 0.305
```

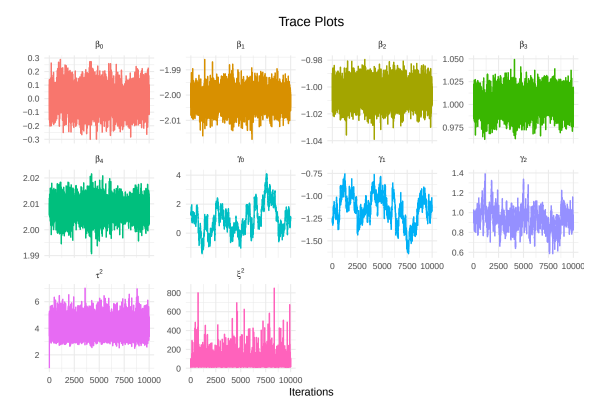


Figure 1: Trace Plots for all Coefficients, data: toy data

### 2.1.2 Graphical Analysis

Most statistical analyses start with an exploratory phase. To obtain a graphical overview of the simulations, the mult\_plot() function is convenient to display trace plots and/or density plots for all model coefficients.

The free\_scale argument is often useful to obtain a meaningful graphical output, if the parameters are on different numerical scales. Setting latex = TRUE transforms the coefficient names to their corresponding greek symbols, which, although being a purely aesthetic feature, required a surprisingly nontrivial implementation.

```
mult_plot(samples = toy_fit, type = "trace", free_scale = TRUE, latex = TRUE)
```

Figure 1 shows trace plots for all coefficients of the toy\_fit model. Due to the high number of simulations, the plot facets appear a bit overloaded. Considering the y-axis scales, the samples for the  $\beta$  vector show a small variance and fast convergence, whereas the samples for  $\gamma_0$  and  $\gamma_1$  indicate a significant autocorrelation and no sign of convergence.

The log argument can be useful for displaying variance parameters such as  $\tau^2$  and  $\xi^2$ , which are strictly positive. Figure 2 illustrates the effect of this option on both, the y-axis of the trace plots as well as the x-axis of the density plots.

```
variance_samples <- cbind(
  abdom_fit$sampling_matrices$tau_samples,
  abdom_fit$sampling_matrices$xi_samples
)

mult_plot(
  samples = variance_samples, type = "both", free_scale = TRUE,
  log = TRUE, latex = TRUE
)</pre>
```

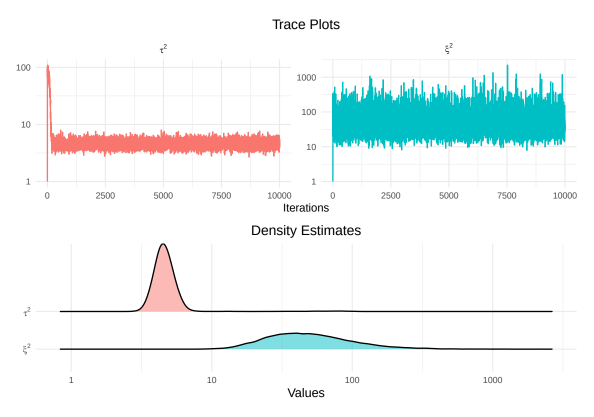


Figure 2: Trace and Density Plots for the Prior Variances, data: abdom

Note that the mult\_plot() function accepts various kinds of input data, such as a complete lmls model in the first case or simply one or multiple sampling matrices in the latter case, and correctly extracts the corresponding samples.

To focus on a single Markov chain, the diagnostic\_plots() function is useful:

```
diagnostic_plots(
   samples = abdom_fit$sampling_matrices$beta_samples[, "beta_1", drop = FALSE],
   lag_max = 100, latex = TRUE
)
```

Figure 3 clearly shows the need for a burnin and thinning step for  $\beta_1$ , since the chain varies heavily among the first iterations and the samples are strongly autocorrelated. The lag\_max argument controls the number of lags that are included in the autocorrelation plot.

In order to confront these issues, we remove the first 500 samples and additionally only keep every 10th iteration:

```
abdom_fit$sampling_matrices$beta_samples[, "beta_1", drop = FALSE] %>%
burnin(num_burn = 500) %>%
thinning(freq = 10) %>%
diagnostic_plots(lag_max = 100, latex = TRUE)
```

The diagnostic plots for  $\beta_1$  in Figure 4 look much better. The chain has converged and there is little residual autocorrelation left after thinning out the samples. Additionally, the posterior density approximately follows a normal distribution.

Although removing posterior estimates in this way (and arguably losing valuable information) is not uncontroversial in the Bayesian community, statistical estimates from the remaining, well behaved chain are usually

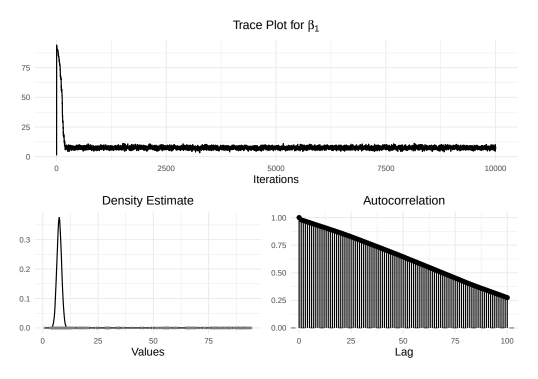


Figure 3: Diagnostic Plots for  $\beta_1$  before thinning, data: abdom

more stable and reliable. In any case, the above procedure clearly emphasizes the usefulness of a graphical diagnosis of the sampling results as a preliminary step before drawing any far reaching conclusions.

## 2.1.3 Statistical Analysis

At the end of a Bayesian statistical analysis, summary statistics of the posterior distribution are of major interest. These include estimates for *centrality*, such as the Posterior Mean or the Posterior Median, estimates for the *spread*, e.g. the Posterior Variance, and quantile estimates in the tails of the posterior distribution as bounds for credible intervals.

The asp21bridge package offers two options to quickly access this information: If only a quick look at the numerical results without further investigation is desired, the summary() function can be used. The lmls package adapts this generic S3 method to the lmls class with an additional type argument. Specifying mcmc\_ridge displays the estimates of the mcmc\_ridge() function. Note, that this option is only applicable for the toy\_fit model, which is based on the lslm class:

```
summary(toy_fit, type = "mcmc_ridge")
##
## Call:
  lmls(location = y \sim x1 + x2 + z1 + z2, scale = \sim z1 + z2, data = toy_data,
##
##
       light = FALSE)
##
## Pearson residuals:
##
       Min. 1st Qu.
                                         3rd Qu.
                        Median
                                   Mean
                                                      Max.
   -3.29800 -0.38660 0.11770 -0.01354
                                          0.57410
                                                   2.54600
##
## Location coefficients (identity link function):
                            2.5%
##
                Mean
## beta 0 0.0003161 -0.1488290 -0.0013337
## beta 1 -2.0015679 -2.0082464 -2.0015799 -1.995
```

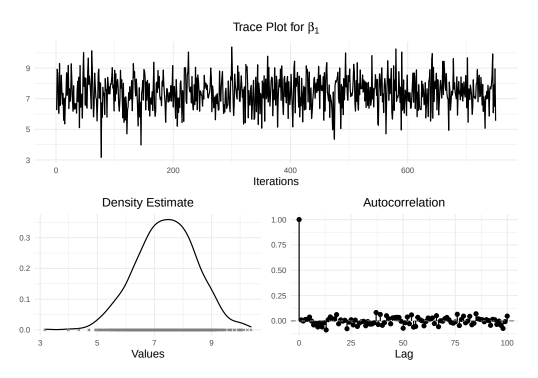


Figure 4: Diagnostic Plots for  $\beta_1$  after thinning, data: abdom

```
## beta 2 -1.0044943 -1.0179012 -1.0044382 -0.991
## beta 3 1.0014607
                     0.9802516
## beta_4 2.0080638 2.0015876
                                2.0081264
##
##
  Scale coefficients (log link function):
##
              Mean
                      2.5%
                               50%
##
   gamma_0
           0.9373 -0.7053
                           0.8277
                                    3.074
  gamma_1 -1.1471 -1.5038 -1.1357 -0.862
  gamma_2 0.9325
                   0.7238
                           0.9305
##
## Residual degrees of freedom: 42
## Log-likelihood: 32.28
## AIC: -48.57
## BIC: -33.27
```

One downside of this approach is that the displayed values are not saved anywhere by default and, thus, cannot be immediately accessed. This issue is solved by the more informative summary\_complete() function, which conveniently saves all relevant quantities in a data frame, the central object for data analysis in R.

In the following section we focus on the toy\_data model, since there the true coefficient values are known, and use the popular dplyr package for further data frame manipulations. We are primarily interested in Posterior Mean estimates for all coefficients, such that we only select a subset of the output columns and add a new column containing the true data generating values:

```
library(dplyr)

true_beta <- c(0, -2, -1, 1, 2)

true_gamma <- c(0, -1, 1)

summary_complete(samples = toy_fit) %>%
```

```
filter(stringr::str_detect(Parameter, pattern = "beta|gamma")) %>%
 mutate(Truth = c(true_beta, true_gamma)) %>%
 select(Parameter, `Posterior Mean`, Truth, `Standard Deviation`)
## # A tibble: 8 x 4
##
     Parameter `Posterior Mean` Truth `Standard Deviation`
##
     <chr>
                           <dbl> <dbl>
                                                        <db1>
## 1 beta_0
                        0.000316
                                                     0.0769
                                     0
## 2 beta 1
                       -2.00
                                     -2
                                                     0.00339
## 3 beta 2
                       -1.00
                                     -1
                                                     0.00676
## 4 beta 3
                        1.00
                                     1
                                                     0.0104
## 5 beta 4
                        2.01
                                     2
                                                     0.00315
## 6 gamma 0
                        0.937
                                      0
                                                     0.967
## 7 gamma_1
                       -1.15
                                     -1
                                                     0.163
## 8 gamma_2
                        0.933
                                      1
                                                     0.109
```

Except for  $\gamma_0$ , all coefficients are estimated very accurately with a slightly larger deviation form the true values for the remaining  $\gamma$  vector, which is sampled by the Metropolis-Hastings algorithm, compared to the  $\beta$  vector, which is drawn directly from a multivariate normal distribution.

Similar to the plotting functions introduced in section 2.1.2, the summary\_complete() function is very robust with respect to its data input: Besides the whole model object as in the example above, just the toy\_fit\$mcmc\_ridge list entry or even simply the sampling matrices toy\_fit\$mcmc\_ridge\$sampling\_matrices are valid inputs, all leading to the same output.

Further, the function has the optional <code>include\_plot</code> argument. If set to TRUE, one additional <code>Plot</code> column is added to the data frame, which leverages the <code>diagnostic\_plots()</code> function with some default settings and, thus, contains diagnostic plot objects for all coefficients. This option comes in handy in an interactive workflow: Particularly interesting findings from the <code>summary\_complete()</code> output can be quickly extracted from the data frame without interrupting the current thought process by using the <code>diagnostic\_plots()</code> function separately.

In the example above, the gamma\_0 row shows a large posterior variance in addition to the large deviation of the Posterior Mean estimate from the true value. In order to investigate, if the chain suffers from a lack of convergence or a significant autocorrelation, we could create the desired (yet here omitted) output with the following simple command:

```
summary_complete(toy_fit, include_plot = TRUE) %>%
filter(Parameter == "gamma_0") %>%
pull(Plot)
```

This introductory tutorial is not intended to validate the sampler's performance nor to interpret the obtained coefficient estimates, but rather to provide an overview of the various applications, where the asp21bridge package turns out to be useful. A more in depth analysis of the mcmc\_ridge() results, also in comparison to alternatives like the lmls() and mcmc() functions of the underlying lmls package, can be found in chapter 3 after explaining some of the internal implementations in section 2.2.

# 2.2 Implementation of the Markov Chain Monte Carlo Sampler

Of all components that the asp21bridge package is built upon, the implementation of the mcmc\_ridge() function is of special interest from a statistical perspective. Thus, section 2.2.1 explains both the overall structure of the mcmc\_ridge() function and the Gibbs Sampler that is responsible for inducing the Ridge penalty. Section 2.2.2 is dedicated to the integrated Metropolis-Hastings step, which is used for sampling  $\gamma$  and, if desired,  $\beta$  as well.

### 2.2.1 Iterative Parameter Sampling

### 2.2.2 Metropolis Hastings Step

# 2.3 Package Development

This section is dedicated to more niche aspects, that come along with the package development process. Part 2.3.1 focuses on more formal components of the package, whereas some of the ideas behind the asp21bridge functions are discussed in section 2.3.2.

### 2.3.1 Code Coverage

Arguably the most important component of a package for new users is the **documentation**. All exported functions of the <code>asp21bridge</code> package as well as the <code>toy\_data</code> data set are fully documented according to common standards for R packages.

Specifically, we put effort into precise, yet not excessively long descriptions of the overall function and their parameters with highlighted default settings. The help pages are designed uniformly across all functions. The examples section usually starts with the most basic applications signaling the function's *intent* and proceeds with more complex use cases that cover many of the function's optional arguments.

A more elaborate, publicly accessible tutorial similar to section 2.1 of this report can be found in the README file or the front page of the asp21bridge GitLab website.

A second major aspect, that contributes to the quality of a package, are **unit tests**. Up to this point, a total of 251 unit tests have been written for the lmls and the asp21bridge packages combined. The number of implemented tests alone is, of course, not a meaningful metric, since the tests might be highly redundant and capture only a small fraction of the entire package's functionality. Thus, we prioritized both *width* and *depth* of the test coverage.

More specifically, unit tests are written for every single exported function with a larger focus on the more complex and more central functions such as the main mcmc\_ridge() function.

Further, not only simple input checks (which are important nonetheless!), but also more esoteric edge cases and in particular very common use cases of combining inputs and outputs of multiple asp21bridge functions are covered. Whenever discovering and fixing an unexpected error during the development phase, we implemented corresponding unit tests, such that this specific error will not reoccur in the future. Tests are continuously written and updated to ensure a stable and enjoyable user experience.

At a larger scope, the R CMD CHECK, or equivalently the devtools::check() command, produces 0 errors, 0 warnings and 0 messages. Besides working unit tests, this includes correctly specified DESCRIPTION and NAMESPACE files for all imported and exported functions as well as functioning examples in the documentation. The master branch of the asp21bridge project will maintain this standard in the foreseeable future; possible new features that could involve breaking changes will first be implemented on separate Git branches and merged into the master branch at a later stage after a thorough testing procedure.

### 2.3.2 Design Choices

Throughout the development process, we tried to adhere to some 'best practices' for general software development. These include several different aspects:

#### **Default Settings:**

For many asp21bridge functions, the user has to specify very few inputs manually, since many input options are set to sensible default arguments. These inputs are chosen in a *neutral* way, such that the output aligns as best as possible with the user's expectation.

As an example, the default starting values for  $\beta$  and  $\gamma$  are the Maximum Likelihood estimates from the lmls() function, if a model object is provided as input to the mcmc ridge() function. This serves as guidance

for users with little prior knowledge about their model. It is worth noting, however, that there are multiple input parameters which *can* be set manually if desired, enabling fine control over the sampling procedure.

# Input Flexibility / Defensive Programming:

As already mentioned in section 2.1, all exported functions are designed to be as flexible as possible with respect to their input arguments. Thus, many functions accept multiple data structures like (nested) lists, data frames, matrices or even simple vectors as data input and internally transforms the input into the desired format.

These transformations, however, are only performed as long as the user's *intention* is clear, e.g. providing the samples as input to a plotting functions regardless of the exact data structure, where the samples are stored. Whenever there is ambiguity or the input is simply not valid / incomplete, we have put effort into writing informative error messages.

To illustrate this last point, assume that we had forgotten to provide the second design matrix Z as input to our second application example from section 2.1 using the abdom data set. This might happen quite frequently, especially if the user is not particularly familiar with the structure of location-scale regression models:

```
y <- abdom$y
X <- as.matrix(abdom$x)

abdom_fit <- mcmc_ridge(
    y = y, X = X, beta_start = 1, gamma_start = 1, num_sim = 10000
)

## Error in validate_input(m = m, X = X, Z = Z, y = y, beta_start = beta_start, :
## At least either all model matrices (X, Z, y) and coefficients
## (beta_start, gamma_start) or a model object (m) must be given.</pre>
```

From the error message, it is immediately obvious which part of the input is missing.

One further quite neat feature is implemented for the generic summary() function. Since the possible values for the type argument are specific to the lmls class and therefore a potential reason for confusion, the error message provides some guidance in case of spelling mistakes, suggesting the closest valid input option:

```
summary(toy_fit, type = "mcmc-ridge")
## Error: `type` must be one of "ml", "boot", "mcmc", or "mcmc_ridge".
## Did you mean "mcmc_ridge"?
```

### Modularity:

The lmls package served as a fantastic example how to design and compose functions in a modular way, leading to independent applications of single small functions in various contexts, easier debugging and arguably better code transparency and readability.

The asp21bridge package similarly splits large functions into multiple pieces according to the well known premise, that functions should be doing one thing only, but one thing well. Therefore the  $mcmc_ridge_helpers$  file contains various helper functions for the main  $mcmc_ridge()$  function, which perform tasks like input validation or inclusion of the Metropolis-Hastings step for the  $\gamma$  vector.

However, there is certainly a trade off to keep in mind: Since naming functions and variables is notoriously challenging, excessive modularity with poor naming choices can hinder code comprehension by inducing a wrong mental model of the function's task. Moreover, it can make sense to keep related functions close together in a physical sense and avoid spreading them across multiple files across the project.

### Coding Style:

Before starting our work on different parts of the code base, we agreed to a consistent coding style. This includes using only lowercase letters and underscores in function and variable names and a limit of 80

characters per line. Here, we mostly conformed to the recommendations from the tidyverse style guide by Hadley Wickham. Moreover, we leveraged the styler package for consistent and aesthetically pleasing code formatting.

To extend consistent naming schemes even further, we chose the same argument names for all exported functions whenever possible. For instance, the data input for all functions that expect the MCMC simulations is called samples and all shared arguments of the 5 plotting functions are assigned the same name and the same functionality. Hence, it often suffices to be familiar with one function of a certain family, since that knowledge can be easily transferred to all related functions.

These conventions allow both new and experienced users of the package easier file navigation and orientation as well as a more enjoyable programming experience overall. Finally, there are two design choices, that are very specific to the asp21bridge context.

## Operation Chaining:

First, we aimed to allow for frictionless function composition via the popular %>% operator. As illustrated in section 2.1, this comes in particularly handy when including burnin() and thinning() operations in a larger pipeline without the need to define dummy variables for temporary objects:

