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multibridge: An R Package To Evaluate Multinomial Order Constraints	
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9 Abstract

The **multibridge** package has been developed to efficiently compute Bayes factors for binomial and multinomial models, that feature inequality constraints, equality constraints, free parameters and mixtures between them. By using the bridge sampling algorithm to compute the Bayes factor, **multibridge** facilitates the evaluation of large models with many constraints and models with small parameter spaces. The package was developed in the R programming language and is freely available from the Comprehensive R Archive Network (CRAN). We illustrate the functions based on two empirical examples.

multibridge: An R Package To Evaluate Multinomial Order Constraints

18 Introduction

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We present **multibridge**, an R package to evaluate informed hypotheses in
multinomial models and models featuring independent binomials using Bayesian inference.
This package allows users to specify constraints on the underlying category proportions
including inequality constraints, equality constraints, free parameters and mixtures between
them. This package is available from the Comprehensive R Archive Network (CRAN) at
https://CRAN.R-project.org/package=multibridge. Here we introduce the methodology
used to evaluate informed hypotheses on categorical variables and show how to use the
implementations provided in **multibridge** through fully reproducible examples.

The most common way to analyze categorical variables is to test whether the 27 underlying category proportions are exactly equal or whether they are fixed and follow a 28 predicted pattern (what is generally known as either chi-square goodness of fit tests, or 29 binomial or multinomial tests). These null hypotheses are then tested against an encompassing hypothesis which places no constraints on the category proportions. This 31 analytic strategy has been criticized, since the null hypotheses might reflect an unrealistic expectation about the real world and the encompassing hypothesis is too uninformative 33 (Hoijtink, Klugkist, & Boelen, 2008). In addition, this strategy cannot or only insufficiently test the specific predictions derived by scientific theories researchers are interested in. A simple example for this are theories that predict ordinal relations among the underlying category proportions, such as increasing or decreasing trends. For instance, to check for irregularities in audit data, one could test whether the leading digits in the data are distributed according to an expected Benford distribution or whether they deviate from it, for example, by showing a general decreasing trend. Here, the Benford distribution can be tested with standard methods, however, the general decreasing trend cannot be tested, since

we cannot derive fixed underlying proportions for the leading digits. Theories can also generate more complex predictions, including ones that feature combinations of equality and 43 inequality constraints, as well as predictions that let some category proportions free to vary. In the following, we will denote such predictions as informed hypotheses, since they "add theoretical expectations to the traditional alternative hypothesis, thus making it more informative' (Hoijtink et al., 2008, p. 2). Such an informed hypothesis was expressed, for instance, by Nuijten, Hartgerink, Assen, Epskamp, and Wicherts (2016) who studied the prevalence of statistical reporting errors in articles published in different areas of psychological science. Nuijten et al. (2016) hypothesized that articles published in social psychology journals would have higher error rates than articles published in other psychological journals while not expressing expectations about the error rate distribution among the other journals. Here again no fixed proportions can be derived, such that this hypothesis cannot not be evaluated using standard tests. Generally, if researchers can utilize statistical methods for testing informed hypotheses, they are able to specify hypotheses that relate more closely to their theories.

In the Bayesian framework, researchers can compare models that instantiate the
hypotheses of interest by means of Bayes factors (Jeffreys, 1935; Kass & Raftery, 1995). To
compute Bayes factors for informed hypotheses several R packages are already available. For
instance, with the package multinomineq (Heck & Davis-Stober, 2019) users can specify
inequality constrained hypotheses but also more general linear inequality constraints for
multinomial models as well as models that feature independent binomials. The BAIN
package (Gu, Hoijtink, Mulder, & Rosseel, 2019) allows for the evaluation of inequality
constraints in structural equation models. Outside of R, the Fortran 90 program BIEMS
(Mulder, Hoijtink, Leeuw, & others, 2012) allows for the evaluation of order constraints for
multivariate linear models such as MANOVA, repeated measures, and multivariate
regression. All these packages rely on one of two methods to approximate order constrained
Bayes factors: the encompassing prior approach (Gu, Mulder, Deković, & Hoijtink, 2014;

Hoijtink, 2011; Hoijtink et al., 2008; Klugkist, Kato, & Hoijtink, 2005) and the conditioning method (Mulder, 2014, 2016; Mulder et al., 2009). However, even though these methods are currently widely used, they are known to become increasingly unreliable and inefficient as the number of constraints increases or when the parameter space of the constrained model is small (Sarafoglou et al., 2020).

In contrast to these available packages, multibridge uses a bridge sampling routine that enables users to compute Bayes factors for informed hypotheses more reliably and 75 efficiently (Bennett, 1976; Meng & Wong, 1996; Sarafoglou et al., 2020). The workhorse for this analysis, the bridge sampling algorithm, constitutes a special case of the algorithm implemented in the R package bridgesampling (Gronau, Singmann, & Wagenmakers, 2020). The **bridgesampling** package, allows users to estimate the marginal likelihood for a wide variety of models, including models implemented in Stan (Stan Development Team, 2020). However, the algorithm implemented in **bridgesampling** is not suitable for models 81 that include constraints on probability vectors and hence is unsuitable for the analysis of 82 categorical data. Therefore, in **multibridge**, we tailored the bridge sampling algorithm such that it accommodates the specification of informed hypotheses on probability vectors. The 84 package then produces an estimate for the Bayes factor in favor of or against the informed hypothesis. The resulting Bayes factor compares the evidence for the informed hypotheses to the encompassing hypothesis that imposes no constraints on the underlying category 87 proportions. Alternatively, the informed hypothesis can be tested against the null hypothesis that all underlying category proportions are exactly equal. Given this result, users can then either receive a visualization of the posterior parameter estimates under the encompassing hypothesis using the plot-method, or get more detailed information on how the Bayes factor is composed using the summary-method. For hypotheses that include mixtures between equality and inequality constrained hypotheses the bayes_factor method shows the conditional Bayes factor for the inequality constraints given the equality constraints and a Bayes factor for the equality constraints. The general workflow of **multibridge** is illustrated

in Figure 1. Table 1 summarizes all S3 methods currently available in **multibridge**.

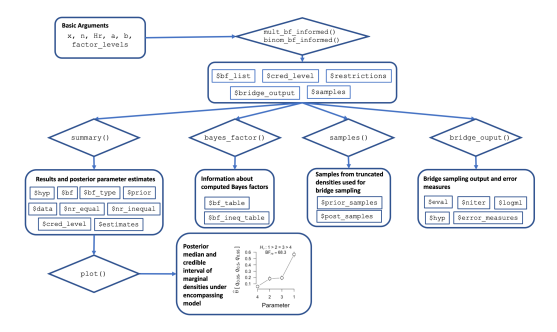


Figure 1. The multibridge workflow. The user specifies the data values (x and n for binomial models and x for multinomial models, respectively), the informed hypothesis (Hr), the α and β parameters of the Binomial prior distributions (a and b) or the concentration parameters for the Dirichlet prior distribution (a), respectively, and the factor levels (factor_levels). The functions mult_bf_informed and binom_bf_informed then produce an estimate for the Bayes factor of the informed hypothesis versus the encompassing or the null hypothesis. Based on these results different S3 methods can be used to get more detailed information on the individual components of the analysis (e.g., summary, bayes_factor), and parameter estimates of the encompassing distribution (plot).

