Estimating Causal Effects using Bayesian Methods with the R Package BayesCACE

by Jincheng Zhou, Jinhui Yang, James S. Hodges, Lifeng Lin, and Haitao Chu

Abstract Noncompliance, a common problem in randomized clinical trials (RCTs), complicates the analysis of the causal treatment effect, especially in meta-analysis of RCTs. The complier average causal effect (CACE) measures the effect of an intervention in the latent subgroup of the population that complies with its assigned treatment (the compliers). Recently, Bayesian hierarchical approaches have been proposed to estimate the CACE in a single RCT and a meta-analysis of RCTs. We develop an R package, BayesCACE, to provide user-friendly functions for implementing CACE analysis for binary outcomes based on the flexible Bayesian hierarchical framework. This package includes functions for analyzing data from a single study and for performing a meta-analysis with either complete or incomplete compliance data. The package also provides various functions for generating forest, trace, posterior density, and auto-correlation plots, which can be useful to review noncompliance rates, visually assess the model, and obtain study-specific and overall CACEs.

1 Introduction

Noncompliance in randomized clinical trials and causal effect

Randomized clinical trials (RCTs) are often used to evaluate healthcare-related interventions. An RCT typically compares an experimental treatment to a standard treatment or to a placebo. A common problem in RCTs is that not all patients fully comply with the allocated treatments. Although RCT investigators control the randomization process, the actual treatments received by study participants may not follow the randomization allocation; this is called noncompliance. For example, in trials of a therapist-led intervention, noncompliance occurs when individuals randomized to the intervention fail to take the intervention (e.g., due to severe adverse events), or when some patients assigned to the control figure out a way to take the intervention. In some cases, investigators can collect outcome data on all of these patients, regardless of whether they followed interventions. When compliance status is incompletely observed, it is more complicated to evaluate the causal treatment effect.

Conventionally, researchers use the intention-to-treat (ITT) analysis, in which data are analyzed based on treatments originally allocated rather than treatments actually received. The ITT method estimates the effect of being offered the intervention, namely, the overall effect in the real world in which the intervention is made available. However, our interest may lie in a different question, namely the causal effect of actually receiving the treatment. When using ITT, the treatment effect tends to be diluted by including people who do not receive the treatment to which they were randomly allocated (Freedman 1990).

To identify a treatment's causal effect, the principal stratification framework (Frangakis and Rubin 2002) is proposed, which stratifies subjects on the joint potential post-randomization variables. This causal inference method is widely used in handling various intercurrent events (also called an intermediate variable) in areas like vaccine effect (Hudgens and Halloran 2006; J. Zhou et al. 2016), pain relief use (Baccini, Mattei, and Mealli 2017), surrogate endpoint evaluation (Gilbert et al. 2015), noncompliance (J. Zhou et al. 2019), etc. An estimator called the "complier average causal effect" (CACE) has been proposed, in which patients are classified into different principal strata (compliers, never-takers, always-takers, and defiers) based on their potential behavior after assignment to both the treatment and control arms. Compliers are patients who receive the treatment as assigned; never-takers are those who do not receive treatment, regardless of treatment assignment; always-takers are those who receive treatment regardless of treatment assignment; and patients who always do the opposite of their treatment assignment are called *defiers*. The CACE is the causal effect of the intervention estimated from compliers. Because patients are assumed to be compliers (or not) before the randomization, the CACE retains the benefit of the randomization. Specifically, CACE is an unbiased estimate of the difference in outcomes for compliers in the intervention group compared to those in the control group, who would have engaged with treatment had they been randomized to the intervention group.

The biggest challenge in estimating the CACE is that we cannot actually identify which participants are compliers. Some of those receiving the treatment in the intervention group are compliers, but the rest are always-takers. Similarly, some of those not receiving the treatment in the control arm are compliers, but others are never-takers. Several R packages are available to perform CACE analysis in a single study. For example, the **noncomplyR** package (Coggeshall 2017) provides convenient functions for using Bayesian methods to perform inferences on the CACE. The package **eefAnalytics** (Kasim et al.

CACE in meta-analysis

All of the above methods are framed in a single study setting. However, for analyzing multiple trials in the presence of noncompliance, no software is available for causal effect analysis, specifically for meta-analysis. When noncompliance data are reported in each trial, one could intuitively implement a two-step approach by first estimating CACE for each study and then combining the study-specific estimates using a fixed-effect or random-effects model to estimate the population-averaged CACE. Recently, J. Zhou et al. (2019) proposed a Bayesian hierarchical model to estimate the CACE in a meta-analysis of randomized trials where compliance may be heterogeneous between studies. It is also common that noncompliance data are not available for some trials. Simply excluding trials with incomplete noncompliance data from a meta-analysis can be inefficient and potentially biased. J. Zhou, Hodges, and Chu (2021) proposed an improved flexible Bayesian hierarchical CACE framework to account simultaneously for heterogeneous noncompliance and incomplete noncompliance data. More recently, T. Zhou et al. (2021) used a generalized linear latent and mixed model to estimate CACE, which accounts for between-study heterogeneity with random effects. The package BayesCACE focuses on providing user-friendly functions to estimate CACE in either a single study or metaanalysis using models based on J. Zhou et al. (2019), Baker (2020), J. Zhou, Hodges, and Chu (2020) and J. Zhou, Hodges, and Chu (2021).

This article introduces the R package **BayesCACE**, which performs CACE analysis for binary outcomes in a single study, and meta-analysis with either complete or incomplete noncompliance information. The package **BayesCACE** is available from the Comprehensive R Archive Network (CRAN). It uses Markov chain Monte Carlo (MCMC) methods on the R platform through JAGS. JAGS is a program for analyzing Bayesian hierarchical models using MCMC simulation, which is available for diverse computer platforms including Windows and Mac OS X. Convergence of the MCMC routine can be assessed by the function outputs. The package also provides functions to generate posterior trace plots, density plots, and auto-correlation plots. For meta-analysis, the package provides a forest plot of study-specific CACE estimates with 95% credible intervals as well as the overall CACE estimate, to visually display the causal treatment effect comparisons.

This article is organized as follows. The next section defines CACE in mathematical notation that will be used throughout the paper. We also describe the assumptions needed to make the CACE a valid causal effect estimator. Following that, we present an overview of the Bayesian hierarchical models for CACE implemented in the **BayesCACE** package. Then, we illustrate use of the package with a case study example and discuss the output structures. Finally, we provide a brief discussion with potential future improvements.

Assumptions and definition of CACE

The CACE is a measure of the causal effect of a treatment or intervention on patients who received it as intended by the original group allocation. It is an unbiased causal effect estimate based on five standard assumptions commonly used in causal inference research. First, it assumes that potential outcomes for each participant are independent of the potential outcomes for other participants, known as the Stable Unit Treatment Value Assumption (SUTVA). Second, it assumes that assignment to treatment is random, so that the proportion of compliers should be the same in the intervention and control groups, thus allowing the estimation of one of the core unobserved parameters needed to derive a CACE estimate. Third, it assumes that treatment assignment has an effect on the outcome only if it changes the actual treatment taken, an assumption known as exclusion restriction. For never-takers, for instance, it assumes that simply being assigned to treatment does not affect their outcomes, as they do not actually receive the treatment assigned to them. Fourth, it assumes that assigning the study treatment to participants in the intervention group induces at least some participants to receive the treatment, so the compliance rate is not zero. Finally, it assumes that there is a monotonic relationship between treatment assignment and treatment receipt, which implies that there are no individuals for whom assignment to treatment actually reduces the likelihood of receiving treatment (i.e., no defiers). This assumption reduces the number of compliance types for which estimates are derived, permitting a properly identified model.

We follow J. Zhou et al. (2019) and introduce notation both on the individual level and on the study level. Suppose a meta-analysis reviews I two-armed RCTs, and N_i is the number of subjects in

On the individual level, notation is defined as follows for subject *j* in trial *i*.

