# The asp21bridge Package

# 1 The asp21bridge Package

This chapter introduces numerous facets of the asp21bridge package.

Section 1.1 explains how to use and combine the functions that are contained in the package most efficiently.

Section 1.2 focuses on the implementation of the Markov Chain Monte Carlo Sampler with Ridge Penalty, links the source code to the mathematical model from chapter ?? 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 1.3.

### 1.1 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 1.2.1.
- 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.

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

As explained in section 1.2.1, 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 ??), 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 ??, 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>
```

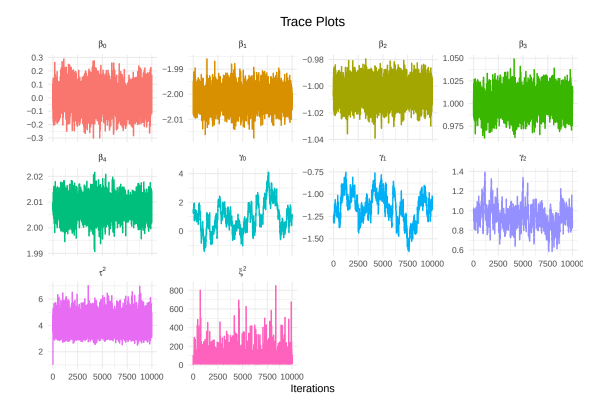


Figure 1: Trace Plots for all Coefficients, data: toy data

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
```

### 1.1.2 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

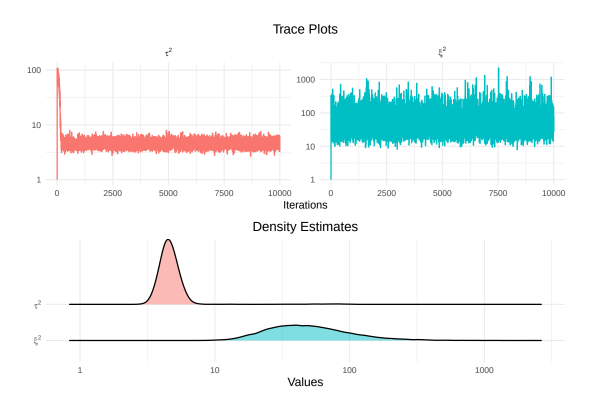


Figure 2: Trace and Density Plots for the Prior Variances, data: abdom

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>
```

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
)
```

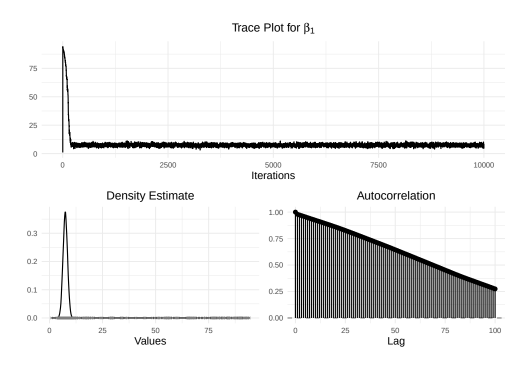


Figure 3: Diagnostic Plots for  $\beta_1$  before thinning, data: abdom

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 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.