The remainder of this article is organized as follows: In the methods section, we
describe the Bayes factor identity for informed hypotheses in binomial and multinomial
models, and present the bridge sampling routine implemented in the **multibridge** package
including details of the necessary transformations required for this routine. In Section 3, we
will schematically introduce the most relevant functions in **multibridge** and their
arguments. Section 4 illustrates how to use the **multibridge** package to estimate

parameters, and compute Bayes factors using two examples.

104 Methods

In this section we formalize multinomial models and models that feature independent 105 binomial probabilities as we have implemented them in **multibridge**. In the multinomial 106 model, we assume that the vector of observations \mathbf{x} in the K categories follow a multinomial 107 distribution in which the parameters of interest, θ , represent the underlying category 108 proportions. Since we assume a dependence between the K categories, the vector of 109 probability parameters is sum-to-one constrained, such that $\sum_{k=1}^{K} (\theta_1, \dots, \theta_K) = 1$. 110 Therefore, a suitable choice for a prior distribution for θ is the Dirichlet distribution with 111 concentration parameters α : 112

$$x_1, \dots, x_K \sim \text{Multinomial}(\sum_{k=1}^K x_k, \theta_1, \dots, \theta_K)$$
 (1)

$$\theta_1, \cdots, \theta_K \sim \text{Dirichlet}(\alpha_1, \cdots, \alpha_K),$$
 (2)

where α can be interpreted as vector of a priori category counts. Since the 113 multinomial model constitutes a generalization of the binomial model (for $K \geq 2$), the 114 formalization of a model that features independent binomial probabilities is very similar. In 115 the binomial model, we assume that the elements in the vector of successes \mathbf{x} and the 116 elements in the vector of total number of observations \mathbf{n} in the K categories follow independent binomial distributions. As in the multinomial model, the parameter vector of 118 the binomial success probabilities θ contains the underlying category proportions, however, 119 in this model we assume that categories are independent which removes the sum-to-one 120 constraint. Therefore, a suitable choice for a prior distribution for θ is a vector of 121 independent beta distributions with parameters α and β : 122

$$x_1 \cdots x_K \sim \prod_{k=1}^K \text{Binomial}(\theta_k, n_k)$$
 (3)
 $\theta_1 \cdots \theta_K \sim \prod_{k=1}^K \text{Beta}(\alpha_k, \beta_k),$ (4)

$$\theta_1 \cdots \theta_K \sim \prod_{k=1}^K \text{Beta}(\alpha_k, \beta_k),$$
 (4)

where α can be interpreted as vector of a priori successes that observations fall withing the various categories and β can be interpreted as vector of a priori failures.

Bayes factor

With **multibridge** package, it is possible to collect evidence for informed hypotheses 126 on a parameter vector $\boldsymbol{\theta}$ by means of the Bayes factor. Bayes factors compare the relative 127 evidence of two hypotheses in the light of the data. It is defined as the ratio of marginal 128 likelihoods of the respective hypotheses. For instance, the Bayes factor for the informed 129 hypothesis versus a hypothesis that lets all parameters free to vary is defined as:

$$\mathrm{BF}_{re} = \frac{\overbrace{p(\mathbf{x} \mid \mathcal{H}_r)}^{\mathrm{Marginal likelihood}}_{\mathbf{p}(\mathbf{x} \mid \mathcal{H}_e)}^{\mathbf{p}(\mathbf{x} \mid \mathcal{H}_r)},}{\underbrace{p(\mathbf{x} \mid \mathcal{H}_e)}_{\mathrm{Marginal likelihood}}^{\mathrm{Marginal likelihood}}_{\mathbf{under} \ \mathcal{H}_e},}$$

where the subscript r denotes the informed hypothesis and e denotes a hypothesis 131 which predicts that all parameters free to vary. In **multibridge** we use two different 132 methods to compute Bayes factors, one method evaluates hypotheses that feature equality constraints on θ and one method evaluates hypotheses that feature inequality constraints on $\boldsymbol{\theta}$. Both methods will be outlined below. In cases where informed hypotheses feature 135 mixtures between inequality and equality constraints, we compute the corresponding Bayes 136 factor BF_{re} , by multiplying the individual Bayes factors for both constrait types with each 137 other: 138

$$BF_{re} = BF_{1e} \times BF_{2e} \mid BF_{1e}$$

where the subscript 1 denotes the hypothesis that only features equality constraints and the subscript 2 denotes the hypothesis that only features inequality constraints. A Bayes factor for mixtures thus factors into a Bayes factor for the equality constraints, BF_{1e} , and a conditional Bayes factor for the inequality constraints given the equality constraints $BF_{2e} \mid BF_{1e}$ (for the proof, see Sarafoglou et al., 2020).

144 The Bayes Factor For Equality Constraints

The Bayes factor for the equality constraints can be computed analytically both for binomial and multinomial models. For binomial models, the function $binom_bf_equality$ is available to compute BF_{0e} . Assuming that the first i binomial probabilities in a model are equality constrained, the Bayes factor is defined as:

$$BF_{0e} = \frac{\prod_{i < k} B(\alpha_i, \beta_i)}{\prod_{i < k} B(\alpha_i + x_i, \beta_i + n_i - x_i)} \times \frac{B(\alpha_+ + x_+ - i + 1, \beta_+ + n_+ - x_+ - i + 1)}{B(\alpha_+ - i + 1, -i + 1)}$$

where B() denotes the beta function and $\alpha_+ = \sum_{i < k} \alpha_i$, $\beta_+ = \sum_{i < k} \beta_i$, $x_+ = \sum_{i < k} x_i$ and $n_+ = \sum_{i < k} n_i$. The latter factor introduces a correction for marginalizing which stems from the change in degrees of freedom, when we collapse i equality constraint parameters: For i collapsed categories, i-1 degrees of freedom are lost which are subtracted from the prior parameters in the corresponding Binomial distribution.

For multinomial models, the function multBayes_bf_equality is available. Assuming again that the first i category probabilities in a model are equality constraint, the Bayes factor $BF_{=e}$ is defined as:

$$BF_{0e} = \frac{B(\boldsymbol{\alpha} + \mathbf{x})}{B(\boldsymbol{\alpha})} \left(\frac{1}{i}\right)^{\sum_{i < k} x_i} \frac{B(\sum_{i < k} \alpha_i - i + 1, \alpha_k, \dots, \alpha_K)}{B(\sum_{i < k} \alpha_i + x_i - i + 1, \alpha_k + x_k, \dots, \alpha_K + x_K)},$$

50 The Bayes Factor For Inequality Constraints

To approximate the Bayes factor for informed hypotheses, Klugkist et al. (2005)
derived the following identity:

$$BF_{re} = \frac{p(\boldsymbol{\theta} \in \mathcal{R}_r \mid \mathbf{x}, \mathcal{H}_e)}{p(\boldsymbol{\theta} \in \mathcal{R}_r \mid \mathcal{H}_e)}.$$
Proportion of prior parameter space consistent with the restriction
$$(5)$$

Recently, Sarafoglou et al. (2020) showed that the Bayes factor BF_{re} can also be interpreted as ratio of two marginal likelihoods:

$$BF_{re} = \frac{\overbrace{p(\boldsymbol{\theta} \in \mathcal{R}_r \mid \mathbf{x}, \mathcal{H}_e)}^{\text{Marginal likelihood of}}}{\underbrace{p(\boldsymbol{\theta} \in \mathcal{R}_r \mid \mathbf{x}, \mathcal{H}_e)}_{\text{Marginal likelihood of}}}.$$
(6)

In this identity, $p(\theta \in \mathcal{R}_r \mid \mathbf{x}, \mathcal{H}_e)$ denotes the marginal likelihood of the constrained 155 posterior distribution and $p(\theta \in \mathcal{R}_r \mid \mathcal{H}_e)$ denotes the marginal likelihood of the constrained 156 prior distribution. Even though both identities are mathematically equivalent, the methods 157 to estimate these identities are very different. In the first case, for instance, the number of 158 samples from the encompassing distribution in accordance with the inequality constrained hypothesis, serve as an estimate for the proportion of prior parameter space consistent with the restriction. On the flip side, however, this means that the accuracy of this estimate is 161 strongly dependent on the number of the constrained parameters in the model and the size 162 of the constrained parameter space. That is, as the constraints become stronger, the 163 constrained parameter space decreases. As a result it becomes less likely that draws from the 164

encompassing distribution will fall into the constrained region, so that in some cases the estimation of the Bayes factor becomes practically impossible (Sarafoglou et al., 2020).