- 1. Let $R_{ij} = r$ index the randomization assignment with r = 0 for those randomized to control and r = 1 for those randomized to the intervention.
- 2. Let $T_{ij}^r = t \in \{0, 1\}$ be the indicator of whether the individual received the intervention. This is a *potential* outcome under the randomization assignment $r \in \{0, 1\}$, i.e., what the value of treatment *t* would be for individual (i, j) if r = 0 or r = 1, respectively.
- 3. Let $Y_{ij}^{r,t} = o \in \{0,1\}$ be the *potential* binary outcome under randomization assignment *r* and treatment received *t*. Note that the *exclusion restriction* assumption allows us to define $Y_{ij}^t \equiv Y_{ij}^{r,t}$.
- 4. The sets of $\{Y_{ij}^{r,t}\}$ and $\{T_{ij}^r\}$ are the *potential* outcome and treatment-received status respectively under possible *r* and *t*, but for each subject in a trial, only one of the possible values of each set can be observed. Therefore, we denote the observed response and received treatment variables as Y_{ij} and T_{ij} .
- 5. We allow $T_{ij}^{j} = *$ if the actual received treatment is not recorded. Then let $M_{ij} = m$ be the missing indicator corresponding to whether subject *j* has actual treatment received status on record (*m* = 0) or missing (*m* = 1).
- 6. Using these potential outcomes, we can define the compliers and the CACE. Let *C_{ij}* be the latent compliance class of individual *j* in trial *i*, defined as follows:

$$C_{ij} = \begin{cases} 0, & \text{for never-taker with } (T_{ij}^0, T_{ij}^1) = (0, 0) \\ 1, & \text{for complier with } (T_{ij}^0, T_{ij}^1) = (0, 1) \\ 2, & \text{for always-taker with } (T_{ij}^0, T_{ij}^1) = (1, 1) \\ 3, & \text{for defier with } (T_{ij}^0, T_{ij}^1) = (1, 0) \end{cases}$$

A subject's compliance status C_{ij} is not observable because in a two-arm trial, only one of T_{ij}^1 and T_{ij}^0 can be observed. Based on the observed randomization group and actual treatment received, the compliance classes can be only partially identified.

Now, the complier average causal effect of the *i*-th trial is the average difference between potential outcomes for compliers. In this case, the CACE in study *i* is $\theta_i^{\text{CACE}} = E(Y_{ij}^1 - Y_{ij}^0 | C_{ij} = 1)$, where the patients for whom $C_{ij} = 1$ are the compliers.

On the study level, n_{irto} denotes the observed number of individuals in study *i*, randomization group *r*, actual received treatment group *t*, and outcome *o*. If the compliance status of individual *j* in trial *i* is not on record, $T_{ij} = t = *$ so the corresponding count is n_{ir*o} , which is the sum of the two unobserved counts n_{ir0o} and n_{ir1o} .

2 Estimating CACE

This section briefly describes the Bayesian hierarchical models used to estimate CACE. These models form the basis of the framework proposed by J. Zhou et al. (2019) and underlie the **BayesCACE** package. In addition to the notation defined in the previous section, we define the following parameters for study *i*.

- 1. Let π_{ia} and π_{in} be the probabilities of being an always-taker and a never-taker, respectively. Because defiers are ruled out by the monotonicity assumption, each trial has at most only three compliance classes. Thus the probability of being a complier in study *i* is $\pi_{ic} = 1 - \pi_{ia} - \pi_{in}$.
- 2. Define these response probabilities: u_{i1} for a complier randomized to the treatment group; v_{i1} for a complier randomized to the control/placebo group; s_{i1} for a never-taker; and b_{i1} for an always-taker. Thus for study *i*, the parameters included in the model are $\beta_i = (\pi_{ia}, \pi_{in}, u_{i1}, v_{i1}, s_{i1}, b_{i1})$.

As the outcome is binary, the expected difference between outcomes from the two treatment groups among compliers is just the risk difference between u_{i1} and v_{i1} . Therefore, the CACE can be written as $\theta_i^{\text{CACE}} = E(Y_{ij}^1 - Y_{ij}^0 | C_{ij} = 1) = u_{i1} - v_{i1}$.

CACE for a single trial with noncompliance

Consider first a single trial with noncompliance, i.e., I = 1, so all notation and parameters defined earlier are reduced to the version without subscript *i*. According to J. Zhou et al. (2019), each observed n_{rto} has a corresponding probability that can be written in terms of parameters defined in $\beta = (\pi_a, \pi_n, u_1, v_1, s_1, b_1)$, thus the vector $(n_{000}, n_{001}, n_{010}, n_{011}, n_{100}, n_{101}, n_{111})$ follows a multinomial distribution. The likelihood is available in the Supplemental Materials.

The CACE for a single study is $u_1 - v_1$, so the posterior of θ^{CACE} is the posterior of $u_1 - v_1$.

CACE for a meta-analysis with complete compliance information

This section introduces two methods for performing a meta-analysis of the CACE when noncompliance data are reported in each trial.

The two-step approach

As described in the previous section, using the observed data n_{irto} , θ_i^{CACE} is identified for study~*i*. Therefore, to estimate the population-average CACE in a meta-analysis, we propose combining the study-specific estimates and standard errors using a standard meta-analysis method such as the fixed-effect (Laird and Mosteller 1990) or random-effects model (Hedges and Vevea 1998; Hedges and Olkin 1985). We call this a "two-step" approach. As the CACE measure is a risk difference, a transformation may be necessary to ensure that the normal distribution assumption is approximately true. Building upon the well-developed R package metafor, various estimators suggested in the literature can be estimated to account for potential between-study heterogeneity in the CACE, e.g., the Hunter–Schmidt estimator, the Hedges estimator, or the empirical Bayes estimator (Viechtbauer 2010).

The Bayesian hierarchical model

In a meta-analysis, the CACE can also be estimated using the joint likelihood from the Bayesian hierarchical model. This method is systematically introduced in J. Zhou et al. (2019). The log likelihood contribution of trial *i* is denoted by adding a subscript *i* to each parameter. Then the log likelihood for all trials in the meta-analysis is $\log \mathcal{L}(\beta) = \sum_i \log L_i(\beta_i)$. Because the studies are probably not exactly identical in their eligibility criteria, measurement techniques, study quality, etc., differences in methods and sample characteristics may introduce heterogeneity to the meta-analysis. One way to model the heterogeneity is to use a random-effects model.

To guarantee the desired properties of study *i*'s latent compliance classes and to account for possible between-study heterogeneity in the compliance class and response probabilities, we use these transformations:

- 1. $\pi_{in} = \frac{\exp(n_i)}{1 + \exp(n_i) + \exp(a_i)}, \pi_{ia} = \frac{\exp(a_i)}{1 + \exp(n_i) + \exp(a_i)}, \text{ where } n_i = \alpha_n + \delta_{in}, a_i = \alpha_a + \delta_{ia}, \text{ and } (\delta_{in}, \delta_{ia})^\top \sim N(0, \Sigma_{ps}), \Sigma_{ps} = \begin{pmatrix} \sigma_n^2 & \rho\sigma_n\sigma_a \\ \rho\sigma_n\sigma_a & \sigma_a^2 \end{pmatrix}.$ 2. We also define random effect means on the transformed scale of each response properties.
- 2. We also define random effect models on the transformed scale of each response probability $s_{i1}, b_{i1}, u_{i1}, v_{i1}$: $g(s_{i1}) = \alpha_s + \delta_{is}, g(b_{i1}) = \alpha_b + \delta_{ib}, g(u_{i1}) = \alpha_u + \delta_{iu}, g(v_{i1}) = \alpha_v + \delta_{iv}$, where $g(\cdot)$ is a link function such as the logit or probit, $\delta_{is} \sim N(0, \sigma_s^2), \delta_{ib} \sim N(0, \sigma_b^2), \delta_{iu} \sim N(0, \sigma_u^2), \delta_{iv} \sim N(0, \sigma_v^2)$.