```
lmls(
  location = y ~ x1 + x2 + z1 + z2, scale = ~ z1 + z2,
  data = toy_data, light = FALSE
) %>%
  mcmc_ridge(num_sim = 1000) %>%
  purrr::pluck("mcmc_ridge", "sampling_matrices") %>%
  burnin(num_burn = 100) %>%
  thinning(freq = 5) %>%
  mult_plot(type = "both", free_scale = TRUE, latex = TRUE)
```

This workflow proved to be so useful, that we provided access to the pipe operator directly by loading the asp21bridge package, i.e. we reexported %>% from the magrittr package. Alternatively, one could use the native |> pipe introduced in R 4.1. However, since the release of this R version was so recent, it might induce a stronger dependency than just relying on the %>% operator, which is ubiquitous at least in the R community that is related to the data science field.

There are two conditions that functions must fulfill to chain multiple operations:

- The first argument must be chosen uniformly for all (except the first) involved commands. In case of the asp21bridge package, all exported functions share the samples argument at the first position.
- Functions that serve as intermediary steps inside a chain should return the same object type as their input. When given a lmls model object, the mcmc\_ridge() function returns a modified (copy of the same) lmls model object. Even more flexible are the burnin() and thinning() functions. They can take a list of matrices, a single matrix or a numeric vector as input and always return the same provided input data structure.

### **Object-Oriented Programming:**

Secondly, we thought about initializing our own S3 class when calling the mcmc\_ridge() function, but finally decided against it for two reasons: We always considered the asp21bridge package as a strict extension to the lmls package. Thus, we did not want to make the underlying lmls() results less accessible after using mcmc\_ridge(), i.e. generic functions like coef(), plot() or print() should in our mind create the same output before and after extending the original model.

One exception is the summary() function, as we have already mentioned. Here, we added one more option to the type argument, which has to be specified anyway, as soon as the boot() and/or mcmc() function of the lmls package are used. Apart from that, we could not see much additional value in implementing wrapper functions, which adapt e.g. the print() output to the MCMC results with penalty, beyond the already existing tools, that were introduced in section 2.1.

# 3 Simulation Studies

This chapter investigates the effect of manipulating some of the mcmc\_ridge() inputs and thereby sampling parameters in a controlled environment, such that changes in the estimation outcome can be directly linked to respective changes in the model inputs. The simulation studies discussed in this chapter can be categorized into three groups:

1. Evaluating the **quality** and **robustness** of the sampler.

Sections 3.1 and 3.4 create data configurations that can be challenging for estimation. The variation in the input parameters is therefore focused on the function argument m or alternatively the combination of X, Z and y. Section 3.2 is dedicated to manipulating the sample size through the input parameter n. Here, we are looking for a possible stabilization process with increasing size of the input data hinting at asymptotic/convergence properties.

2. Investigating the **penalty** effect.

Sections 3.3 and 3.4 replicate real world estimation scenarios, where the mcmc\_ridge() sampler is primarily chosen for the desired regularization effect. To emphasize the shrinkage effect on the coefficient estimates, the mcmc\_ridge() model is compared to the lmls() and the mcmc() models from the lmls package, which do not leverage a penalty.

3. Exploring the effect of the **hyperparameters**.

Section 3.5 analyzes the connection between variation of the input parameters a\_tau, b\_tau, a\_xi and b\_xi of the Inverse Gamma prior distribution of the variance parameters  $\tau^2$  and  $\xi^2$ , as specified in chapter 1. The effect of hyperparameters in a hierarchical Bayesian model can be difficult to predict based on pure logical reasoning. Therefore simulations are a useful tool to either confirm prior assumptions or discover unexpected behaviour.

Since the resulting simulation studies serve such diverse purposes (e.g. diagnostic vs. explorative), they demand for different approaches in the simulation settings, the implementation as well as the analysis and presentation of the results. For that reason, we decided against forcing all of the following sections into one common rigid framework. Instead, each section individually motivates, explains and interprets the methods chosen for its particular use case.

In order to keep the analysis in this chapter compact and succinct, there will be almost no code included. It it worth noting though that the R Markdown document itself as well as all R Scripts used for the simulations are contained in the simulation-studies folder of the asp21bridge package. Thus, each figure as well as all numerical results are fully reproducible and can be repeated and extended by the reader.

### 3.1 Correlated Predictor Variables

Up to this point, we have often illustrated the usage and results of the mcmc\_ridge() sampler with simulated data from the built-in toy\_data set. As stated in section 2.1, each regressor variable is independently sampled from a normal distribution and the outcome variable is simulated based on the correctly specified location-scale regression model introduced in chapter 1. All these conditions lead to an excellent performance of the mcmc\_ridge() sampler, but might arguably not represent the most challenging task.

Sections 3.1 and 3.4 analyze the sampler's performance on simulated data, which might be closer to data found in the real world. First, we will induce correlation among the predictor variables, whereas in a later part the distributional assumptions are considerably changed. Further, the mcmc\_ridge() performance is compared to the Maximum Likelihood based lmls() estimates and the Markov Chain Monte Carlo mcmc() sampler.

#### Simulation Setting

Briefly stated, we simulated data from a three dimensional multivariate normal distribution with three different values for the pairwise correlation  $\rho$ . Then, we fitted the three models mcmc\_ridge(), mcmc() and

# Empirical 90% Confidence Intervals for Posterior Mean Estimates True coefficient values are marked by grey circles

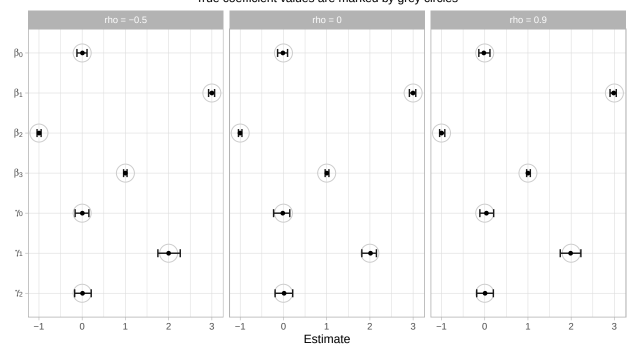


Figure 5: Comparison of Correlation Structures - 50 Simulation Cycles

lmls(), where the number of simulations was set to 10000 for the Bayesian Samplers.

This simulation study is intended to evaluate the *accuracy* of the three models. Therefore, the same (standardized) values of the covariates are used for both, the generation of the outcome variable **y** and the model fit. This entails that the true coefficient values are known and can be compared to the resulting estimates of all three and in particular the mcmc\_ridge() model. The exact simulation setting is described in full detail in the Appendix.

Moreover, we compared the performance of the classical  $mcmc\_ridge()$  implementation, which draws  $\beta$  from the closed form full conditional (multivariate normal) distribution, with an alternative sampling process that uses a Metropolis-Hastings approach for both, the location parameter  $\beta$  as well as the scale parameter  $\gamma$ .

### **Simulation Results**

All three models perform very similarly for each correlation structure: The estimates for  $\beta$  are almost exactly equal to the true values, whereas there are slight deviations for the  $\gamma$  vector. These errors, however, are consistent in all three sampling scenarios and, thus, do not depend on the strength nor the direction of the correlation between the covariates. Due to this similarity between the three models, the remaining analysis of this simulation study is dedicated to the mcmc\_ridge() model.

There is one additional aspect to be noted: The performance of the  $mcmc_ridge()$  function is considerably worse, when  $\beta$  is sampled by means of the Metropolis-Hastings algorithm. Although the acceptance rates of the proposed values are in acceptable ranges for a random walk proposal (between 35% and 40%), the chains are strongly correlated and did not fully converge even after 10000 iterations. For that reason, we limit the Metropolis-Hastings sampling process for  $\beta$  to this one example and will focus on the classical  $mcmc_ridge()$  implementation in the remaining parts of chapter 3.

The corresponding graphical display for the comparison between the three models and between the two sampling procedures for  $\beta$  is omitted here for brevity reasons, since the findings are not that surprising. The

code as well as all plots can be found and fully reproduced in the regressor-correlation.R file located inside of the simulation-studies folder of the project directory.