However, when we interpret the Bayes factor BF_{re} as ratio of marginal likelihoods and 167 we are able to sample from the constrained prior and posterior distributions, we can utilize 168 numerical sampling methods such as bridge sampling to obtain the estimates. Crucially, in 169 this approach, it does not matter how small the constrained parameter space is in proportion to the encompassing density. This gives the method a decisive advantage over the 171 encompassing prior approach in terms of accuracy and efficiency especially (1) when binomial and multinomial models with relatively high number of categories (i.e., K > 10) are 173 evaluated and (2) when relatively little posterior mass falls in the constrained parameter 174 space. 175

176 The Bridge Sampling Method

Bridge sampling is a method to estimate the ratio of two marginal likelihoods (Bennett, 177 1976; Meng & Wong, 1996). In **multibridge**, we are using bridge sampling to estimate the 178 identity presented in Equation 6. But instead of estimating the ratio of marginal likelihoods 179 directly, we implemented a version of bridge sampling that estimates one marginal likelihood 180 at the time. This approach has the benefit that it increases the accuracy of the method 181 without considerably increasing its computational efficiency (Overstall & Forster, 2010). 182 Specifically, we subsequently estimate the marginal likelihood for the constrained prior 183 distribution and the marginal likelihood of the constrained posterior distribution. 184

When applying this modified version of the bridge sampling method, we estimate each marginal likelihood by means of a so-called proposal distribution. In **multibridge** this proposal distribution is the multivariate normal distribution. To estimate the marginal likelihood, bridge sampling only requires samples from the distribution of interest—the

so-called target distribution—and samples from the proposal distribution.

Samples from the target distribution—that is the constrained prior and posterior
Dirichlet distribution for multinomial models and constrained prior and posterior beta
distributions for binomial models—are drawn through the Gibbs sampling algorithms
proposed by Damien and Walker (2001). For binomial models, we apply the suggested Gibbs
sampling algorithm for constrained beta distributions. In the case of the multinomial models,
we apply an algorithm that simulates values from constrained Gamma distributions which
are then transformed into Dirichlet random variables (for details, see Appendix C in
Sarafoglou et al., 2020). To sample efficiently from these distributions, we implemented a
C++ routine for this algorithm in the package.

Samples from the proposal distribution can be generated using the standard 199 rmvnorm-function from the R package stats. The vector of means and the covariance matrix of this distribution are derived from one part of the samples of the probit transformed target 201 distribution. The reason for this approach is that the efficiency of the bridge sampling 202 method is optimal only if the target and proposal distribution operate on the same 203 parameter space and have sufficient overlap. We therefore probit transform the samples of 204 the constrained distributions to move the samples from the probability space to the entire 205 real line. Subsequently, we use half of these draws to construct the proposal distribution 206 using the method of moments. Details on the probit transformations are provided in the 207 appendix. Thus, for the marginal likelihood of the constrained prior distribution, the 208 modified bridge sampling identity is then defined as: 200

$$p(\boldsymbol{\theta} \in \mathcal{R}_r \mid \mathcal{H}_e) = \frac{\mathbb{E}_{g(\boldsymbol{\theta})} \left(p(\boldsymbol{\theta} \mid \mathcal{H}_e) \mathbb{I}(\boldsymbol{\theta} \in \mathcal{R}_r) h(\boldsymbol{\theta}) \right)}{\mathbb{E}_{\text{prior}} \left(g(\boldsymbol{\theta}) h(\boldsymbol{\theta}) \right)}, \tag{7}$$

where the term $h(\boldsymbol{\theta})$ refers to the bridge function proposed by Meng and Wong (1996) and $g(\boldsymbol{\theta})$ refers to the proposal distribution. The numerator evaluates the unnormalized density

for the constrained prior distribution with samples from the proposal distribution. The
denominator evaluates the normalized proposal distribution with samples from the
constrained prior distribution. Using this identity, we receive the bridge sampling estimator
for the marginal likelihood of the constrained prior distribution by applying the iterative
scheme proposed by Meng and Wong (1996):

$$\hat{p}(\boldsymbol{\theta} \in \mathcal{R}_r \mid \mathcal{H}_e)^{(t+1)} \approx \frac{\frac{1}{N_2} \sum_{m=1}^{N_2} \frac{\ell_{2,m}}{s_1 \ell_{2,m} + s_2 p(\tilde{\boldsymbol{\theta}}_m \in \mathcal{R}_r \mid \mathcal{H}_e)^{(t)}}}{\frac{1}{N_1} \sum_{n=1}^{N_1} \frac{1}{s_1 \ell_{1,n} + s_2 p(\boldsymbol{\theta}_n^* \in \mathcal{R}_r \mid \mathcal{H}_e)^{(t)}}},$$

where N_1 denotes the number of samples drawn from the constrained distribution, that is, $\theta^* \sim p(\theta \mid \mathcal{H}_r), N_2 \text{ denotes the number of samples drawn from the proposal distribution, that}$ is $\tilde{\boldsymbol{\theta}} \sim g(\boldsymbol{\theta}), s_1 = \frac{N_1}{N_2 + N_1}$, and $s_2 = \frac{N_2}{N_2 + N_1}$. The quantities $\ell_{1,n}$ and $\ell_{2,m}$ are defined as follows:

$$\ell_{1,n} = \frac{q_{1,1}}{q_{1,2}} = \frac{p(\boldsymbol{\theta_n^*} \mid \mathcal{H}_e)\mathbb{I}(\boldsymbol{\theta_n^*} \in \mathcal{R}_r)}{g(\boldsymbol{\xi_n^*})},\tag{8}$$

$$\ell_{2,m} = \frac{q_{2,1}}{q_{2,2}} = \frac{p(\tilde{\boldsymbol{\theta}}_m \mid \mathcal{H}_e) \mathbb{I}(\tilde{\boldsymbol{\theta}}_m \in \mathcal{R}_r)}{g(\tilde{\boldsymbol{\xi}}_m)},\tag{9}$$

where $\boldsymbol{\xi_n}^* = \Phi^{-1}\left(\frac{\boldsymbol{\theta_n^*} - 1}{\mathbf{u} - \mathbf{l}}\right)$, and $\tilde{\boldsymbol{\theta}}_m = ((\mathbf{u} - \mathbf{l})\Phi(\tilde{\boldsymbol{\xi}}_m) + \mathbf{l}) |J|)$. The quantity $q_{1,1}$ refers to the evaluations of the constrained distribution for constrained samples and $q_{1,2}$ refers to the proposal evaluations for constrained samples, respectively. The quantities $q_{2,1}$ refers to evaluations of the constrained distribution for samples from the proposal and $q_{2,2}$ refers to the proposal evaluations for samples from the proposal, respectively. Note that the quantities $\ell_{1,n}$ and $\ell_{2,m}$ have been adjusted to account for the necessary parameter transformations to create overlap between the constrained distributions and the proposal distribution. **multibridge** runs the iterative scheme until the tolerance criterion suggested

by Gronau et al. (2017) is reached, that is:

$$\frac{\mid \hat{p}(\boldsymbol{\theta} \in \mathcal{R}_r \mid \mathcal{H}_e)^{(t+1)} - \hat{p}(\boldsymbol{\theta} \in \mathcal{R}_r \mid \mathcal{H}_e)^{(t)} \mid}{\hat{p}(\boldsymbol{\theta} \in \mathcal{R}_r \mid \mathcal{H}_e)^{(t+1)}} \leq 10^{-10}.$$