Here we allow correlation between n_i and a_i , and assign random effect variables to all parameters. However, if a parameter does not vary between trials, it can be modeled as a fixed effect. Let $f(\beta_i | \beta_0, \Sigma_0)$ be the distributions described above of all parameters $\beta_i = (\pi_{ia}, \pi_{in}, s_{i1}, b_{i1}, u_{i1}, v_{i1})$, where β_0 is the vector of mean hyper-parameters $(\alpha_n, \alpha_a, \alpha_s, \alpha_b, \alpha_u, \alpha_v)$, and Σ_0 is the diagonal covariance matrix containing $\Sigma_{ps}, \sigma_s^2, \sigma_u^2$ and σ_v^2 .

If we specify $f(\beta_0)$ and $f(\Sigma_0)$ as the prior distributions for the hyper-parameters, then the joint posterior distribution is proportional to the likelihood multiplied by the priors, i.e., $\prod_i L_i(\beta_i) f(\beta_i | \beta_0, \Sigma_0) f(\beta_0) f(\Sigma_0)$.

As stated earlier, $\theta_i^{CACE} = u_{i1} - v_{i1}$ for study *i*, so for the meta-analysis, the overall CACE is $\theta^{CACE} = E(\theta_i^{CACE}) = E(u_{i1}) - E(v_{i1})$. When a random effect δ_{iu} or δ_{iv} is not assigned in the model, $E(u_{i1}) = g^{-1}(\alpha_u)$ and $E(v_{i1}) = g^{-1}(\alpha_v)$. Otherwise, $E(u_{i1})$ and $E(v_{i1})$ can be estimated by integrating out the random effects, e.g., $E(u_{i1}) = \int_{-\infty}^{+\infty} g^{-1}(\alpha_u + t)\sigma_u^{-1}\phi(\frac{t}{\sigma_u})dt$, where $\phi(\cdot)$ is the standard Gaussian density. If the function $g(\cdot)$ is the probit link, this expectation has a closed form:

 $E(u_{i1}) = \Phi(\frac{\alpha_u}{\sqrt{1+\sigma_u^2}})$. If the link function $g(\cdot)$ is logit, a well-established approximation $E(u_{i1}) \approx \log t^{-1}(\frac{\alpha_u}{\sqrt{1+C^2\sigma_u^2}})$ can be used, where $C = \frac{16\sqrt{3}}{15\pi}$ (Zeger, Liang, and Albert 1988). The above formulas also apply to $E(v_{i1})$, the expected response rate of a complier in the control group.

The two-step approach, stated by Lin and Zeng (2010), can be viewed as asymptotically equivalent to the model using the joint likelihood. However, as the two-step approach requires the whole set of parameters to be estimated independently for each study, the total number of effective parameters tends to be larger than the Bayesian hierarchical model, so estimates using our method are likely to be more efficient.

CACE for meta-analysis with incomplete compliance information

Another advantage of the Bayesian hierarchical model is that it can include trials with incomplete compliance data. Commonly, some trials do not report noncompliance data because study investigators do not collect actual received treatment status for some subjects or simply do not report compliance. The two-step approach needs counts for all of the groups defined by randomized assignment, treatment received, and outcome in order to estimate the study-specific θ_i^{CACE} . Thus, by using this method, trials with incomplete compliance data are simply excluded, making estimation less efficient and potentially biased.

J. Zhou, Hodges, and Chu (2021) proposed a comprehensive framework to incorporate both heterogeneous and incomplete noncompliance data for estimating the CACE in a meta-analysis of RCTs. Here we present the data structure needed for binary outcomes. For study *i*, randomization group $r \in \{0, 1\}$ and output $o \in \{0, 1\}$, if the compliance information is reported, then values of n_{ir0o} and n_{ir1o} are reported, so we assign the marginal count $n_{ir*o} = 0$. Otherwise, we do not have data on outcomes for groups defined by actually received treatment, so only the marginal n_{ir*o} is observed, where n_{ir*o} is the number of patients randomized to treatment arm *r* who had outcome *o*. In this situation, the two unobserved counts n_{ir0o} and n_{ir1o} are assigned as 0. In the Supplemental Materials, a table for the observed counts data with corresponding probabilities is presented. The log likelihood is also obtained from the multinomial distribution. The CACE for this meta-analysis incorporating incomplete compliance data is $\theta^{CACE} = E(\theta_i^{CACE}) = E(u_{i1}) - E(v_{i1}) = \Phi(\frac{\alpha_u}{\sqrt{1+\sigma_u^2}}) - \Phi(\frac{\alpha_v}{\sqrt{1+\sigma_v^2}})$ if the probabilities is the structure of the structure and r_i are assigned as 0. In the supplemental Materials, a table for the observed counts data with corresponding probabilities is presented. The log likelihood is also obtained from the multinomial distribution. The CACE for this meta-analysis incorporating incomplete compliance data is $\theta^{CACE} = E(\theta_i^{CACE}) = E(u_{i1}) - E(v_{i1}) = \Phi(\frac{\alpha_u}{\sqrt{1+\sigma_u^2}}) - \Phi(\frac{\alpha_v}{\sqrt{1+\sigma_v^2}})$ if the probabilities is the function is used for r_i and r_i

probit link function is used for u_{i1} and v_{i1} .

3 Using the R package BayesCACE

The primary objective of the **BayesCACE** package is to provide a user-friendly implementation of the Bayesian method for estimating the CACE. The package is now available to download and install via CRAN at https://CRAN.R-project.org/package=BayesCACE. It can be installed within R using the command install.packages("BayesCACE"). The latest version of the package is 1.2.3.

The **BayesCACE** package depends on the R packages **rjags** (Plummer 2018), **coda** (Plummer et al. 2006), and **forestplot** (Gordon and Lumley 2017). Users need to install JAGS separately from its homepage http://mcmc-jags.sourceforge.net as the **BayesCACE** package does not include a copy of the JAGS library. The current version of JAGS is 4.3.0, which is the version of the package that **BayesCACE** requires; earlier versions of JAGS may not guarantee exactly reproducible results.

Data structure for estimating the CACE

We introduce the data structures through the illustrative example included in the package **BayesCACE**: epidural_c and epidural_ic. These two data sets were obtained from Bannister-Tyrrell et al. (2015), who conducted an exploratory meta-analysis of the association between using epidural analgesia in labor and the risk of cesarean section. The dataset epidural_c contains 10 trials with full compliance information; each trial has 8 observed counts, denoted by n_{irto} and presented in columns nirto for i = 1, ..., 10 and $r, t, o \in \{0, 1\}$. These data were re-analyzed by J. Zhou et al. (2019) in a meta-analysis using their proposed Bayesian hierarchical model to estimate the CACE. The function cace.meta.c() performs this analysis. The column study.id contains IDs for the 10 studies, and study.name labels each study by its first author's surname and its publication year.

The data can be loaded and printed using these commands:

```
library("BayesCACE")
data("epidural_c", package = "BayesCACE")
epidural_c
```

#>		study.id	study	.name	n000	n001	n010	n011	n100	n101	n110	n111
#>	1	1	Bofill,	1997	37	2	11	1	2	0	42	5
#>	2	2	Clark,	1998	72	6	68	16	7	2	134	13
#>	3	3	Halpern,	2004	62	5	44	7	0	0	112	12
#>	4	4	Head,	2002	51	7	2	0	3	0	43	10
#>	5	5	Jain,	2003	72	11	0	0	0	2	36	7
#>	6	6	Nafisi,	2006	179	19	0	0	0	0	173	24
#>	7	7	Nikkola,	1997	6	0	4	0	0	0	10	0
#>	8	8	Ramin,	1995	546	17	95	8	230	2	393	39
#>	9	9	Sharma,	1997	336	16	5	0	114	1	231	12
#>	10	10	Volmanen,	2008	23	1	3	0	1	0	23	1

The other dataset epidural_ic represents the situation in which not all trials report complete compliance data. It contains 27 studies, only 10 of which have full compliance information and are included in epidural_c. This dataset is also drawn from Bannister-Tyrrell et al. (2015), and represents studies with incomplete compliance information when estimating the CACE. The function cace.meta.ic() performs this analysis.