In order to make any conclusions about bias and variance, the above procedure is repeated 50 times with 1000 simulations each. The black points in Figure 5 represent the mean of these 50 Posterior Mean estimates from the penalized Ridge sampler. For a better visual comparison, the true values for each coefficient are indicated by grey circles. Since we cannot rely on distributional theory for the standard errors, the variability of the estimates is displayed by nonparametric 'confidence' intervals, which are simply given by the range from the empirical 0.05 quantile to the 0.95 quantile of the 50 estimated values.

These intervals are very narrow and centered around the true value in all cases with slightly larger variability for the  $\gamma$  vector, which is sampled via the Metropolis-Hastings algorithm. Overall, these results impressively solidify the sampler's stability in presence of correlated covariates. The bias towards zero that is typical for the Ridge penalty cannot be observed in this case; all coefficients are estimated with both high accuracy and high precision.

It is worth noting that the results are partially affected by the standardization of the covariates. The standardization generally leads to a stabilized fit for all three models. Compared to the performance for unstandardized data (as used for the second report) the variability of the Posterior Mean estimates of  $\beta_0$  in particular is significantly reduced.

# 3.2 Sample Size

This simulation study analyzes the effect of the sample size n on the means of the posterior distribution for the coefficients of  $\beta$  and  $\gamma$ . There are two main goals of this simulation study: On the one hand, we want to investigate whether the posterior means of large samples are closer to the true values than the posterior means of small samples. On the other hand, we want to analyze whether the  $mcmc_ridge()$  penalty affects the location of the posterior means.

### Simulation Setting

• The design matrix  $\mathbf{X} = \begin{pmatrix} \mathbf{1}_n & \mathbf{x}_1 & \mathbf{x}_2 \end{pmatrix}$  contains two independently sampled regressor variables plus one intercept column:

$$- \mathbf{x}_1 \overset{iid}{\sim} \mathcal{N}(1,1),$$
  
$$- \mathbf{x}_2 \overset{iid}{\sim} \mathcal{N}(2,1).$$

• The design matrix  $\mathbf{Z} = \begin{pmatrix} \mathbf{1}_n & \mathbf{z}_1 & \mathbf{z}_2 \end{pmatrix}$  is structured in the same way with the regressor variables:

$$- \mathbf{z}_1 \stackrel{iid}{\sim} \mathcal{N}(1,1),$$
  
$$- \mathbf{z}_2 \stackrel{iid}{\sim} \mathcal{N}(2,1).$$

- The true coefficient vectors are given by  $\boldsymbol{\beta} = \begin{pmatrix} \beta_0 & \beta_1 & \beta_2 \end{pmatrix}^T = \begin{pmatrix} 1 & -1 & 4 \end{pmatrix}^T$  and  $\boldsymbol{\gamma} = \begin{pmatrix} \gamma_0 & \gamma_1 & \gamma_2 \end{pmatrix}^T = \begin{pmatrix} 0 & -0.5 & 1 \end{pmatrix}^T$ .
- The posterior means are analyzed with respect to 6 different sample sizes:  $n \in \{0, 50, 100, 200, 300, 500\}$ .
- In the next step, the outcome vector  $y \in \mathbb{R}^n$  is simulated and passed to the mcmc\_ridge() function with nsim = 500 simulations.
- To make the results more stable, the above procedure is repeated 100 times. For each coefficient, the mean value of the Posterior Mean estimates of each coefficient is calculated as well as the Mean Absolute Error (MAE) with respect to the true values of  $\beta$  and  $\gamma$ .

### Simulation Results

The means of the Posterior Mean estimates are displayed in Figure 6. For larger sample sizes ( $n \ge 200$ ) none of the six parameters are extremely biased.

### **Mean of Posterior Means**

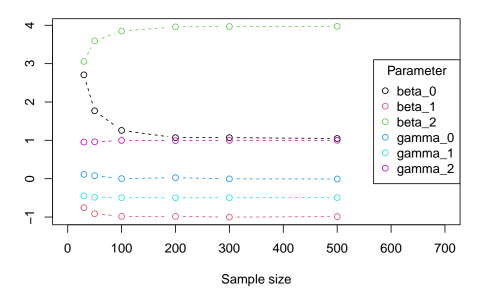


Figure 6: Mean value of 100 Posterior Mean Estimates

Moreover, for n = 30,  $\beta_0$  and  $\beta_2$  are significantly biased, which might be caused by the high mcmc\_ridge() penalty for  $\beta_2 = 4$ . The significant bias of  $\beta_0$  might be explained by a counteract of the  $\beta_2$  bias.

After getting an impression about empirical biases of the coefficients, we now focus on the variability of the posterior means of the coefficients, which are measured by the MAE based on the results of the 100 repetitions. Figure 7 points out that the posterior means of  $\beta_0$  have significantly larger errors than the posterior means of  $\beta_2$  for n = 30. However, this might also be explained by the fact that for n = 30,  $\beta_0$  has a greater empirical bias than  $\beta_2$  as could be observed in Figure 6.

In addition, for increasing sample sizes, the MAE of the Posterior Means tend to zero for all coefficients except  $\beta_0$ . Nevertheless, also the errors of  $\beta_0$  seem to become smaller with increasing sample size.

### 3.3 Redundant Covariates

Similar to the simulation study from section 3.1, this section investigates the sampler's behaviour in presence of strong *pairwise* correlation among the covariates. However, in contrast to all other simulation studies, we add additional *redundant* regressors to the model, that were not used for generating the outcome variable **y**.

The statistical theory of (Ridge) penalization suggests, that the coefficient estimates from the  $mcmc_ridge()$  function are affected by the shrinkage effect induced by the coefficient's prior distributions in presence of high (compared to the sample size n) number of correlated covariates. Thus, the magnitude of the estimated  $\beta$  vector should be smaller compared to models without regularizing penalty component.

### Simulation Setting

The model contains 10 pairs of covariates drawn from a multivariate normal distribution with correlation  $\rho = 0.9$ . For each pair, the first covariate contributes to generating the outcome variable  $\mathbf{y}$  with a true coefficient value of 1. The second covariate has no impact on  $\mathbf{y}$ , but is included nonetheless in the model fitting stage. Similar to section 3.4, the lmls(), mcmc() and mcmc\_ridge() functions are used to generate estimates.

Since the purpose of this study is to demonstrate the Ridge regularization and not primarily the accuracy of the three models, we used the covariates on their original scale for generating y and standardized them

### **MAE of Posterior Means**

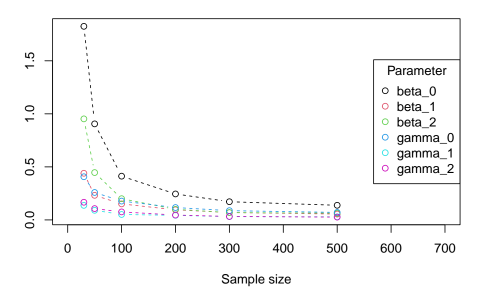


Figure 7: Mean Absolute Error Estimates

afterwards before fitting the models. Thus, the true coefficient values are not known and conclusions about the model *quality* are not valid. This scenario resembles a real world application, where the data generating process is latent and estimates are obtained based on purely observational data. The exact design of the simulation study can again be found in the Appendix.

### Simulation Results

The following analysis is exclusively focused on the location parameter  $\beta$ . As mentioned before, there are two aspects of interest we wish to investigate further:

- 1. Can each model differentiate between the relevant and the redundant covariate for each pair of regressor and, thus, isolate the true effect on  $\mathbf{y}$  (without paying too much attention to the *absolute* size of the estimated values)?
- 2. Due to the high number of 20 covariates compared to the low sample size of 50 observations and the presence of correlated and partially redundant regressors, the setup of this study is prone to overfitting. Can we therefore observe the hypothesized shrinkage effect on the coefficient estimates for the mcmc\_ridge() model compared to the models without penalty?

Figure 8 illustrates the results obtained by fitting all three models.

We start with the answer to the first question: For each of the three models and each coefficient pair the absolute value of the coefficient corresponding to relevant regressor is larger than the magnitude of the paired quantity. Thus all three models always contribut the major impact on the outcome variable to the correct covariate. Further, the differences between estimations for those coefficients with true value 1 and those with true value 0 do not differ in an *obvious* manner across the models. Since the variance of the estimates seems to be largest for the lmls() model and smallest for the mcmc\_ridge() model, the ratio (instead of the difference) tends to be largest for the regularized model.

The reduced variance of the mcmc\_ridge() model can be explained by the Ridge penalty: Since all estimates are shrunken towards zero, fewer 'outlier' estimates that are far away from zero will be observed. This property is beneficial for the red points, since the estimates clutter around their true value in this case. However, the coefficients corresponding to the green points are penalized as well such that a bias towards

### Posterior Means / MLE for pairwise correlated Covariates

True Coefficient Value: - 0 - 1

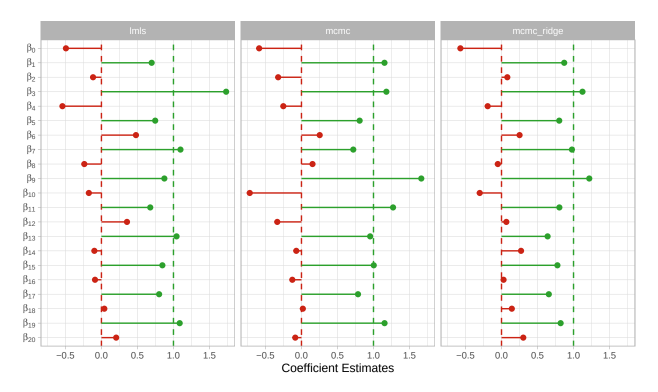


Figure 8: Shrinkage Effect of Ridge Penalty in Presence of Redundant Covariates

zero is induced, which is not present for the two unpenalized models. Hence, the hypothesized shrinkage effect can indeed be observed.

It is insightful to compare the observed effect from the Ridge penalty to the expected effect of the *LASSO* penalty. The LASSO penalty encourages *sparse* solutions: each coefficient is assigned the same weight such that shrinkage of large estimates is not preferred over shrinkage of small estimates. Thus, small estimates are often set to exactly zero, while large estimates, although reduced, can still be significantly different from zero.