The bridge sampling estimate for the log marginal likelihood of the constrained distribution and its associate relative mean square error, the number of iterations, and the quantities $q_{1,2}$, $q_{1,2}$, $q_{1,2}$, and $q_{1,2}$ are included in the standard output in **multibridge**. The function to compute the relative mean square error was taken from the R package bridgesampling.

Is this important enough to mention it here?

Not sure where to include it otherwise

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Usage and Examples

The **multibridge** package can be installed from the Comprehensive R Archive
Network (CRAN) at https://CRAN.R-project.org/package=multibridge:

```
install.packages('multibridge')
library('multibridge')
```

A list of all currently available functions and datasets is given in Table 3. Additional
examples are available as vignettes (see https://cran.r-project.org/package=multibridge, or
vignette(package = "multibridge")). The two core functions of multibridge—the
mult_bf_informed-function and the binom_bf_informed-function—can be illustrated
schematically as follows:

```
mult_bf_informed(x, Hr, a factor_levels)
binom_bf_informed(x, n, Hr, a, b, factor_levels)
```

The basic required arguments for these functions are listed in Table 2. In the following,
we will outline two examples on how to use **multibridge** to compare an informed hypothesis
to a null or encompassing hypothesis. In addition, the first example shows how two informed
hypotheses can be compared to each other.

Example 1: Applying A Benford Test to Greek Fiscal Data

The first digit phenomenon, otherwise known as Benford's law (Benford, 1938;
Newcomb, 1881) states that the expected proportion of leading digits in empirical data can
be formalized as follows: for any given leading digit $d, d = (1, \dots, 9)$ the expected proportion
is approximately equal to

$$\mathbb{E}_{\theta_d} = \log_{10}((d+1)/d).$$

This means that a number in a empirical dataset has leading digit 1 in 30.1% of the cases, and leading digit 2 in 17.61% of the cases; leading digit 9 is the least frequent digit with an expected proportion of only 4.58% (see Table 4 for an overview of the expected proportions). Empirical data includes, for instance, data on population sizes, death rates, baseball statistics, atomic weights of elements, and physical constants (Benford, 1938). In contrast, 248 generated data, such as telephone numbers, do in general not obey Benford's law (Hill, 1995). 249 Since Benford's law proved to be highly suitable to discriminate between empirical data and 250 generated data, a so-called Benford test can be used to check whether certain observed 251 frequencies of first digits, obey Benford's law and therefore can be considered an empirical 252 dataset. Benford's tests are used in fields like accounting and auditing to check for 253 indications for poor data quality, for instance, in fiscal statements [for an overview,] see e.g., 254 Durtschi, Hillison, and Pacini (2004), Nigrini and Mittermaier (1997), Nigrini (2012). Data 255 that do not pass the Benford test, should raise audit risk concerns, meaning that it is 256 recommended that the data undergo additional follow-up checks (Nigrini, 2019). 257

In the following, we discuss three possible Bayesian adaptations of the Benford's test.

258

In a first scenario we simply conduct Bayesian multinomial test in which we test the 259 point-null hypothesis \mathcal{H}_0 which predicts a Benford distribution against the encompassing 260 hypothesis \mathcal{H}_e which leaves all model parameters free to vary. Testing against the 261 encompassing hypothesis is considered standard practice, yet, it leads to an unfair 262 comparison to the detriment of the null hypothesis. In general, if we are dealing with a 263 high-dimensional parameter space and the competing hypotheses differ largely in their 264 complexity, the Bayes factor generally favors the less complex hypothesis (i.e., \mathcal{H}_e) even if 265 the data follow the predicted trend of the more complex hypothesis considerably well. In a 266 second scenario we therefore test the null hypothesis against an alternative hypothesis, 267 denoted as \mathcal{H}_{r1} , which predicts a decreasing trend in the proportions of leading digits. The 268 hypothesis \mathcal{H}_{r1} implies considerably more constraints than \mathcal{H}_e and is a suitable choice if our 269 primary goal is to distinguish whether data comply with Benford's law or whether the data only follow a similar trend. In a third scenario, where the main goal is to identify fabricated 271 data, we could test the null hypothesis against an hypothesis, which predicts a trend that is characteristic for manipulated data. This hypothesis, which we denote as \mathcal{H}_{r2} , could be 273 derived from empirical research on fraud or be based on observed patterns from former fraud 274 cases. For instance, Hill (1988) instructed students to produce a series of random numbers; 275 in the resulting data the proportion of the leading digit 1 occurred most often and the digits 276 8 and 9 occurred least often which is consistent with the general pattern of Benford's law. 277 However, the proportion for the remaining leading digits were approximately equal. We do 278 want to note that the predicted distribution derived from Hill (1988) is not currently used as 279 a test to detect fraud. However, for the sake of simplicity, if we assume that this pattern 280 could be an indication for fabricated auditing data, the Bayes factor could quantify the 281 evidence of whether the proportion of first digits resemble authentic or fabricated data. 282

Data and Hypothesis. The data we use to illustrate the computation of Bayes
factors were originally published by the European statistics agency "Eurostat" and served as
basis for reviewing the adherence to the Stability and Growth Pact of EU member states.

Rauch, Göttsche, Brähler, and Engel (2011) conducted a Benford test on data related to 286 budget deficit criteria, that is, public deficit, public dept and gross national products. This 287 data used for this example contains fiscal data from Greece related in the years between 1999 288 and 2010; a total of N = 1.497 numerical data were included in the analysis. We choose this 289 data, since the Greek government deficit and debt statistics states has been repeatedly 290 criticized by the European Commission in this timespan (European Commission, 2004, 2010). 291 In particular, the commission has accused the Greek statistical authorities to have 292 misreported deficit and debt statistics. For further details on the dataset see Rauch et al. 293 (2011). The observed proportions are displayed in Table 4, the figure displaying the observed 294 versus the expected proportions are displayed in Figure 2. 295

In this example, the parameter vector of the multinomial model, $\theta_1, \dots, \theta_K$, reflects the probabilities of a leading digit in the Greek fiscal data being a number from 1 to 9. Thus, we can formalize the discussed hypotheses as follows. The null hypothesis specifies that the proportions of first digits obeys Benford's law:

$$\mathcal{H}_0: \boldsymbol{\theta}_0 = (0.301, 0.176, 0.125, 0.097, 0.079, 0.067, 0.058, 0.051, 0.046).$$

We are testing the null hypothesis against the following alternative hypotheses:

$$\mathcal{H}_e: \boldsymbol{\theta} \sim \text{Dirichlet}(\boldsymbol{\alpha}),$$

$$\mathcal{H}_{r1}: \theta_1 > \theta_2 > \theta_3 > \theta_4 > \theta_5 > \theta_6 > \theta_7 > \theta_8 > \theta_9,$$

$$\mathcal{H}_{r2}: \theta_1 > (\theta_2 = \theta_3 = \theta_4 = \theta_5 = \theta_6 = \theta_7) > (\theta_8, \ \theta_9).$$

In cases, in which we are interested in computing two informed hypotheses with each other, we need to make use of the transitivity property of the Bayes factor. For instance, if we would like to compare the two informed hypotheses \mathcal{H}_{r1} and \mathcal{H}_{r2} with each other, we would first compute BF_{er1} and BF_{er2} and then yield BF_{r1r2} as follows:

$$BF_{r1e} \times BF_{er2} = BF_{r1r2}$$
.