Each study is represented by one row in the dataset; the columns study.id and study.name have the same meanings as in the dataset epidural_c. Each study's data are summarized in 12 numbers (columns) denoted by n_{irto} and n_{ir*o} . For a particular randomization group $r \in \{0, 1\}$, the observed counts are presented either as n_{irto} or n_{ir*o} depending on whether the compliance information is available; values for other columns are denoted by 0. The corresponding column names in the dataset are nirto and nirso, respectively.

The first 6 rows of the dataset epidural_ic are printed below.

```
data("epidural_ic", package = "BayesCACE")
head(epidural_ic)
```

#>		study	y.id	study	.name	n000	n001	n010	n011	n0s0	n0s1	n100	n101	n110	n111
#>	1		1	Bofill,	1997	37	2	11	1	0	0	2	0	42	5
#>	2		2	Clark,	1998	72	6	68	16	0	0	7	2	134	13
#>	3		3	Dickinson,	2002	0	0	0	0	428	71	0	0	0	0
#>	4		4	Evron,	2008	40	4	0	0	0	0	0	0	0	0
#>	5		5	El Kerdawy,	2010	0	0	0	0	12	3	0	0	0	0
#>	6		6	Gambling,	1998	0	0	0	0	573	34	206	10	371	29
#>		n1s0	n1s1												
#>	1	0	6)											
#>	2	0	6)											
#>	3	408	85	5											
#>	4	129	19)											
#>	5	11	4	ł											
#>	6	0	e)											

Note that NA is not allowed in a dataset for the package **BayesCACE**, but some trials may have 0 events or 0 noncompliance rates.

Plotting noncompliance rates

Before performing the CACE analysis, one might want a visual overview of study-specific noncompliance rates in both randomization arms. The function plt.noncomp provides a forest plot of noncompliance rates in an R plot window. The function can be simply called as

plt.noncomp(data, overall = TRUE)

where data is a dataset with structure like epidural_c or epidural_ic. Specifically, the dataset should contain the following columns: study.id, study.name, and 8 or 12 columns of data represented by n_{irto} , or n_{irto} and n_{ir*o} (see previous section for more details). Each row corresponds to one study. Only studies with full compliance information are included in this plot because noncompliance rates cannot be calculated without compliance data. Figure 1 shows the resulting plot, where the red dot with its horizontal line shows the study-specific noncompliance rate with its 95% exact confidence interval for the patients randomized to the treatment arm, and the blue square with its horizontal line represents that rate and interval for those in the control arm. The confidence intervals are calculated



Figure 1: Noncompliance rates plot generated by the function plt.noncomp(). The red dots and lines show the study-specific noncompliance rate with its 95% confidence interval randomized to the treatment arm, and the blue squares and lines refer to those in the control arm.

by the Clopper–Pearson exact method (Clopper and Pearson 1934), which is based on the cumulative distribution function of the binomial distribution. Using the default overall = TRUE, the figure also gives a summary estimate of the compliance rates per randomization group. This overall rate is estimated using a logit generalized linear mixed model. Otherwise, if the argument overall is FALSE, the plot shows only study-specific noncompliance rates. Any additional parameters passed to the function will be automatically used in the forestplot function from the forestplot package.

CACE analysis for a single study or in a meta-analysis

The major functions in **BayesCACE** are cace.study(), cace.meta.c(), and cace.meta.ic(), which implement the models introduced earlier to perform Bayesian CACE analysis for different data structures. In particular, cace.study() performs CACE analysis for a single study. The function cace.meta.c() performs CACE analysis for a meta-analysis when each trial reports noncompliance information. Users can choose to do the analysis either by the two-step approach or using the Bayesian hierarchical model. When some trials do not report noncompliance data, the function cace.meta.ic() can be applied to perform a CACE meta-analysis using the likelihood provided in the Supplemental Materials. Each function may take 1–15 minutes to run. Generally the two-step approach using the function cace.meta.c() takes longer because MCMC chains are run on the studies one by one. The actual run time depends on the amount of data and the user's processor.

Function cace.study() for a study-specific analysis or a two-step meta-analysis

For the default interface, the arguments of the function cace.study() are

```
cace.study(data, param = c("CACE", "u1", "v1", "s1", "b1", "pi.c", "pi.n",
  "pi.a"), re.values = list(), model.code = '', digits = 3, n.adapt = 1000,
  n.iter = 100000, n.burnin = floor(n.iter/2), n.chains = 3, n.thin =
  max(1,floor((n.iter-n.burnin)/1e+05)), conv.diag = FALSE, mcmc.samples =
  FALSE, two.step = FALSE, method = "REML")
```

where users need to input data with the same structure as epidural_c, containing either one row of observations for a single study, or multiple rows referring to multiple studies in a meta-analysis. This function fits a model for a single study. If the data includes more than one study, the study-specific CACEs will be estimated by retrieving data row by row.

The argument param is a character string vector indicating the parameters to be tracked and estimated. By default all parameters are included: θ^{CACE} (CACE), u_1 (u1), v_1 (v1), s_1 (s1), b_1 (b1), π_a (pi.a), π_n (pi.n), and $\pi_c = 1 - \pi_a - \pi_n$ (pi.c). Users can modify the string vector to only include parameters of interest besides θ^{CACE} . Users can specify the prior distributions (mean and standard deviation) of $n, a, \alpha_s, \alpha_b, \alpha_u, \alpha_v$ with the revalues parameter. By default, the revalues list is empty,

and they are assigned to the transformed scale of the following parameters: $\pi_n = \frac{\exp(n)}{1+\exp(n)+\exp(a)}$,

 $\pi_a = \frac{\exp(a)}{1+\exp(n)+\exp(a)}$, $\operatorname{logit}(s_1) = \alpha_s$, $\operatorname{logit}(b_1) = \alpha_b$, $\operatorname{probit}(u_1) = \alpha_u$, and $\operatorname{probit}(v_1) = \alpha_v$, where $n, a \sim N(0, 2.5^2)$ and $\alpha_s, \alpha_b, \alpha_u, \alpha_v \sim N(0, 2^2)$. With these settings, a 95% prior probability interval for any of the probabilities π_{in}, π_{ia} , and π_{ic} ranges from about 0.001 to 0.91, and a 95% prior interval for the probabilities s_1, b_1, u_1 , and v_1 ranges approximately from 0.01 to 0.98. The prior parameters are passed into the model.study function to get the model code, which first calls the prior.study to get the custom prior distribution. Here we give an example output of prior.study when assigning $N(0, 10^{-2})$ to every parameter:

out.string < "# priors
 n ~ dnorm(0, 0.01)
 a ~ dnorm(0, 0.01)
 alpha.s ~ dnorm(0, 0.01)
 alpha.b ~ dnorm(0, 0.01)
 alpha.u ~ dnorm(0, 0.01)
 alpha.v ~ dnorm(0, 0.01)
 "</pre>

To customize the model fully, the user can pass their complete model string to the cace.study() function with the parameter model.code. The arguments n.adapt, n.iter, n.burnin, n.chains, and n. thin control the MCMC algorithm run by the R package rjags (Plummer 2018). The argument n.adapt is the number of iterations for adaptation; it is used to maximize the sampling efficiency, and the default is set as 1,000. The argument n.chains determines the number of MCMC chains (the default is 3); n.iter is the number of iterations of each MCMC chain; n.burnin is the number of burn-in iterations to be discarded at the beginning of each chain; n. thin is the thinning rate for MCMC chains, which is used to avoid potential high auto-correlation and to save computer memory when n.iter is large. The default of n.thin is set as 1 or the largest integer not greater than ((n.iter – n.burnin)/1e+05)), whichever is larger. The argument conv.diag specifies whether to compute the Gelman and Rubin convergence statistic (\hat{R}) of each parameter as a convergence diagnostic (Brooks and Gelman 1998; Gelman and Rubin 1992). The chains are considered well-mixed and converged to the target distribution if $\hat{R} \leq 1.1$. If the argument mcmc.samples = TRUE, the function saves each chain's MCMC samples for all parameters, which can be used to produce trace, posterior density, and auto-correlation plots by calling the functions plt.trace, plt.density, and plt.acf.