In contrast, the Ridge penalty generally does not induce sparsity. In fact, quite the opposite can be the case: Small coefficient estimates such as  $\beta_{14}$  in Figure 8 are sometimes *increased* in magnitude compared to the unpenalized models. Large estimates such as  $\beta_4$  or  $\beta_{12}$ , however, are reduced in size quite heavily. The Ridge penalty tries to put all coefficients on the same scale and, thus, prioritizes shrinkage of large estimates compared to small ones! Note also, that the intercept term is typically excluded from the penalization, wich is the reason why the estimates of  $\beta_0$  are of approximately equal size for all three models.

In order to quantify the observed shrinkage effect from Figure 8 numerically, we calculate the sum of squared coefficient estimates, i.e. the squared Euclidean norm  $\|\beta\|^2$ , for each model. Moreover, these magnitudes are compared separately for all coefficients corresponding to relevant regressors (those with an *odd* subscript) and those corresponding to redundant regressors (*even* subscript). Table 1 summarizes the results.

The most interesting and reassuring finding is obtained from the mcmc\_ridge() vector norms: In agreement with the visual illustration, the Ridge regularization effect can be numerically detected for all coefficient estimates, independent of the true value. While this property induces a bias that is not desired for the second column (underestimating the true values on average), it does indeed prevent overfitting by shrinking the coefficients corresponding to the redundant covariates, which is indicated by the lowest value in the first column.

Table 1:  $\|\boldsymbol{\beta}\|^2$  excluding  $\beta_0$ 

	True Value $= 0$	True Value $= 1$
lmls	1.05	10.09
mcmc	1.26	12.13
$mcmc\_ridge$	0.72	7.87

Table 2:  $\|\boldsymbol{\beta}\|^2 + \|\boldsymbol{\gamma}\|^2$  excluding  $\beta_0$  and  $\gamma_0$ 

	Normal	t	Uniform
lmls mcmc	1906 1903	2024 $2054$	2088 2100
mcmc_ridge	1222	1609	950

In that sense, the Ridge penalty increases the model *robustness* in presence of a high number of (correlated) regressors and often leads to an overall lower Mean Squared Error due to the reduction in variance.

# 3.4 Challenging the Model Assumptions

This simulation study is structured in a very similar way to the study considered in section 3.1. Instead of varying the correlation structure among the regressors in the underlying data set, both the regressors and the outcome variable y are sampled from distributions that are more challenging for estimation than the normal distribution.

### Simulation Setting

The covariates of the design matrices  $\mathbf{X}$  and  $\mathbf{Z}$  are generated by a mix of distributions including the Normal, Exponential, Uniform, Bernoulli and t - distribution. Given these values, the outcome variable  $\mathbf{y}$  is drawn from three different distributions (Normal, t and Uniform) corresponding to three separate scenarios, which are compared in the following analysis. As usual, the full simulation design is stated in the Appendix.

Note that the lmls(), mcmc() and  $mcmc\_ridge()$  models are built upon the assumption of a Gaussian distribution for y. Hence, we expect all three estimation procedures to perform well under the first outcome specification, which they were designed for. The remaining two cases analyze the performance in presence of a mild (t distribution) and a moderately strong (Uniform distribution) violation of this model assumption.

### Simulation Results

Just as in section 3.1 the results of one complete simulation cycle (each of the  $3 \cdot 3 = 9$  models was fitted once / each data point represents one estimate) are displayed in Figure ??. Note that the second facet is labeled by  $y \sim t$ , although it is formally sampled from an affine transformation of a t-distributed random variable, which does not follow an exact t distribution.

The differences within each facet as well as between the facets are significant. All three models seem to estimate the  $\beta$  vector well, when there are no or only mild violations of the normal assumption for y. If y is sampled from a uniform distribution, there are major differences for  $\beta_0$ ,  $\beta_2$  and  $\beta_4$  (notice the extended x-scale in the third facet). Interestingly, the  $\gamma$  vector is estimated very well in the latter case with more deviations in the setting, where y is based on the t distribution.

To gain insights beyond this single simulation cycle, which could well be disturbed by random noise, we repeat the sampling process 50 times. The resulting means as well as empirical confidence intervals (analogous to section 3.1) are plotted in Figure 9.

## Empirical 90% Confidence Intervals for Posterior Mean Estimates

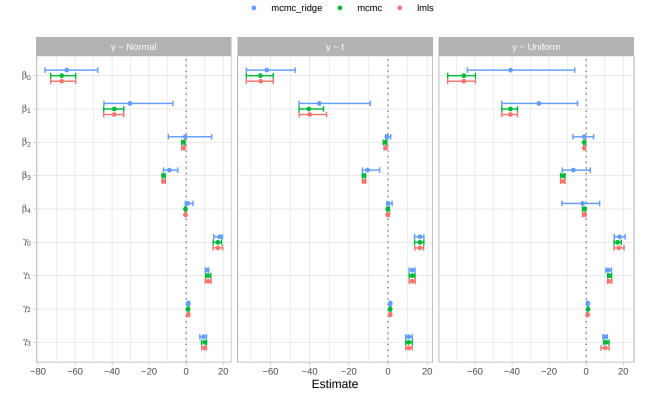


Figure 9: Comparison of different Outcome Distributions - 50 Simulation Cycles

This plot (literally) paints a drastically different picture, emphasizing the necessity of repeating experiments multiple times whenever possible. Across all 50 simulations the deviation of the estimates for  $\beta_2$  (corresponding to the regressor variable from the exponential distribution) is huge for all three distributional specifications of y.

The small bias induced by all three models is negligible compared to the wide confidence intervals, which is particularly interesting, when y stems from a normal distribution. In this case all models should perform well, however the lmls() and the mcmc() Posterior Mean / Maximum Likelihood estimates vary wildly across the simulation cycles. A similar effect can be observed for  $\beta_0$  in case of the uniform distribution. Here, all models overestimate the true value on average, while the  $mcmc_ridge()$  function again shows the smallest variability. In contrast, the estimates for  $\gamma_0$  in the right facet are fairly stable across simulation cycles, but also consistently wrong at the same time.

The standard error estimates displayed in Table ?? clearly indicate the worst performance of the lmls() and the mcmc() model for  $\beta_2$ . In almost all cases (and sometimes very significantly), the  $mcmc_ridge()$  sampler has the smallest standard error. This finding nicely confirms the underlying mathematical theory: The present prior specifications in the Bayesian setting, which induces the equivalent form of a frequentist Ridge penalty, can lead to biased estimation.

However, this loss in accuracy can be (as it is in this case) dominated by the gain in precision by the shrinkage effect of the penalty. Note that (except for  $\gamma_0$  in the most right facet in Figure 9) the mcmc\_ridge() sampler slightly overestimates coefficients with true negative values and underestimates those with true positive values. This again is caused by the Ridge penalty leading to estimated coefficients close to zero.

In summary, the following conclusions can be drawn:

- 1. All three models are affected by changes in the regressor and/or outcome distributions. In the former case the regressor variables sampled from the Exponential and the Bernoulli distributions were the greatest challenge, in the latter case the outcome variable from the Uniform distribution. This is generally not surprising, since these distributions deviate most from the nicely behaved normally distributed case.
- 2. As expected, the lmls() function is affected strongly by violating the model assumptions, since its estimation process is based on the normal (log) Likelihood. Surprisingly, the mcmc() sampler without penalty often did not perform much better.
- 3. While the mcmc\_ridge() function does not excel at estimation accuracy, it does lead to the most stable estimation with smallest standard errors in the vast majority of cases. As emphasized above, this behaviour nicely agrees with the mathematical theory of Ridge penalization.
- 4. Had we not conducted repeated experiments, our conclusions would have been quite different. Simulation results are therefore always worth repeating many times to consolidate the correct interpretations.

### **Technical Aspects**

As outlined in the previous paragraph, a total of  $50 \cdot 3 \cdot 3 = 450$  models were fitted to analyze the performance differences. In order to speed up the involved computations of this specific and some of the other simulation studies in this report, we used the *parallel computing* capabilities of R.

There are many options from various packages to choose from. We decided to use the furrr package, which is built on top of the future package specialized on parallel processing. As the name suggests, furrr provides a convenient way to use many functions from the popular purrr package, while using multiple cores at the same time. This functional programming based approach (similar to the apply() family in 'base R') is particularly well suited for simulation studies and provides some structural as well as minor performance advantages compared to the classical for-loop approach.

The following (slightly modified) code snippet provides a brief insight into the implementation:

```
plan(multisession, workers = 8)

full_results <- tibble(id = 1:50) %>%
  mutate(samples = future_map(
    .x = id,
    .f = ~ show_results(n = 50, num_sim = 1000),
    .options = furrr_options(seed = 1)
  ))
```

The plan() function borrowed from the future package initializes the parallel computing process and determines the number of cores/workers available for computation. The show\_results() helper function fits all three models mcmc\_ridge(), mcmc() and lmls() for each outcome distribution in a single simulation cycle.

This entire procedure is repeated 50 times in parallel using the future\_map() function from the furrr package, where the results of all 450 models are saved in a well organized structure inside of a list column. This new column of the data frame contains complete information about all simulations, such that any required element for the further analysis can be easily extracted and post processed.

Finally, the .options() argument allows the specification of a random seed. Random number generation in the context of parallel computing is slightly more involved compared to the sequential approach. This additional complexity is automatically handled by the future\_map() function, such that all results are sampled in a statistically valid and fully reproducible manner.

# 3.5 Hyperparameters

In the past, we have been sampling data with the mcmc\_ridge() function without having a closer look on the effect of the hyperparameters and model inputs a\_tau, b\_tau, a\_xi and b\_xi. However, they affect the Full Conditional Distributions of  $\tau^2$  and  $\xi^2$ , as stated in sections 1.3.1 and 1.3.2 in chapter 1.

Moreover, the mean vector  $\boldsymbol{\mu}_{beta}$  and covariance matrix  $\boldsymbol{\Sigma}_{beta}$  of the  $\boldsymbol{\beta}$  vector both depend on  $\tau^2$  and, thus, implicitly on the hyperparameters  $a_{\tau}$  and  $b_{\tau}$  (see section 1.3.3). Analogously, section 1.3.4 illustrates the direct effect of the Full Conditional distribution of  $\boldsymbol{\gamma}$  on  $\xi^2$ , which in turn depends on the hyperparameters  $a_{\xi}$  and  $b_{\xi}$ .

Finally, cross effects can be observed, since  $f(\beta \mid \cdot)$  depends on  $\gamma$  through the quantities **W** and **u** as defined in chapter 1 and  $f(\gamma \mid \cdot)$  directly depends on  $\beta$ . These dependencies are reflected in the mcmc\_ridge() sampler by the iterative sampling procedure which is discussed in great detail in the previous sections 2.2.1 and 2.2.2.

Thus, the hyperparameter choice of  $a_{\tau}$ ,  $b_{\tau}$ ,  $a_{\xi}$  and  $b_{\xi}$  inevitably impacts the result of *all* coefficient estimates contained in the model in a nontrivial way, such that pure analytical reasoning might be misleading. For this reason, this section investigates these effects based on a simulation approach.

### Simulation Setting

- The design matrix  $\mathbf{X} = \begin{pmatrix} \mathbf{x}_1 & \mathbf{x}_2 \end{pmatrix}$  is simulated from a two dimensional normal distribution  $\mathcal{N}_2(\mu, \Sigma)$  with mean vector  $\boldsymbol{\mu} = \begin{pmatrix} 1 & 2 \end{pmatrix}^T$  and identity covariance matrix  $\boldsymbol{\Sigma} = \mathbf{I}_2$ . The same holds true for the design matrix  $\mathbf{Z} = \begin{pmatrix} \mathbf{z}_1 & \mathbf{z}_2 \end{pmatrix}$  with mean vector  $\boldsymbol{\mu} = \begin{pmatrix} 5 & 3 \end{pmatrix}^T$  and identity covariance matrix.
- In both design matrices intercept columns are added for estimation purposes. The true coefficient vectors are given by  $\boldsymbol{\beta} = \begin{pmatrix} \beta_0 & \beta_1 & \beta_2 \end{pmatrix}^T = \begin{pmatrix} 0 & -1 & 4 \end{pmatrix}^T$  and  $\boldsymbol{\gamma} = \begin{pmatrix} \gamma_0 & \gamma_1 & \gamma_2 \end{pmatrix}^T = \begin{pmatrix} 0 & -2 & 1 \end{pmatrix}^T$ .
- For sampling the location parameter, the full conditional multivariate normal distribution of  $\beta$  is chosen, i.e.  $mcmc\_ridge(..., mh\_location = FALSE)$  is used. Therefore, the location estimate is directly affected by the hyperparameters.
- For simulating the influence of the hyperparameters, nine different values are chosen:  $a_{\tau}, b_{\tau}, a_{\xi}, b_{\xi} \in \{-1, 0, 0.5, 1, 2, 10, 50, 100, 200\}$ . Since for statistical properties like the mean of an Inverse Gamma distribution  $\frac{b}{a-1}$  the condition a > 1 is required, particular attention is given to larger values. However, it is an aim to inspect the performance of the sampler for smaller hyperparameter values than 1 as well.

#### Simulation Results

The first two plots of Figures 10 and 11 display the absolute deviations of the Posterior Mean estimates from the true parameters with the stated different values for  $a_{\tau}$  and  $b_{\tau}$ . For each estimate, the Posterior Mean averages over 1000 simulations of the mcmc\_ridge() sampler. Note, that location and scale parameters are plotted separately, according to the relationship mentioned above. For a better overview, the dotted line displays the linear trend of all estimate deviations.

The x - axis is transformed by a pseudo logarithm in order to clearly visualize the deviations in the range of -1 to 10, which would not be possible on original scales. Since -1 and 0 are also part of the hyperparameter values, the pseudo\_log\_trans() function of the scales package is applied, log-transforming positive values only.

It can be observed, that the intercept estimates in each plot show the largest deviations from their true value. In Figure 10, however, the overall deviations of  $\beta$  estimates from their corresponding true value are small in absolute value. In contrast, deviations of the  $\gamma$  estimates in Figure 11 are fairly significant, especially for  $\gamma_0$ .

The functional chain that applies to the estimates of  $\beta$  can be described by the effect of the mean of the inverse gamma distribution on  $\tau^2$ : A larger value for  $b_{\tau}$  leads to larger values of  $\tau^2$ , which are again affecting the full posterior parameters of  $\beta$  and, thus, potentially increase the absolute deviation of the corresponding

# Absolute deviations from true beta (1000 simulations)

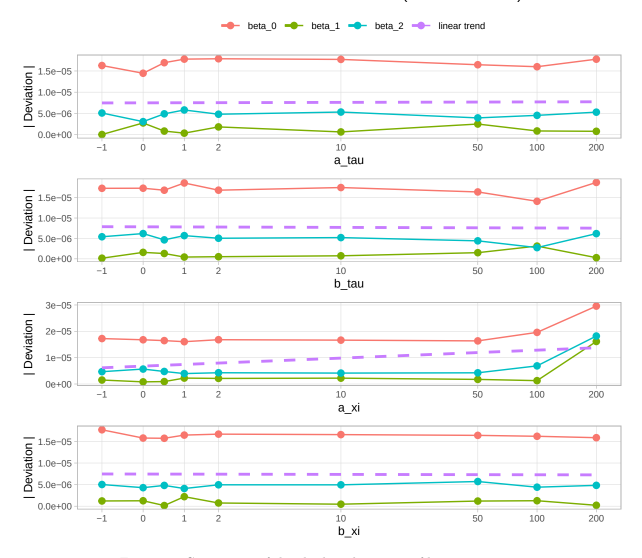


Figure 10: Comparison of the absolute deviations of beta parameters

estimates from their true values.  $a_{\tau}$  causes the opposite effect. This numerically observable effect, however, is hided by the overall small deviation in the first two plots of Figure 10.

It is remarkable, that the deviation of  $\beta$  estimates is smallest when  $a_{\tau}, b_{\tau} \in \{50, 100\}$ . For values of  $a_{\tau} \leq 1$ , one obtains wider variances of absolute deviations, since the Posterior Mean requires values larger than one.

In the upper two plots of Figure 11, there is no clear impact of  $\tau^2$  and its parameters. Rooted in no direct effect of  $\tau^2$  on  $\gamma$  according to our underlying mathematical model, one observes cross-effects through the sampling procedure of the mcmc\_ridge() sampler, where the full posterior  $f(\gamma \mid \cdot)$  depends on  $\beta$ .

Anyway, our sampler produces the lowest deviation of  $\gamma$  estimates for  $a_{\tau}, b_{\tau} \in \{0, 0.5, 200\}$ , where 0.5 is chosen by coincidence for  $a_{\tau}$  here, since wide variations for  $a_{\tau} \leq 1$  of absolute deviations are observable again.

The lower two plots of Figures 10 and 11 are constructed analogously, but showing the impact of  $a_{\xi}$  and  $b_{\xi}$  on the location and scale parameters respectively. Again, the overall absolute deviations for the  $\beta$  estimates from their true values are small, whereas the deviations for the  $\gamma$  estimates are considerably larger. Once again, the intercept estimates display the largest deviations from their true value.

## Absolute deviations from true gamma (1000 simulations)

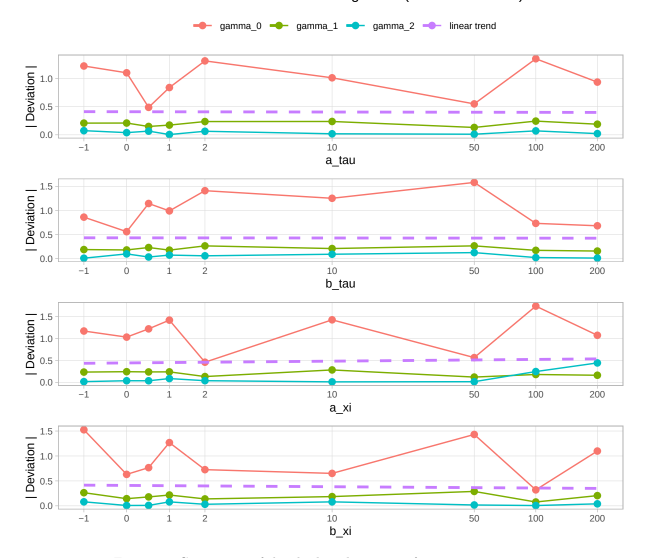


Figure 11: Comparison of the absolute deviations of gamma parameters

Arguing with the mean of the Inverse Gamma distribution of  $\xi^2$  in a similar way, one obtains larger mean values for  $b_{\xi}$ , while  $a_{\xi}$  lowers them. The impact of  $\xi^2$  on  $\gamma$  is assumed to decrease  $f(\gamma \mid \cdot)$  according to our underlying theoretical model. This effect is indicated by the linear trend lines in the second half of Figure 11.