Method. We can compare \mathcal{H}_0 and \mathcal{H}_e by means of a Bayesian multinomial test, that 304 is, we stipulate equality constraints on the entire parameter vector $\boldsymbol{\theta}$. The corresponding 305 Bayes factor is thus computationally straightforward; we can calculate BF_{0e} by applying the 306 function mult_bf_equality. To evaluate \mathcal{H}_0 , we only need to specify (1) a vector with 307 observed counts, (2) a vector with concentration parameters of the Dirichlet prior 308 distribution, and (3) the vector of predicted proportions. Since we have no specific 309 expectations about the distribution of leading digits in the Greek fiscal data, we set all 310 concentration parameters to one which corresponds to a uniform Dirichlet distribution. 311

```
# Observed counts
x <- c(509, 353, 177, 114, 77, 77, 53, 73, 64)
# Concentration parameters
a <- rep(1, 9)
# Expected proportions
p <- log10((1:9 + 1)/1:9)
# Execute the analysis
results_HO_He <- mult_bf_equality(x = x, a = a, p = p)</pre>
```

Since the hypotheses \mathcal{H}_{r1} and \mathcal{H}_{r2} contain inequality constraints, we use the function mult_bf_informed to compute the Bayes factor of the informed hypotheses to the encompassing hypothesis. In this function, we need to specify (1) a vector with observed counts, (2) the informed hypothesis \mathcal{H}_{r1} or \mathcal{H}_{r2} (e.g., as character vector), (3) a vector with concentration parameters of the Dirichlet prior distribution, and (4) labels for the categories of interest (i.e., leading digits):

```
# Labels for categories of interest
factor_levels <- 1:9
# Specifying the informed Hypothesis
Hr1 <- c('1 > 2 > 3 > 4 > 5 > 6 > 7 > 8 > 9')
```

```
318 ## BFType LogBF
319 ## 1 LogBF0e -17.67150
320 ## 2 LogBF0r1 -25.08832
321 ## 3 LogBF0r2 154.57114
```

As the evidence is extreme in all three cases, we report all Bayes factors on the log scale. The log Bayes factor $\log(\mathrm{BF}_{0e})$ suggests extreme evidence against the hypothesis that the first digits in the Greek fiscal data follow a Benford's distribution; $\log(\mathrm{BF}_{0e}) = -17.67$. The log Bayes factor $\log(\mathrm{BF}_{0r1})$ indicates extreme evidence in favor for a decreasing trend, $\log(\mathrm{BF}_{0r1}) = -25.09$. Even though the Bayes factor suggests extreme evidence against the hypothesis that the Greek fiscal data are an empirical dataset, there is no support the

hypothesis that the data are fabricated. The log Bayes factor $\log(BF_{0r1})$ indicates extreme evidence against \mathcal{H}_{r2} with $\log(BF_{0r2}) = 154.57$. The proportions of leading digits is best characterized by a monotonously decreasing trend, compared to all parameters varying freely ($\log(BF_{r1e}) = 7.42$), and compared to a distribution that one could expect from fabricated data ($\log(BF_{r1r2}) = 180$).

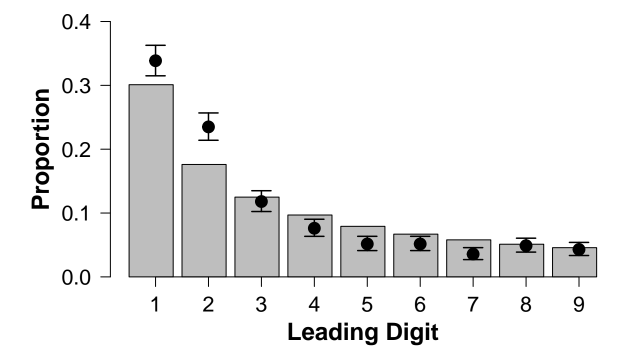


Figure 2. The bargraph displays the expected proportions of leading digits according to Benford's law. The black dots indicate for the actual fiscal statistics from Greece the posterior estimates for the proportion of leading digits and the corresponding 95% credible intervals based on the encompassing model. Only three out of nine estimates cover the expected proportions.

Discussion. In this example we tested the data quality of Greek fiscal data in the years 1999 to 2009 by conducting three variations of a Bayesian Benford test. More precisely, we evaluated the null hypothesis that β Greek fiscal data conform to Benfords law. We tested this hypothesis against three alternatives. The first alternative hypothesis, \mathcal{H}_e relaxed the constraints imposed by the null hypothesis and left all model parameters free to vary. The

second alternative hypothesis, \mathcal{H}_{r1} predicted a decreasing trend in the proportion of leading 338 digits. The third alternative hypothesis \mathcal{H}_{r2} predicted a trend that Hill (1988) observed 339 when humans tried to generate random numbers. Our result suggest that the leading digits 340 in the fiscal statistics do not follow a Benford distribution; in fact, we collected extreme 341 evidence against Benford's law compared to two out of three of the alternative hypotheses. 342 When comparing the alternative hypotheses directly to each other, the data show most 343 evidence in favor for a decreasing trend. A Benford test of fiscal statements can be a helpful 344 tool to detect poor data quality and suspicious numbers. In follow-up checks of these 345 numbers, it could then be examined for instance, whether financial statements were actually 346 materially misstated, for isntance, by rounding up or down numbers, avoiding certain 347 thresholds etc., Nigrini (2019). 348

Example 2: Prevalence of Statistical Reporting Errors

In any scientific article that uses null hypothesis significance testing, there is a chance 350 that the reported test statistic and degrees of freedom do not match the reported p-value. In 351 most cases this is because researchers copy the relevant test statistics by hand into their 352 articles and there are no automatic checks to detect these mistakes. Therefore, Epskamp and 353 Nuijten (2014) developed the R package statcheck, which only requires the PDF of a given 354 scientific article to detect these reporting errors automatically and efficiently. This package 355 allowed Nuijten et al. (2016) to get an overview about the prevalence of statistical reporting 356 errors in the field of psychology. In total, the authors investigated a sample of 30,717 articles (which translates to over a quarter of a million p-values) published in eight major psychological journals between 1985 to 2013: Developmental Psychology (DP), the Frontiers 359 in Psychology (FP), the Journal of Applied Psychology (JAP), the Journal of Consulting and 360 Clinical Psychology (JCCP), Journal of Experimental Psychology: General (JEPG), the 361 Journal of Personality and Social Psychology (JPSP), the Public Library of Science (PLoS), 362