#### 1.1.3 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

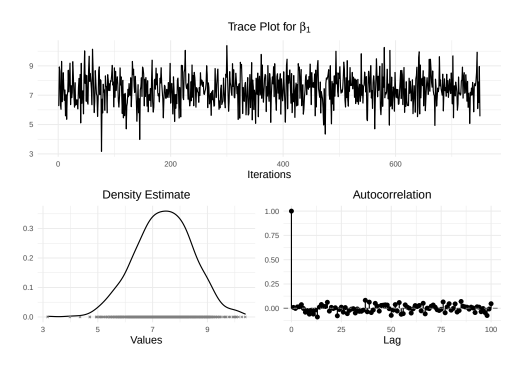


Figure 4: Diagnostic Plots for  $\beta_1$  after thinning, data: abdom

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.
                       Median
                                  Mean
                                        3rd Qu.
                                                    Max.
  -3.29800 -0.38660 0.11770 -0.01354
                                        0.57410
                                                  2.54600
##
##
## Location coefficients (identity link function):
##
                Mean
                           2.5%
                                       50%
## beta 0 0.0003161 -0.1488290 -0.0013337
## beta_1 -2.0015679 -2.0082464 -2.0015799 -1.995
## beta_2 -1.0044943 -1.0179012 -1.0044382 -0.991
## beta_3 1.0014607 0.9802516
                                            1.022
                                 1.0017075
## beta_4 2.0080638 2.0015876
                                2.0081264
##
## Scale coefficients (log link function):
                      2.5%
                               50% 97.5%
##
              Mean
## 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 \leftarrow c(0, -2, -1, 1, 2)
true_gamma \leftarrow 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
                       -2.00
                                     -2
## 2 beta_1
                                                      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
                                      0
## 6 gamma_0
                        0.937
                                                      0.967
## 7 gamma_1
                       -1.15
                                     -1
                                                      0.163
## 8 gamma 2
                                                      0.109
                        0.933
                                      1
```

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 1.1.2, 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 Plot 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 ?? after explaining some of the internal implementations in section 1.2.

# 1.2 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 1.2.1 explains both the overall structure of the mcmc\_ridge() function and the Gibbs Sampler that is responsible for inducing the Ridge penalty. Section 1.2.2 is dedicated to the integrated Metropolis-Hastings step, which is used for sampling  $\gamma$  and, if desired,  $\beta$  as well.

### 1.2.1 Iterative Parameter Sampling

The mcmc\_ridge() function is a Markov chain Monte Carlo (MCMC) algorithm allowing to simulate from parameters of a location scale regression model or from matrices and parameters that are assigned by hand. Since multiple input types are allowed, the mcmc\_ridge() function does not only contain the simulation process itself, but also input and output modalities, especially in a sense of defensive programming.

# **Overall Structure:**

As broached in section 1.1.1, the mcmc\_ridge() can take two different valid input types for simulating data. Next to them, however, input parameters for number of simulations, starting values for the simulation process, strengths of ridge penalties and allowance as well as proposal variances for simulating with a *Metropolis-Hastings algorithm* (see section 1.2.2 for the latter). Their default settings are discussed in section 1.3.2. The following enumeration gives a detailed overview of each parameter:

- m: Model object containing a beta and gamma predictor, as well as the according model matrices. The model is used in second stage, if one of X, Z, y, beta\_start or gamma\_start is not given. A model object of the S3 Class lmls from the lmls() can be used here smoothly. Default: NULL.
- X: Matrix containing the data according to the beta\_start coefficient. Default: NULL.
- Z: Matrix containing the data according to the gamma\_start coefficient. Default: NULL.
- v: Response vector;  $y = X * \beta + Z * \gamma$ . Default: NULL.
- num\_sim: Number of simulations. Default: 1000.
- beta\_start: Starting vector for simulation, where  $\beta$  is a linear predictor for the mean (i.e. the location), that is normally distributed. Default: NULL.

- gamma\_start: Starting vector for simulation, where  $\gamma$  is a linear predictor for the standard deviation (i.e. the scale), that is normally distributed. Default: NULL.
- tau\_start: Starting value for the variance of the normal distribution of the beta parameter, where tau\_start is inverse gamma (IG) distributed. Regularization parameter in a Bayesian ridge setting. Default: 1.