In general, one obtains smaller deviations for larger values of  $b_{\xi}$  and lower ones of  $a_{\xi}$ , where especially lots of randomness occurs in the deviations of  $\gamma_0$ . Therefore, the impact of  $a_{\xi}$  and  $b_{\xi}$  on the scale intercepts is overshadowed by the randomness induced by the Metropolis Hastings algorithm. The same wide variations exclusively for  $a_{\xi} \leq 1$  cannot be obtained in the same manner as for  $a_{\tau}$ .

The sampler exhibits the best results for the *scale* estimates for  $a_{\xi} = 2$  and  $b_{\xi} = 100$ . However, due to the wide overall variation, these results must be taken with care.

The effect of  $a_{\xi}$  and  $b_{\xi}$  on  $\beta$  can be explained through the cross-effects of the matrix **W** and the vector **u** introduced at the beginning of this section, both containing  $\gamma$ . These diminish with increasing values of the  $\gamma$  entries.

The matrix **W** affects the variance of the full conditional distribution of  $\beta$  negatively, while the mean is positively affected. Hence, larger values of  $a_{\xi}$  cause higher Posterior Means of the location parameters. The

positive linear trend in the second half of Figure 10 for values of  $a_{\xi}$  is particularly interesting. For values of  $b_{\xi}$ , the trend comes off inferior. The wider variations of deviations for  $a_{\xi} \leq 1$  is again not observable here. Nonetheless, the randomness observable for scale estimates does not show up for location estimates anymore.

The smallest deviations of the *location* estimates can be detected for  $a_{\xi} = 1$  and  $b_{\xi} = 200$ .

Shortly noted, the acceptance rates of the Metropolis Hastings algorithm for sampling  $\gamma$  are always between 0.31 and 0.53. For the value range of  $a_{\tau}$ ,  $b_{\tau}$  and  $a_{\xi}$ , no distinct pattern is observable in this regard. With growing values of  $b_{\xi}$ , however, acceptance rates are more likely to grow. Since the acceptance rates are in reasonable ranges enabling statistically valid estimation, these results are not further investigated here.

# Conclusion

# **Appendix**

The following paragraphs contain additional information about the **simulation studies** that were covered in chapter 3.

# Section 3.1: Correlated Predictor Variables

The simulation design was chosen in the following way:

- The design matrix  $\mathbf{X} = \begin{pmatrix} \mathbf{x}_1 & \mathbf{x}_2 & \mathbf{x}_3 \end{pmatrix}$  is simulated from a three dimensional normal distribution  $\mathcal{N}_3(\boldsymbol{\mu}, \boldsymbol{\Sigma})$  with mean vector  $\boldsymbol{\mu} = \begin{pmatrix} -5 & 2 & 0 \end{pmatrix}^T$  and covariance matrix  $\begin{pmatrix} 1 & \rho & \rho \\ \rho & 3 & \rho \\ \rho & \rho & 5 \end{pmatrix}$ . Hence, the dependence among the regressors is fully determined by the parameter  $\rho$ .
- The design matrix  $\mathbf{Z} = (\mathbf{z}_1 \quad \mathbf{z}_2)$  consists of linear combinations of the regressors  $\mathbf{x}_1$  up to  $\mathbf{x}_3$ , more specifically  $\mathbf{z}_1 = 0.8 \cdot \mathbf{x}_1 + 0.2 \cdot \mathbf{x}_2$  and  $\mathbf{z}_2 = \mathbf{x}_2 0.5 \cdot \mathbf{x}_3$ . In both design matrices intercept columns are added for estimation purposes.
- The true coefficient vectors are given by  $\boldsymbol{\beta} = \begin{pmatrix} \beta_0 & \beta_1 & \beta_2 & \beta_3 \end{pmatrix}^T = \begin{pmatrix} 0 & 3 & -1 & 1 \end{pmatrix}^T$  and  $\boldsymbol{\gamma} = \begin{pmatrix} \gamma_0 & \gamma_1 & \gamma_2 \end{pmatrix}^T = \begin{pmatrix} 0 & 2 & 0 \end{pmatrix}^T$ .
- The outcome variable  $\mathbf{y}$  is generated according to the correctly specified location-scale model  $y_i \stackrel{iid}{\sim} \mathcal{N}\left(\mathbf{x}_i^T\boldsymbol{\beta}, \exp\left(\mathbf{z}_i^T\boldsymbol{\gamma}\right)^2\right)$  for  $i=1,\ldots,n$  with sample size n=50. Before data generation all columns in  $\mathbf{X}$  and  $\mathbf{Z}$  are standardized, i.e. mean-centered around 0 and scaled to unit variance.
- Three different values were chosen for  $\rho \in \{0, -0.5, 0.9\}$  to compare the 'nice' case of uncorrelated predictors with the performance for negative and positive dependence. For each covariance structure the three models  $mcmc_ridge(), mcmc()$  and lmls() are fitted to the standardized covariates, where each Posterior Mean estimate from both of the Markov Chain Monte Carlo samplers is based on 10.000 samples.

# Section 3.2: Sample Size

### Section 3.3: Redundant Covariates

We again state the conditions that the simulation study is based on:

- The design matrix  $\mathbf{X} = \begin{pmatrix} \mathbf{1}_n & \mathbf{x}_1 & \cdots & \mathbf{x}_{20} \end{pmatrix}$  consists of one intercept column plus 10 pairs of successive regressors, starting with the pair  $(\mathbf{x}_1, \mathbf{x}_2)$ . Each pair  $(\mathbf{x}_i, \mathbf{x}_{i+1})$  for  $i \in \{1, 3, \dots, 19\}$  is (independently from all remaining pairs) drawn from a bivariate normal distribution with mean vector  $\boldsymbol{\mu} = \begin{pmatrix} 0 & 0 \end{pmatrix}^T$  and correlation matrix  $\begin{pmatrix} 1 & 0.9 \\ 0.9 & 1 \end{pmatrix}$ .
- The design matrix  $\mathbf{Z} = \begin{pmatrix} \mathbf{1}_n & \mathbf{x}_1 & \mathbf{x}_3 \end{pmatrix}$  is of minor interest in this case and consists of an intercept column plus two uncorrelated columns chosen from  $\mathbf{X}$ .
- The true coefficients of  $\beta$  are determined by the pattern  $\beta_i = 0$ , if i is even and  $\beta_i = 1$ , if i is odd. Thus, all covariates with even subscript are redundant, whereas those with odd subscript contribute to  $\mathbf{y}$ . The true  $\gamma$ , again of minor interest here, is given by  $\gamma = \begin{pmatrix} 0 & 1 & 1 \end{pmatrix}^T$ .
- The outcome variable  $\mathbf{y}$  is generated according to the correctly specified location-scale model  $y_i \stackrel{iid}{\sim} \mathcal{N}\left(\mathbf{x}_i^T\boldsymbol{\beta}, \exp\left(\mathbf{z}_i^T\boldsymbol{\gamma}\right)^2\right)$  for  $i = 1, \dots, n$ , where the covariates are used on their original scale.
- The sample size n=50 is deliberately chosen small compared to the number of regressors. Before fitting each of the three models, all columns of **X** except the intercept column is standardized to zero mean and unit variance. Both of the Bayesian models generate 10.000 values for each coefficient.

## Section 3.4: Challenging the Model Assumptions

The data for this simulation study is generated by the following conventions:

- The design matrix  $\mathbf{X} = \begin{pmatrix} \mathbf{1}_n & \mathbf{x}_1 & \mathbf{x}_2 & \mathbf{x}_3 & \mathbf{x}_4 \end{pmatrix}$  contains four independently sampled regressor variables plus one intercept column:
  - $-\mathbf{x}_1 \stackrel{iid}{\sim} \mathcal{N}(5, 16),$  $-\mathbf{x}_2 \stackrel{iid}{\sim} \mathrm{Exp}(5),$  $- \mathbf{x}_3 \stackrel{iid}{\sim} \mathcal{U}([-2, 12]),$  $-\mathbf{x}_4 \stackrel{iid}{\sim} \mathrm{Ber}(0.3).$
- The design matrix  $\mathbf{Z} = \begin{pmatrix} \mathbf{1}_n & \mathbf{x}_1 & \mathbf{x}_2 & \mathbf{z}_3 \end{pmatrix}$  contains the additional regressor variable  $\mathbf{z}_3 \stackrel{iid}{\sim} t_{10}$ , which is independently sampled from all other columns.
- The true coefficient vectors are given by  $\boldsymbol{\beta} = \begin{pmatrix} \beta_0 & \beta_1 & \beta_2 & \beta_3 & \beta_4 \end{pmatrix}^T = \begin{pmatrix} 0 & -3 & -1 & -1 & 2 \end{pmatrix}^T$  and  $\boldsymbol{\gamma} = \begin{pmatrix} \gamma_0 & \gamma_1 & \gamma_2 & \gamma_3 \end{pmatrix}^T = \begin{pmatrix} 0 & 1 & 2 & 3 \end{pmatrix}^T$ .
- Three different specifications for the outcome distribution were chosen:
  - $-y_i \sim \mathcal{N}(\mu, \sigma^2),$
  - $-y_i \sim \mu + \left(\sigma \cdot \sqrt{\frac{3}{5}}\right) T$ , where  $T \sim t_5$ ,  $-y_i \sim \mu + \sigma \cdot U$ , where  $U \sim \mathcal{U}([0, 1])$ . In all cases, the outcome vectors are generated with the covariates on their original (unstandardized) scale.
- In order to isolate the impact of the different shapes of the three probability distributions from the effect of varying moment structures, the mean  $\mu = \mathbf{x}_i^T \boldsymbol{\beta}$  and the variance  $\sigma^2 = \exp(\mathbf{z}_i^T \boldsymbol{\gamma})^2$  are held constant across the models.
- All three models mcmc ridge(), mcmc() and lmls() are fitted with standardized covariates. The sample size is set to n = 50 and the result of both Bayesian samplers are based on 10.000 simulations.

### Section 3.5: Hyperparameters