363 Psychological Science (PS).

Besides the overall prevalence of statistical reporting errors across these journals, the 364 authors were interested whether there is a higher prevalence for reporting inconsistencies in 365 certain subfields in psychology compared to others. In this context the possibility was raised 366 that there exists a relationship between the prevalence for reporting inconsistencies and 367 questionable research practices. Specifically, the authors argued that besides honest mistakes when transferring the test statistics into the manuscript, statistical reporting error occur 369 when authors misreport p-values, for instance, by incorrectly rounding them down to or 370 below 0.05. Based on this assumption, Nuijten et al. (2016) predicted that the proportion of 371 statistical reporting errors should be highest in articles published in the Journal of Personality and Social Psychology (JPSP), compared to other journals, since researchers in 373 social psychology were shown to have the highest prevalence for questionable research 374 practices (John, Loewenstein, & Prelec, 2012). Specifically, John et al. (2012) found that 375 researchers from the area of social psychology assessed questionable research practices both 376 as more defensible and more applicable for their research compared to other research areas in 377 psychology. 378

Data and Hypothesis. Here, we are reusing the original data published by Nuijten et al. (2016), which we also distribute with the package multibridge under the name journals.

data(journals)

The hypothesis of interest, \mathcal{H}_r , formulated by Nuijten et al. (2016) states that the prevalence for statistical reporting errors for articles published in social psychology journals (i.e., JPSP) is higher than for articles published in other journals. Note that Nuijten et al. (2016) did not make use of inferential statistics since their sample included the entire population of articles from the eight flagship journals in psychology from 1985 to 2013. For

demonstration purposes, however, we will test the informed hypothesis stated by the authors.
We will test \mathcal{H}_r against the the null hypothesis \mathcal{H}_0 that all journals have the same prevalence
for statistical reporting errors. In this example, the parameter vector of the binomial success
probabilities, $\boldsymbol{\theta}$, reflects the probabilities of a statistical reporting error in one of the 8
journals. Thus, we can formalize the discussed hypotheses as follows:

$$\mathcal{H}_r: (\theta_{\mathrm{DP}}, \theta_{\mathrm{FP}}, \theta_{\mathrm{JAP}}, \theta_{\mathrm{JCCP}}, \theta_{\mathrm{JEPG}}, \theta_{\mathrm{PLoS}}, \theta_{\mathrm{PS}}) < \theta_{\mathrm{JPSP}}$$

$$\mathcal{H}_0: \theta_{\mathrm{DP}} = \theta_{\mathrm{FP}} = \dots = \theta_{\mathrm{JPSP}}.$$

Method. To compute the Bayes factor BF_{0r} we need to specify (1) a vector with observed successes (i.e., number of articles that contain a statistical reporting error), and (2) a vector containing the total number of observations, (3) the informed hypothesis, (4) a vector with prior parameter α_i for each binomial proportion, (5) a vector with prior parameter β_i for each binomial proportion, and (6) the categories of interest (i.e., journal names). Since we have no specific expectations about the distribution of statistical reporting errors across journals, we set all parameters α_i and β_i to one which corresponds to uniform Beta distributions. With this information, we can now conduct the analysis with the function binom_bf_informed.

```
# Prior specification
# We assign a uniform beta distribution to each binomial proportion
a \leftarrow rep(1, 8)
b < - rep(1, 8)
# Specifying the informed Hypothesis
Hr <- c('JAP , PS , JCCP , PLOS , DP , FP , JEPG < JPSP')</pre>
# Category labels
journal names <- journals$journal</pre>
# Execute the analysis
results_HO_Hr <- binom_bf_informed(x = x, n = n, Hr = Hr, a = a, b = b,
                                factor levels = journal names,
                                bf type = 'BF0r', seed = 2020)
BFre <- results HO Hr$bf list$bfr table['BFre']
BFe0 <- results_HO_Hr$bf_list$bf0_table['BFe0']
BFr0 <- results HO Hr$bf list$bf['BFr0']
```

The Bayes factor BF_{r0} suggests extreme evidence for the informed hypothesis that the social psychology journal JPSP has the highest prevalence for statistical reporting errors compared to the null hypothesis that the statistical reporting errors are equal across journals; $BF_{r0} = -158$. When taking a closer look at the Bayes factors, we also see that the data suggest that the null hypothesis that the statistical reporting errors are equal across journals is highly unlikely compared to the encompassing hypothesis; with BF_{e0} of 7.38×10^{67} . In addition, the results suggest that the data are 7.43 more likely under the informed

hypothesis than under the hypothesis that the ordering of the journals can vary freely.

In order to get a clearer picture about the ordering of the journals, we can investigate
the posterior estimates under the encompassing model as the next step. The posterior
median and 95% credible interval are returned the summary-method and can be plotted, too:

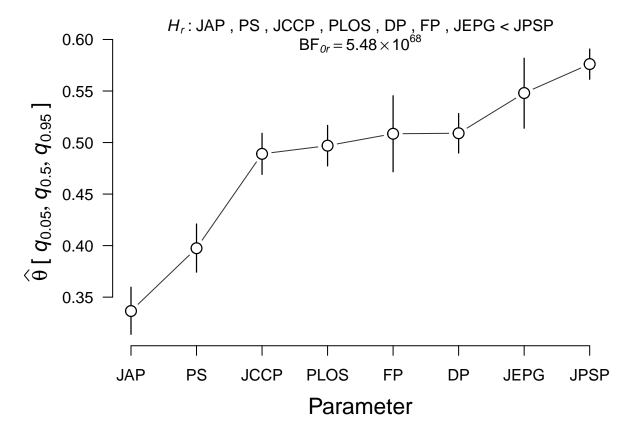


Figure 3. The figure displays for each journal the posterior estimates for the prevalence that an article includes a statistical reporting error and the corresponding 95% credible intervals based on the encompassing model. It appears that all journals show a relatively similar prevalence for statistical reporting errors, with the exception of the Journal of Applied Psychology (JAP) and Psychological Science (PS), whose prevalence is much lower.

Discussion. In this example, we tested whether the prevalence for statistical
reporting errors for articles published in social psychology journals (i.e., JPSP) is higher
than for articles published in other journals. We tested this hypothesis against the null
hypothesis that the prevalence for statistical reporting errors is equal across all journals. The

resulting Bayes factor of $BF_{r0} = 1.82 \times 10^{-69}$ provides extreme evidence for the informed hypothesis. However, this result should be interpreted with caution and be considered more differentiated. It seems that the result is above all an indication that the null hypothesis is highly misspecified and that the prevalence for a statistical reporting error varies greatly from journal to journal. Evidence that JPSP stands out and has a higher prevalence than the other journals is relatively small; the data provided only moderate evidence against the encompassing hypotheses.