- xi\_start: Starting value for the variance of the normal distribution of the gamma parameter, where xi\_start is inverse gamma (IG) distributed. Regularization parameter in a Bayesian ridge setting, Default: 1.
- a\_tau: Fix shape parameter of the IG distribution of tau\_start. Default: 50.
- b\_tau: Fix scale parameter of the IG distribution of tau\_start. Default: 200.
- a\_xi: Fix shape parameter of the IG distribution of xi\_start. Default: 2.
- b\_xi: Fix scale parameter of the IG distribution of xi\_start. Default: 100.
- prop\_var\_scale: Variance of proposal distribution for γ sampling by a Metropolis-Hastings algorithm.
   Default: 3.
- mh\_location: If TRUE, location parameter  $\beta$  is sampled with *Metropolis-Hastings algorithm*. Can be used with every kind of model input. Default: FALSE.
- prop\_var\_loc: Variance of proposal distribution for  $\beta$  sampling. Default: 200.

### 1.2.2 Metropolis Hastings Step

Because the Full Conditional of  $\gamma$  cannot be assigned to a known type of distribution (see section ??), the MCMC update for  $\gamma$  cannot be sampled directly but requires an Metropolis Hastings step. Remind that the Full Conditional of  $\gamma$  is given by the following expression:

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

For numerical reasons, we used the Log-Full Conditionals for the Metropolis Hastings algorithm. The algorithm can be summarized by the following steps:

1. Compute the covariance matrix

$$\Sigma = diag(1/colMeans(Z^2)) * prop\_var\_scale/(n * len(\gamma)).$$

and sample a proposal for  $\gamma$  as

$$\gamma_{prop} \sim N(\gamma_{curr}, \Sigma).$$

with  $\gamma_{curr}$  being the current  $\gamma$  in the Markov chain. Note that the term  $Z^2$  squares all elements of Z elementwise, n denotes the length of the vector y. Furthermore, the function colMeans returns a vector with the column means of a matrix, and the function len simply returns the length of a vector.

2. Compute the Log-Full Conditionals for  $\gamma_{curr}$  and  $\gamma_{prop}$ . In the following, they are denoted as  $LFC_{curr}$  and  $LFC_{prop}$ . Then compute the log-acceptance probability as

$$log\_acc\_prob = min(0, LFC_{prop} - LFC_{curr}).$$

The log-acceptance probability is restricted to be greater than zero to ensure that the acceptance probability does not exceed the value of 1.

3. The acceptance probability is simply computed as

$$acc \ prob = \exp(log \ acc \ prob).$$

4. Simulate a uniform distribution on the interval (0,1). The outcome is denoted as U. Now we consider two different cases: If  $U < acc\_prob$  is true, the Markov chain accepts the proposal and jumps to  $\gamma_{prop}$ . Otherwise, the Markov chain stays at  $\gamma_{curr}$ .

The formula for  $\Sigma$  in step 1 was chosen after a lot simulations in order to make the acceptance probabilities as much independent from Z, y and n as possible. The parameter prop\_var\_scale was created for the mcmc\_ridge() function to control the magnitude of the acceptance probabilities. For the default of prop\_var\_scale = 3, the Metropolis Hastings algorithm should lead to acceptance probabilities between 25% and 50%. If the parameter is increased, the magnitude of the jumps also increases, but the acceptance probability becomes reduced, such that the Markov chain for  $\gamma$  might stay at the same point for a very long time. If the parameter is decreased, the acceptance probability increases. But due to a small magnitude of the jumps within the Markov chain, the Markov chain might look like a random walk and does not fit the Posterior distribution of  $\gamma$  very well.

The Metropolis Hastings algorithm for  $\beta$  works in a similar way but is of minor importance, since the  $\beta$  updates could be also sampled directly from a known distribution.

# 1.3 Package Development

This section is dedicated to more niche aspects, that come along with the package development process. Part 1.3.1 focuses on more formal components of the package, whereas some of the ideas behind the asp21bridge functions are discussed in section 1.3.2.

### 1.3.1 Code Coverage

Arguably the most important component of a package for new users is the **documentation**. All exported functions of the **asp21bridge** package as well as the **toy\_data** 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 1.1 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.

### 1.3.2 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 1.1, 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 1.1 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 m</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 1.1, 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 1.1.