Summary Summary

The R package multibridge facilitates the computation of Bayes factors for informed 424 hypotheses in multinomial models. The underlying algorithm is based on a recently 425 developed bridge sampling routine that is more efficient and reliable than available methods. 426 multibridge can evaluate hypotheses that feature equality constraints, inequality 427 constraints, and free parameters as well as mixtures between them. The core functions of the 428 software package were illustrated with two empirical examples. The **multibridge** package is 429 under continuous development. In the future, we aim to implement methods that extend the functionality of the package to hierarchical binomial and multinomial models. In addition, 431 we want to enable users to specify order constraints that are more complex, including 432 hypotheses on the size ratios of the parameters of interest or the difference between category 433 proportions. 434

reiterate benefits over existing packages

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Table 1 $S3\ methods\ available\ in\ {\it multibridge}$

Function Name(s)	S3 Method	Description	
mult_bf_informed,	print	Prints model specifications and descriptives.	
binom_bf_informed			
	summary	Prints and returns the Bayes factor and associated	
		hypotheses for the full model, and all equality and	
		inequality constraints.	
	plot	Plots the posterior median and 95% credible inter-	
		val of the parameter estimates of the encompassing	
		model.	
	bayes_factor	Contains all Bayes factors and log marginal likeli-	
		hood estimates for inequality constraints.	
	samples	Extracts prior and posterior samples from con-	
		strained distribution (if bridge sampling was ap-	
		plied).	
	bridge_output	Extracts bridge sampling output and associated	
		error measures.	
	restriction_list	Extracts restriction list and associated informed	
		hypothesis.	
binom_bf_inequality,	, print	Prints the bridge sampling estimate for the log	
binom_bf_inequality		marginal likelihood and the corresponding percent-	
		age error.	
	summary	Prints and returns the bridge sampling estimate	
		for the log marginal likelihood and associated error	
		terms.	

Table 2

To estimate the Bayes factor in favor for or against the specified informed hypothesis, the user provides the core functions mult_bf_informed and binom_bf_informed with the following basic required arguments

Argument	Description	
х	a vector with data (for multinomial models) or a vector of counts of	
	successes, or a two-dimensional table (or matrix) with 2 columns,	
	giving the counts of successes and failures, respectively (for binomial	
	models)	
n	numeric. Vector of counts of trials. Must be the same length as \mathbf{x} .	
	Ignored if \mathbf{x} is a matrix or a table	
Hr	string or character. Encodes the user specified informed hypothesis.	
	Users can either use the specified factor_levels or indexes to refer	
	to parameters.	
a	numeric. Vector with concentration parameters of Dirichlet distribu-	
	tion (for multinomial models) or α parameters for independent beta	
	distributions (for binomial models). Default sets all parameters to 1	
b	numeric. Vector with β parameters. Must be the same length as ${\tt x}.$	
	Default sets all β parameters to 1	
factor_levels	character. Vector with category names. Must be the same length	
	as x	

 $\label{thm:condition} \begin{tabular}{ll} Table 3 \\ Core functions available in {\it multibridge} \\ \end{tabular}$

Function Name(s)	Description	
mult_bf_informed	Evaluates informed hypotheses on multinomial parameters.	
mult_bf_inequality	Estimates the marginal likelihood of a constrained prior or	
	posterior Dirichlet distribution.	
mult_bf_equality	Computes Bayes factor for equality constrained multinomial	
	parameters using the standard Bayesian multinomial test.	
mult_tsampling	Samples from truncated prior or posterior Dirichlet density.	
lifestresses, peas	Datasets associated with informed hypotheses in multinomial	
	models.	
binom_bf_informed	Evaluates informed hypotheses on binomial parameters.	
binom_bf_inequality	Estimates the marginal likelihood of constrained prior or pos-	
	terior beta distributions.	
binom_bf_equality	Computes Bayes factor for equality constrained binomial pa-	
	rameters.	
binom_tsampling	Samples from truncated prior or posterior beta densities.	
journals	Dataset associated with informed hypotheses in binomial mod-	
	els.	

 ${\tt generate_restriction_list} \\ Encodes \ the \ informed \ hypothesis.$

Table 4

The Table shows the Observed Counts, Observed Proportions, and Expected Proportions of first digits in Greece governmental data. The total sample size was N=1,497 observations. Note that the observed proportions and counts deviate slightly from those reported in Rauch et al. (2011) (probably due to rounding errors).

Leading digit	Observed Counts	Observed Proportions	Expected Proportions:
			Benford's Law
1	509	0.340	0.301
2	353	0.236	0.176
3	177	0.118	0.125
4	114	0.076	0.097
5	77	0.051	0.079
6	77	0.051	0.067
7	53	0.035	0.058
8	73	0.049	0.051
9	64	0.043	0.046

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Appendix

Transforming An Ordered Probability Vector To The Real Line

Since we choose the multivariate normal as proposal distribution, the mapping between the proposal and target distribution requires us to move θ to the real line. Crucially this 521 transformation needs to retain the inequality constraints imposed on the parameters, that is, 522 it needs to take into account the lower bound l_k and the upper bound u_k of each θ_k . To 523 achieve this goal, **multibridge** uses a probit transformation as proposed in Sarafoglou et al. 524 (2020) which subsequently transforms the elements in θ moving from its lowest to its highest 525 value. In the binomial model, we move all elements in θ to the real line and thus construct a 526 new vector $\boldsymbol{y} \in \mathbb{R}^K$. For multinomial models, however, it follows from the sum-to-one 527 constraint that the vector $\boldsymbol{\theta}$ is completely determined by its first K-1 elements of 528 $\boldsymbol{\theta}: \theta_1 \leq \theta_2 \leq \cdots \leq 1 - \sum_{k=1}^K \theta_k$. Hence, for the transformation we will only consider the first 529 K-1 elements of $\boldsymbol{\theta}$ and we will transform them to K-1 elements of a new vector $y \in \mathbb{R}^{K-1}$. 531

Let ϕ denote the density of a normal variable with a mean of zero and a variance of one, Φ denote its cumulative density function, and Φ^{-1} denote the inverse cumulative density function. Then for each element θ_k , the transformation is

$$\xi_k = \Phi^{-1} \left(\frac{\theta_k - l_k}{u_k - l_k} \right),\,$$

The inverse transformation is given by

$$\theta_k = (u_k - l_k)\Phi(\xi_k) + l_k.$$

The Jacobian of this transformation is:

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$$|J| = \prod_{k=1}^{K-1} (u_k - l_k) \phi(\xi_k).$$

To perform the transformations, we thus need to determine the lower bound l_k and the

upper bound u_k of each θ_k . Assuming $\theta_{k-1} < \theta_k$ for $k \in \{1 \cdots, K\}$ the lower bound for any element in $\boldsymbol{\theta}$ is defined as

$$l_k = \begin{cases} 0 & \text{if } k = 1 \\ \theta_{k-1} & \text{if } 1 < k < K. \end{cases}$$

This definition holds for both binomial models and multinomial models. Differences in 540 these two models appear only when determining the upper bound for each parameter, since 541 parameters in a multinomial models are unit constrained. For binomial models, the upper 542 bound for each θ_k is simply 1. For multinomial models, however, the upper bound for each 543 θ_k depends on the size of smaller elements as well as on the number of remaining larger 544 elements in θ . To determine the upper bound for multinomial parameters we are using a 545 stick-breaking method (Frigyik, Kapila, & Gupta, 2010; Stan Development Team, 2020). 546 The stick-breaking approach represents θ as unit-length stick which we subsequently divide 547 into K elements. By this definition, the upper bound or any θ_k is:

$$u_k = \begin{cases} \frac{1}{K} & \text{if } k = 1\\ \frac{1 - \sum_{i < k} \theta_i}{ERS} & \text{if } 1 < k < K, \end{cases}$$
 (10)

where $1 - \sum_{i < k} \theta_i$ represents the length of the remaining stick, that is, the proportion of the unit-length stick that still needs to be divided among the remaining elements in $\boldsymbol{\theta}$. The elements in the remaining stick are denoted as ERS, and are computed as follows:

$$ERS = K - 1 + k$$

The transformations outlined above are suitable for binomial and multinomial models featuring hypotheses in which all parameters are inequality constrained. However, when hypotheses feature a combination of equality and inequality constrained parameters, as well as parameters that are free to vary we need to modify the formula to compute the upper and lower bounds:

$$l_k = \begin{cases} 0 & \text{if } k = 1\\ \frac{\theta_{k-1}}{e_{k-1}} \times e_k & \text{if } 1 < k < K, \end{cases}$$
 (11)

where e_{k-1} refers to the number of equality constrained parameters that are collapsed in θ_{k-1} .

The upper bound for parameters in the binomial models still remains 1. For multinomial models, the upper bound is then defined as:

$$u_{k} = \begin{cases} \frac{1}{K} - (f_{k} \times l_{k}) & \text{if } k = 1 \\ \left(\frac{1 - \sum_{i < k} \theta_{i}}{ERS} - (f_{k} \times l_{k})\right) \times e_{k} & \text{if } 1 < k < K \text{ and } u_{k} \ge \max(\theta_{i < k}), \\ \left(2 \times \left(\frac{1 - \sum_{i < k} \theta_{i}}{ERS} - (f_{k} \times l_{k})\right) - \max(\theta_{i < k})\right) \times e_{k} & \text{if } 1 < k < K \text{ and } u_{k} < \max(\theta_{i < k}), \end{cases}$$

$$(12)$$

where f_k represents the number of free parameters that share common upper and lower bounds. The elements in the remaining stick are then computed as follows

$$ERS = e_k + \sum_{j>k} e_j \times f_j.$$

The rationale behind these modifications will be described in more detail in the following sections. In **multibridge**, information that is relevant for the transformation of the

parameter vectors is stored in the generated restriction_list which is returned by the
main functions binom_bf_informed and mult_bf_informed but can also be generated
separately with the function generate_restriction_list. This restriction list features the
sublist inequality_constraints which encodes the number of equality constraints
collapsed in each parameter in nr_mult_equal. Similarly the number of free parameters
that share a common bounds are encoded under nr_mult_free.

Equality Constrained Parameters. When informed hypotheses feature a mix of equality and inequality constrained parameters, we collapse in the constrained prior and posterior distributions all equality constrained parameters into one category. When transforming the samples from these distributions, we need to account for the fact that inequality constraints on the collapsed parameters might not hold even though the constraint is valid under the original parameter values. For instance, for $\theta_1 = \theta_2 = \theta_3 \le \theta_4 \le \theta_5$, where the elements in θ take the values (0.15, 0.15, 0.15, 0.25, 0.3), the inequality constraint does not hold for the collapsed parameters (i.e., $\theta^* \not< \theta_4 \le \theta_5$ since $0.45 \nleq 0.25 \le 0.3$). For these cases, the upper and lower bounds for the parameters need to be adjusted as follows:

$$l_k = \begin{cases} 0 & \text{if } k = 1\\ \frac{\theta_{k-1}}{e_{k-1}} \times e_k & \text{if } 1 < k < K, \end{cases}$$
 (13)

where e_{k-1} and e_k refer to the number of equality constrained parameters that are collapsed in θ_{k-1} and θ_k , respectively. The upper bound is defined as

$$u_k = \begin{cases} \frac{1}{ERS} \times e_k & \text{if } k = 1\\ \frac{1 - \sum_{i < k} \theta_i}{ERS} \times e_k & \text{if } 1 < k < K, \end{cases}$$
 (14)

where $1 - \sum_{i < k} \theta_i$ represents the length of the remaining stick and the number of elements in the remaining stick are computed as follows: $ERS = \sum_{k}^{K} e_k$. The upper bound is then multiplied by the number of equality constrained parameters within the current constraint.

Concretely, for the constraint above, that is $\theta^* \leq \theta_4$, the lower bound for θ^* would be 0.

The upper bound is computed by taking into account the number of equality constrained parameters, such that $u_k = 1/5 \times 3 = 0.6$. For θ_4 the lower bound is $\theta^*/3 = 0.15$, since 3 parameters are collapsed in θ^* . The upper bound for θ_4 is then $\frac{(1-\theta^*)}{2} = 0.275$ and θ_5 is $1-\theta^*-\theta_4 = 1-0.45-0.25 = 0.3$.

Corrections for Free Parameters. Different adjustments are required for a sequence of inequality constrained parameters that have shared upper and lower bounds, but can vary freely within certain upper and lower bounds. For instance, the hypothesis

$$\mathcal{H}_r: \theta_1 < \theta_2, \theta_3$$

specifies that θ_2 and θ_3 have the shared lower bound θ_1 and the shared upper bound 1, however, θ_2 can be larger than θ_3 or vice versa. To integrate these cases within the stick-breaking approach one must account for these potential changes of order. For these cases, the lower bounds for the parameters remain unchanged, however the upper bounds need to be adjusted as follows:

$$u_k = \begin{cases} \frac{1}{K} - (f_k \times l_k) & \text{if } k = 1\\ \frac{1 - \sum_{i < k} \theta_i}{ERS} - (f_k \times l_k) & \text{if } 1 < k < K, \end{cases}$$

$$(15)$$

where f_k represents the number of free parameters that share common upper and lower bounds. Here, the number of elements in the remaining stick are computed as follows: $ERS = 1 + \sum_{j>k} f_j$. Subtracting the lower bound for the remaining free parameters from the upper bound of the current parameter secures a minimum stick-length for the remaining free parameters to comply with the constraint. A further correction is required, if a preceding free parameter (i.e., a free parameter that was already accounted for in the stick) is larger than the upper bound of the current parameter. In that case, we need we subtract the difference between the largest preceding free parameter in the sequence with the current

upper bound. Thus if $u_k < \max(\theta_{i < k})$, the upper bound becomes:

$$u_k = u_k - (\max(\theta_{i < k}) - u_k) \tag{16}$$

$$= 2 \times u_k - \max(\theta_{i < k}). \tag{17}$$

To outline when such a correction is necessary, consider the constraint $\theta_1 \leq \theta_2, \theta_3 \leq \theta_4$, where the elements in $\boldsymbol{\theta}$ take on the values (0.1, 0.35, 0.15, 0.40). When transforming the parameters, the lower bound for θ_1 is 0, the upper bound $^1/4$. The parameters θ_2 and θ_3 share the same lower bound, which is, $\theta_1 = 0.1$. The upper bound for θ_2 , is the length of the remaining stick divided by the elements of the remaining stick, that is, $^{0.9}/_2 = 0.45$. From the resulting upper bound, we subtract the lower bound for the remaining free parameters of the sequence, which yields an upper bound for θ_2 of 0.45 - 0.1 = 0.35. Since θ_2 is the first free parameter in the sequence that is evaluated an additional downward correction is not necessary. The upper bound for θ_3 is (1 - 0.1 - 0.35)/2 = 0.275. However, if θ_3 would actually take on the value 0.275, θ_4 would need to be 0.275 too, which would violate the constraint (i.e., $0.1 \leq 0.35, 0.275 \nleq 0.275$). Therefore, we apply the additional correction, such that

$$u_k = 2 \times u_k - \max(\theta_{i < k}) \tag{18}$$

$$= 2 \times 0.275 - 0.35 \tag{19}$$

$$=0.2,$$
 (20)

which secures the proper ordering for the remainder of the parameters, since $\theta_4 = 0.2$ would yield $0.1 \le 0.35, 0.2 \le 0.35$.

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