By default, the function cace.study() returns a list including posterior estimates (posterior mean, standard deviation, median, and a 95% credible interval with 2.5% and 97.5% quantiles as the lower and upper bounds), and the deviance information criterion (DIC) statistic (Spiegelhalter et al. 2002) for each study. The argument two.step is a logical value indicating whether to conduct a two-step meta-analysis. If two.step = TRUE, the posterior mean and standard deviation of study-specific θ_i^{CACE} are used to perform a standard meta-analysis, using the R package metafor. The default estimation method is the REML (restricted maximum-likelihood estimator) method for the random-effects model (Harville 1977). Users can change the argument method to obtain different meta-analysis estimators from either a random-effects model or a fixed-effect model, e.g., method = "DL" refers to the DerSimonian–Laird estimator. More details are available from the documentation of the function metafor : :rma (Viechtbauer 2010). If the input data include only one study, the meta-analysis result is the same as the result from the single study.

Here is an example to demonstrate the function's usage. We call the function cace.study() on the dataset epidural_c as follows:

The following messages are output as the code runs:

% NA is not allowed in the input data set; % the rows containing NA are removed. Compiling model graph Resolving undeclared variables Allocating nodes Graph information: Observed stochastic nodes: 2 Unobserved stochastic nodes: 6 Total graph size: 44

Initializing model

If the dataset contains more than one study, e.g., the epidural_c dataset has 10 trials, then once the JAGS model compiles for the first study, it automatically continues to run on the next study's data. The results are saved in the object out.study, a list containing the model name, posterior information for each monitored parameter, and DIC of each study. We can use parameter names to display the corresponding estimates. The argument digits in the function cace.study() can be used to change the number of significant digits to the right of the decimal point. Here, we used the default setting digits = 3. For example, the estimates of θ^{CACE} for each single study (posterior mean and standard deviation, posterior median, 95% credible interval, and time-series standard error) can be displayed as:

out.study\$CACE

#>		Mean	SD	2.5%	50%	97.5%	Naive SE	Time-series SE
#>	[1,]	0.04980	0.0797	-0.09510	4.46e-02	0.2180	1.45e-04	2.51e-04
#>	[2,]	-0.02490	0.0489	-0.12200	-2.23e-02	0.0785	8.94e-05	1.48e-04
#>	[3,]	-0.02210	0.0606	-0.12700	-2.90e-02	0.1120	1.11e-04	1.94e-04
#>	[4,]	0.07180	0.0758	-0.07550	7.10e-02	0.2230	1.38e-04	2.01e-04
#>	[5,]	0.08250	0.0768	-0.06260	8.11e-02	0.2370	1.40e-04	2.51e-04
#>	[6,]	0.02600	0.0319	-0.03650	2.59e-02	0.0891	5.83e-05	7.55e-05
#>	[7,]	0.01430	0.1580	-0.28200	2.11e-04	0.4050	2.89e-04	4.21e-04
#>	[8,]	0.05030	0.0248	0.00176	5.02e-02	0.0993	4.54e-05	7.34e-05
#>	[9,]	-0.01100	0.0234	-0.05740	-1.09e-02	0.0350	4.27e-05	6.22e-05
#>	[10,]	0.00145	0.0655	-0.13400	-4.36e-06	0.1460	1.20e-04	1.56e-04

If the argument conv.diag is specified as TRUE, the output list contains a sub-list conv.out, which outputs the point estimates of the "potential scale reduction factor" (the Gelman and Rubin convergence statistic, labeled Point est.) calculated for each parameter from each single study, and their upper confidence limits (labeled Upper C.I.). Approximate convergence is diagnosed when the upper limit is close to 1 (Brooks and Gelman 1998; Gelman and Rubin 1992). For example, the first sub-list from conv.out is:

out.study\$conv.out[[1]]

#>		Point est.	Upper C.I.
#>	CACE	1.000025	1.000046
#>	b1	1.000041	1.000129
#>	pi.a	1.000025	1.000094
#>	pi.c	1.000036	1.000134
#>	pi.n	1.000029	1.000067
#>	s1	1.000014	1.000018
#>	u1	1.000016	1.000033
#>	v1	1.000077	1.000185

In this example, we included mcmc.samples = TRUE in the argument, so the output list out.study includes each chain's MCMC samples for all parameters. They can be used with our plotting functions to generate the trace, posterior density, and auto-correlation plots for further model diagnostics.

If the dataset used by the function cace.study() has more than one study, specifying the argument two.step = TRUE causes the two-step meta-analysis for θ^{CACE} to be done. The outcomes are saved as a sub-list object meta. Note that users can obtain different meta-analysis estimators by changing the method argument as described earlier.

out.study\$meta

```
#>
#> Random-Effects Model (k = 10; tau^2 estimator: REML)
#>
#> tau^2 (estimated amount of total heterogeneity): 0.0002 (SE = 0.0008)
#> tau (square root of estimated tau^2 value):
                                                    0.0131
#> I^2 (total heterogeneity / total variability):
                                                    8.10%
#> H^2 (total variability / sampling variability): 1.09
#>
#> Test for Heterogeneity:
#> Q(df = 9) = 5.9353, p-val = 0.7464
#>
#> Model Results:
#>
#> estimate
            se
                      zval
                               pval
                                       ci.lb
                                             ci.ub
    0.0182 0.0143 1.2758 0.2020 -0.0098 0.0462
#>
#>
#> --
#> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Function cace.meta.c() for meta-analysis with complete compliance data

The function cace.meta.c() performs the Bayesian hierarchical model method for meta-analysis when the dataset has complete compliance information for all studies. The function's default arguments are as shown:

```
cace.meta.c(data, param = c("CACE", "ulout", "vlout", "slout", "blout",
  "pic", "pin", "pia"), random.effects = list(), re.values = list(),
  model.code = '', digits = 3, n.adapt = 1000, n.iter = 100000,
  n.burnin = floor(n.iter/2), n.chains = 3, n.thin =
  max(1,floor((n.iter-n.burnin)/100000)), conv.diag = FALSE,
  mcmc.samples = FALSE, study.specific = FALSE)
```

The arguments controlling the MCMC algorithm are mostly similar to those of cace.study(). One major difference is that users need to specify parameters that are modeled as random effects. Earlier, we showed how to specify random effects for each parameter on the transformed scales, namely δ_{in} , δ_{ia} , δ_{iu} , δ_{iv} , δ_{is} , and δ_{ib} , and allowed a non-zero correlation ρ between δ_{in} and δ_{ia} . The model with all of these random effects as well as the correlation ρ is considered the full model. However, this function is flexible, allowing users to choose which random effects to include by specifying the random.effects argument. By default, the list is empty and all of the list values are set to TRUE. Users can customize that by setting delta.n, delta.a, delta.u, delta.v, delta.s, delta.b, and/or cor to FALSE. Note that ρ (cor) can only be included when both δ_{in} (delta.n) and δ_{ia} (delta.a) are set to TRUE. Otherwise, a warning is shown and the model continues running by forcing delta.n = TRUE and delta.a = TRUE. The default parameters to be monitored depend on which parameters are modeled as random effects. For example, ulout refers to $E(u_{i1})$, where for the probit link, $E(u_{i1}) = \Phi(\alpha_u)$ if δ_u is not specified in the model, and $E(u_{i1}) = \Phi(\frac{\alpha_u}{\sqrt{1+\sigma_u^2}})$ when the random effect δ_u is included.

Users can use the re.values parameter to customize the prior distribution. Like the function cace.study(), by default, weakly informative priors α_n , $\alpha_a \sim N(0, 2.5^2)$ and α_s , α_b , α_u , $\alpha_v \sim N(0, 2^2)$ are assigned to the means of these transformed parameters: $\pi_{in} = \frac{\exp(n_i)}{1+\exp(n_i)+\exp(a_i)}$, $\pi_{ia} = \frac{\exp(a_i)}{1+\exp(n_i)+\exp(a_i)}$, where $n_i = \alpha_n + \delta_{in}$, $a_i = \alpha_a + \delta_{ia}$, $\log it(s_{i1}) = \alpha_s + \delta_{is}$, $\log it(b_{i1}) = \alpha_b + \delta_{ib}$, $\operatorname{probit}(u_{i1}) = \alpha_u + \delta_{iu}$, and $\operatorname{probit}(v_{i1}) = \alpha_v + \delta_{iv}$. For the random effects, we have $\delta_{is} \sim N(0, \sigma_s^2)$, $\delta_{ib} \sim N(0, \sigma_b^2)$, $\delta_{iu} \sim N(0, \sigma_u^2)$, and $\delta_{iv} \sim N(0, \sigma_v^2)$, as response rates are assumed to be independent between latent classes. A *Gamma*(2,2) hyper-prior distribution is assigned to the precision parameters σ_s^{-2} , σ_b^{-2} , σ_u^{-2} and σ_v^{-2} , which corresponds to a 95% interval of (0.6, 2.9) for the corresponding standard deviations, allowing moderate heterogeneity in the response rates. In a reduced model with one of δ_{in} or δ_{ia} set to 0, the prior of the other precision parameter is also assumed to be *Gamma*(2, 2), which gives moderate heterogeneity for latent compliance class probabilities, whereas for the full model, $(\delta_{in}, \delta_{ia})^{T} \sim N(0, \Sigma_{ps})$, the prior for the variance-covariance matrix Σ_{ps} is *InvWishart*(I, 3), where I is the identity matrix.

Similar to cace.study(), to customize the model fully, the user can pass their complete model string with the parameter model.code. Because the function cace.meta.c() is more complicated depending on the choice of random effects, we show an example of the customized prior distributions

```
when assigning delta.n = TRUE, delta.a = TRUE, delta.u = TRUE, delta.v = FALSE, delta.s =
TRUE, delta.b = TRUE, and cor = TRUE while keeping default values for re.values.
string <-</pre>
"# priors
alpha.n \sim dnorm(0, 0.16)
alpha.a ~ dnorm(0, 0.16)
alpha.s \sim dnorm(0, 0.25)
alpha.b ~ dnorm(0, 0.25)
alpha.u ~ dnorm(0, 0.25)
alpha.v ~ dnorm(0, 0.25)
II[1,1] <- 1
II[2.2] <- 1
II[1,2] <- 0
II[2,1] <- 0
Omega.rho ~ dwish (II[,], 3)
Sigma.rho <- inverse(Omega.rho)</pre>
sigma.n <- Sigma.rho[1, 1]</pre>
sigma.a <- Sigma.rho[2, 2]</pre>
rho <- Sigma.rho[1, 2]</pre>
ulout <- phi(alpha.u/sqrt(1+sigma.u^2))</pre>
tau.u ~ dgamma(2, 2)
sigma.u <- 1/sqrt(tau.u)</pre>
v1out <- phi(alpha.v)</pre>
CACE <- u1out-v1out
slout <- ilogit(alpha.s/sqrt(1 + (16^2*3/(15^2*pi^2))*sigma.s^2))</pre>
tau.s ~ dgamma(2, 2)
sigma.s <- 1/sqrt(tau.s)</pre>
blout <- ilogit(alpha.b/sqrt(1 + (16^2*3/(15^2*pi^2))*sigma.b^2))</pre>
tau.b ~ dgamma(2, 2)
sigma.b <- 1/sqrt(tau.b)</pre>
```

The epidural_c dataset is used as a real-study example:

The usage of arguments conv.diag and mcmc.samples is the same as for the function cace.study. When the argument study.specific is specified as TRUE, the model will first check the logical status of arguments delta.u and delta.v. If both are FALSE, meaning that neither response rate u_{i1} or v_{i1} is modeled with a random effect, then the study-specific θ_i^{CACE} is the same across studies. The function gives a warning and continues by making study.specific = FALSE. Otherwise, the study-specific θ_i^{CACE} are estimated and saved as the parameter cacei.

In this example, by calling the object smry from the output list out.meta.c, posterior estimates (posterior mean, standard deviation, posterior median, 95% credible interval, and time-series standard error) are displayed.

out.meta.c\$smry

#>		Mean	SD	2.5%	50%	97.5%	Naive SE	Time-series SE
#>	CACE	0.020200	0.0627	-0.10200	0.018900	0.1490	1.14e-04	7.69e-04
#>	b1out	0.128000	0.0459	0.05970	0.121000	0.2370	8.39e-05	4.07e-04
#>	cacei[1]	0.043900	0.0679	-0.08130	0.040700	0.1870	1.24e-04	2.35e-04
#>	cacei[2]	-0.023100	0.0489	-0.11500	-0.025000	0.0822	8.94e-05	1.89e-04
#>	cacei[3]	-0.007630	0.0569	-0.11000	-0.011800	0.1130	1.04e-04	2.16e-04
#>	cacei[4]	0.065000	0.0678	-0.06620	0.064300	0.2010	1.24e-04	1.62e-04
#>	cacei[5]	0.054000	0.0686	-0.07380	0.051500	0.1960	1.25e-04	2.45e-04
#>	cacei[6]	0.026300	0.0309	-0.03390	0.026200	0.0875	5.64e-05	6.87e-05

#>	cacei[7]	0.002770	0.0931	-0.18900	-0.000142	0.2100 1.70	0e-04	3.71e-04
#>	cacei[8]	0.048300	0.0239	0.00171	0.048200	0.0956 4.36	6e-05	6.38e-05
#>	cacei[9]	-0.010600	0.0224	-0.05520	-0.010500	0.0331 4.09	9e-05	5.49e-05
#>	cacei[10]	0.000228	0.0600	-0.12000	-0.001200	0.1280 1.10	0e-04	2.15e-04
#>	pia	0.114000	0.0742	0.02460	0.098200	0.3010 1.35	5e-04	3.92e-03
#>	pic	0.821000	0.0842	0.60500	0.838000	0.9350 1.54	le-04	4.50e-03
#>	pin	0.064200	0.0397	0.01540	0.056700	0.1590 7.25	5e-05	2.11e-03
#>	s1out	0.184000	0.1040	0.04560	0.161000	0.4450 1.91	e-04	9.03e-04
#>	u1out	0.127000	0.0473	0.05520	0.120000	0.2390 8.64	le-05	6.10e-04
#>	v1out	0.107000	0.0406	0.04700	0.100000	0.2040 7.41	e-05	4.55e-04

The posterior estimates of θ_i^{CACE} can be used to make a forest plot by calling the function plt.forest.

Users can manually do model selection procedures by including different random effects and comparing DIC from the outputs. DIC and its two components are saved as an object DIC in the output list.

out.meta.c\$DIC

#> D.bar 204.40801 #> pD 44.74788 #> DIC 249.15590

DIC is the penalized deviance, calculated as the sum of D.bar and pD, where D.bar is the posterior expectation of the deviance, reflecting the model fit, and pD reflects the effective number of parameters in the model. D.bar is usually lower when more parameters are included in the model, but complex models may lead to overfitting. Thus DIC balances the model's fit against the effective number of parameters. Generally a model with smaller DIC is preferred. However, it is difficult to conclude what constitutes an important improvement in DIC. Following Lunn et al. (2012), we suggest that a reduction of less than 5 is not a substantial improvement. When fitting models to a particular dataset, it is usually uncertain which random effect variables should be included in the model. The function cace.meta.c() allows users to specify candidate models with different random effects, and thus to conduct a forward/backward/stepwise model selection procedure to choose the best fitting model.

Function cace.meta.ic() for meta-analysis with incomplete compliance information

Another major function in the package **BayesCACE** is cace.meta.ic(). It also estimates θ^{CACE} using the Bayesian hierarchical model but can accommodate studies with incomplete compliance data. The arguments of this function are:

```
cace.meta.ic(data, param = c("CACE", "u1out", "v1out", "s1out", "b1out",
  "pic", "pin", "pia"), random.effects = list(), re.values = list(),
  model.code = '', digits = 3, n.adapt = 1000, n.iter = 100000,
  n.burnin = floor(n.iter/2), n.chains = 3, n.thin =
  max(1,floor((n.iter-n.burnin)/100000)), conv.diag = FALSE,
  mcmc.samples = FALSE, study.specific = FALSE)
```

The arguments of cace.meta.ic() are mostly similar to those of cace.meta.c(), although the function cace.meta.ic() calls a different built-in model file from the package **BayesCACE**. The major difference in using this function is that users need to create a dataset with the same structure as epidural_ic. As for cace.meta.c(), users can set their customized prior distributions. Here we use the epidural_ic dataset as an example:

The results are saved in the object out.meta.ic, a list containing posterior estimates for monitored parameters, DIC, convergence diagnostic statistics, and MCMC samples. In this example, the argument study.specific is TRUE, so the summary for each study-specific θ_i^{CACE} is displayed in the object out.meta.ic\$smry together with other parameters.

Note that when compiling the JAGS model, the warning "adaptation incomplete" may occasionally occur, indicating that the number of iterations for the adaptation process is not sufficient. The default value of n.adapt (the number of iterations for adaptation) is 1,000. This is an initial sampling phase during which the samplers adapt their behavior to maximize their efficiency (e.g., a Metropolis–Hastings random walk algorithm may change its step size) (Plummer 2018). The "adaptation incomplete" warning indicates that the MCMC algorithm may not achieve maximum efficiency, but it generally has little impact on the posterior estimates of the treatment effects. To avoid this warning, users may increase n.adapt.

Plotting the trace plot, posterior density, and auto-correlation

When compiling the JAGS models, it is helpful to assess the performance of the MCMC algorithm. The functions plt.trace, plt.density, and plt.acf provide diagnostic plots for the MCMC, namely trace plots, kernel density estimation plots, and auto-correlation plots. Both trace plots and auto-correlation plots can be used to examine whether the MCMC chains appear to be drawn from stationary distributions. A posterior density plot for a parameter visually shows the posterior distribution. Users can simply call this function on objects produced by cace.study(), cace.meta.c(), or cace.meta.ic().

The arguments of this plot function are:

```
plt.trace(obj, param = c("CACE"), trialnumber = 1, ...)
plt.density(obj, param = c("CACE"), trialnumber = 1, ...)
plt.acf(obj, param = c("CACE"), trialnumber = 1, ...)
```

We use the objects list obtained from fitting the Bayesian hierarchical model cace.meta.ic() as an example to generate the three plots. To avoid lengthy output we just illustrate how these plots are produced for θ^{CACE} . The relevant code is:

```
plt.trace(obj = out.meta.ic)
plt.density(obj = out.meta.ic)
plt.acf(obj = out.meta.ic)
```

The produced plots are shown in Figures 2–4. The trace plots in Figure 2 show the parameter values sampled at each iteration versus the iteration number. Each chain is drawn as a separate trace plot to avoid overlay. Here we used the default n.chains = 3, so three trace plots are drawn. These plots show evidence that the posterior samples of θ^{CACE} are drawn from the stationary distribution.

The density plot in Figure 3 is smoothed using the R function density(). It shows that the kernelsmoothed posterior of θ^{CACE} is almost symmetric. The posterior mean is not far from 0, indicating that the complier average causal effect of using epidural analgesia in labor on cesarean section is likely not significant.

The auto-correlation plot in Figure 4 is a bar plot displaying the auto-correlation for different lags. At lag 0, the value of the chain has perfect auto-correlation with itself. As the lag becomes greater, the values become less correlated. After a lag of about 50, the auto-correlation drops below 0.1. If the plot shows high auto-correlation, users can run the chain longer or can choose a larger n.thin, e.g., n.thin = 10 would keep only 1 out of every 10 iterations, so that the thinned out chain is expected to have the auto-correlation drop quickly. Any additional parameters passed to the 3 plotting function will be automatically used in the plot function for plt.trace and plt.density, and in the acf function for plt.acf.

Plotting the study-specific CACE in a forest plot

A graphical overview of the results can be obtained by creating a forest plot (Lewis and Clarke 2001). The function plt.forest() draws a forest plot for θ^{CACE} estimated from the meta-analysis. Users can call this function for the objects from cace.meta.c() or cace.meta.ic(). Here is an example using the object out.meta.ic:

plt.forest(data = epidural_ic, obj = out.meta.ic)

Note that in addition to the object out.meta.ic, users also need to specify the dataset used to compute that object, from which the plt.forest() function extracts the study names and publication years for the figure.

Figure 5 is a forest plot of θ_i^{CACE} for each study individually, using the Bayesian method with full random effects and default priors. The summary estimate based on the model cace.meta.ic()



Figure 2: Trace plots for θ^{CACE} from the epidural_ic dataset fit using cace.meta.ic() for a sample of 3 chains. Because there are no strong patterns and the variability is relatively constant, we can conclude that the posterior means are drawn from a stationary distribution.

Density of CACE



Figure 3: The kernel smoothed density for θ^{CACE} from the function cace.meta.ic() applied to the epidural analgesia in labor meta-analysis. The posterior mean is close to 0, indicating that the complier average causal effect may not be significant in this case.



Figure 4: Auto-correlation plot of θ^{CACE} from the model cace.meta.ic() fit to the epidural_ic dataset. As the lag increases, the values become less correlated. Users can choose to address high auto-correlation with a longer chain or a larger n.thin.



Figure 5: Forest plot of study-specific θ^{CACE} from the model cace.meta.ic() with full random effects fit to the epidural_ic dataset. The summary estimate and confidence interval limits based on the model cace.meta.ic() are included in the figure, both in terms of written values and the squares and lines on the right. Overall, it shows that the study-specific θ_i^{CACE} vary from negative to positive in individual studies, while most of the 95% credible intervals cover zero.

4 Discussion

This article provides an overview of the **BayesCACE** package for conducting CACE analysis with R. Bayesian hierarchical models estimating the CACE in individual studies and in meta-analysis are introduced to demonstrate the underlying methods of the functions. Practical usage of various functions is illustrated using real meta-analysis datasets epidural_c and epidural_ic. The package provides several plots for model outputs and model diagnosis.

trial without complete compliance data is not estimable using only data from that single trial, dashed

lines tend to have longer credible intervals than those with complete data (solid lines).

It is important to note that the two-step approach for meta-analysis is included in the package **BayesCACE** because by using the full observed data from a single study *i*, θ_i^{CACE} is identifiable, making it possible to pool the estimated posterior means and standard deviations of the θ_i^{CACE} in a meta-analysis. However, the Bayesian hierarchical-model meta-analysis method for estimating the overall CACE is preferred for two reasons: the conventional two-step approach requires the whole set of parameters to be estimated for each trial, giving a greater total number of parameters than the random effect model, so the estimate of the CACE can be less efficient. Also, when study *i* does not report complete compliance data, it must be excluded from the two-step approach because θ_i^{CACE} is no longer directly estimable by simply using the incomplete data from this individual study, while the function cace.meta.ic() can use the incomplete information and thus help improve the efficacy in estimation.

The Gelman and Rubin convergence statistics, time-series standard errors, trace plots, and autocorrelation plots are provided by the package **BayesCACE** to examine whether the MCMC chains are drawn from stationary distributions. However, in practice, any sample is finite, thus there is no guaranteed way to prove that the sampler has converged (Kass et al. 1998; Cowles and Carlin 1996). Additional techniques may be required to determine the effective sample size for adequate convergence (Robert and Casella 2004). For example, the well-developed R package mcmcse (Flegal et al. 2017) can be used to assess whether MCMC has been run for enough iterations (sufficient chain lengths). To call the functions in mcmcse, users can specify the argument mcmc.samples = TRUE in cace.study(), cace.meta.c(), and cace.meta.ic(), so the MCMC posterior samples of monitored parameters are saved in the output objects.

The current version of **BayesCACE** only applies to binary outcomes. However, the Bayesian hierarchical model can be extended to handle ordinal outcomes $o \in \{1, ..., O\}$. By selecting weighting scores $\{W_1, W_2, ..., W_O\}$ to reflect distances between outcome categories $\{1, ..., O\}$, θ_i^{CACE} is defined as $E(Y_{ij}^1 - Y_{ij}^0|C_{ij} = 1) = \sum_o (W_o \times u_{io}) - \sum_o (W_o \times v_{io})$ (J. Zhou, Hodges, and Chu 2021; J. Zhou et al. 2019). Equally spaced scores $\{1, 2, ..., O\}$, their linear transforms, and midranks are reasonable weight choices (Agresti 2013). Future work will add CACE meta-analysis functions for ordinal outcomes, and allow users to choose their preferred weights $\{W_1, W_2, ..., W_O\}$. Note that ordinal outcomes lead to more complex correlation structures in the parameters related to

References

Agresti, Alan. 2013. Categorical Data Analysis. Third Edition. Hoboken, NJ: John Wiley & Sons.

response rates, so multivariate prior distributions are necessary to analyze such outcomes.

- Baccini, Michela, Alessandra Mattei, and Fabrizia Mealli. 2017. "Bayesian Inference for Causal Mechanisms with Application to a Randomized Study for Postoperative Pain Control." *Biostatistics* 18 (4): 605–17.
- Baker, Stuart G. 2020. "CACE and Meta-Analysis (Letter to the Editor)." Biometrics 76 (4): 1383-84.
- Bannister-Tyrrell, Melanie, Branko Miladinovic, Christine L Roberts, and Jane B Ford. 2015. "Adjustment for Compliance Behavior in Trials of Epidural Analgesia in Labor Using Instrumental Variable Meta-Analysis." Journal Article. *Journal of Clinical Epidemiology* 68 (5): 525–33.
- Brooks, Stephen P, and Andrew Gelman. 1998. "General Methods for Monitoring Convergence of Iterative Simulations." Journal Article. *Journal of Computational and Graphical Statistics* 7 (4): 434–55.

- Clopper, Charles J, and Egon S Pearson. 1934. "The Use of Confidence or Fiducial Limits Illustrated in the Case of the Binomial." *Biometrika* 26 (4): 404–13.
- Coggeshall, Scott. 2017. noncomplyR: Bayesian Analysis of Randomized Experiments with Non-Compliance. https://CRAN.R-project.org/package=noncomplyR.
- Cowles, Mary Kathryn, and Bradley P Carlin. 1996. "Markov Chain Monte Carlo Convergence Diagnostics: A Comparative Review." *Journal of the American Statistical Association* 91 (434): 883– 904.
- Flegal, James M, John Hughes, Dootika Vats, and N Dai. 2017. mcmcse: Monte Carlo Standard Errors for MCMC. https://CRAN.R-project.org/package=mcmcse.
- Frangakis, Constantine E, and Donald B Rubin. 2002. "Principal Stratification in Causal Inference." Biometrics 58 (1): 21–29.
- Freedman, Laurence S. 1990. "The Effect of Partial Noncompliance on the Power of a Clinical Trial." Controlled Clinical Trials 11 (3): 157–68.
- Gelman, Andrew, and Donald B Rubin. 1992. "Inference from Iterative Simulation Using Multiple Sequences." Journal Article. *Statistical Science* 7 (4): 457–72.
- Gilbert, Peter B, Erin E Gabriel, Ying Huang, and Ivan SF Chan. 2015. "Surrogate Endpoint Evaluation: Principal Stratification Criteria and the Prentice Definition." *Journal of Causal Inference* 3 (2): 157–75.
- Gordon, Max, and Thomas Lumley. 2017. forestplot: Advanced Forest Plot Using "Grid" Graphics. https://CRAN.R-project.org/package=forestplot.
- Harville, David A. 1977. "Maximum Likelihood Approaches to Variance Component Estimation and to Related Problems." *Journal of the American Statistical Association* 72 (358): 320–38.
- Hedges, Larry V, and Ingram Olkin. 1985. *Statistical Methods for Meta-Analysis*. Orlando, FL: Academic Press.
- Hedges, Larry V, and Jack L Vevea. 1998. "Fixed- and Random-Effects Models in Meta-Analysis." Psychological Methods 3 (4): 486–504.
- Hudgens, Michael G, and M Elizabeth Halloran. 2006. "Causal Vaccine Effects on Binary Postinfection Outcomes." *Journal of the American Statistical Association* 101 (473): 51–64.
- Jiang, Yang, and Dylan Small. 2014. ivpack: Instrumental Variable Estimation. https://CRAN.Rproject.org/package=ivpack.
- Kasim, Adetayo, ZhiMin Xiao, Steve Higgings, and Ewoud De Troyer. 2017. eefAnalytics: Analysing Education Trials. https://CRAN.R-project.org/package=eefAnalytics.
- Kass, Robert E, Bradley P Carlin, Andrew Gelman, and Radford M Neal. 1998. "Markov Chain Monte Carlo in Practice: A Roundtable Discussion." *The American Statistician* 52 (2): 93–100.
- Laird, Nan M, and Frederick Mosteller. 1990. "Some Statistical Methods for Combining Experimental Results." International Journal of Technology Assessment in Health Care 6 (1): 5–30.
- Lewis, Steff, and Mike Clarke. 2001. "Forest Plots: Trying to See the Wood and the Trees." *BMJ* 322 (7300): 1479–80.
- Lin, D Y, and D Zeng. 2010. "On the Relative Efficiency of Using Summary Statistics Versus Individual-Level Data in Meta-Analysis." *Biometrika* 97 (2): 321–32.
- Lunn, David, Chris Jackson, Nicky Best, David Spiegelhalter, and Andrew Thomas. 2012. *The BUGS Book: A Practical Introduction to Bayesian Analysis*. New York, NY: Chapman; Hall/CRC.
- Plummer, Martyn. 2018. rjags: Bayesian Graphical Models Using MCMC. https://CRAN.R-project. org/package=rjags.
- Plummer, Martyn, Nicky Best, Kate Cowles, and Karen Vines. 2006. "CODA: Convergence Diagnosis and Output Analysis for MCMC." *R News* 6 (1): 7–11.
- Robert, Christian, and George Casella. 2004. *Monte Carlo Statistical Methods*. New York, NY: Springer Science & Business Media.
- Spiegelhalter, David J, Nicola G Best, Bradley P Carlin, and Angelika Van Der Linde. 2002. "Bayesian Measures of Model Complexity and Fit." Journal Article. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)* 64 (4): 583–639.
- Viechtbauer, Wolfgang. 2010. "Conducting Meta-Analyses in r with the metafor Package." Journal of Statistical Software 36 (3): 1–48.
- White, Halbert. 1982. "Instrumental Variables Regression with Independent Observations." *Econometrica* 50 (2): 483–99.
- Zeger, Scott L, Kung-Yee Liang, and Paul S Albert. 1988. "Models for Longitudinal Data: A Generalized Estimating Equation Approach." *Biometrics* 44 (4): 1049–60.
- Zhou, Jincheng, Haitao Chu, Michael G Hudgens, and M Elizabeth Halloran. 2016. "A Bayesian Approach to Estimating Causal Vaccine Effects on Binary Post-Infection Outcomes." Statistics in Medicine 35 (1): 53–64.
- Zhou, Jincheng, James S Hodges, and Haitao Chu. 2020. "Rejoinder to 'CACE and Meta-Analysis (Letter to the Editor)' by Stuart Baker." *Biometrics* 76 (4): 1385.
 - —. 2021. "A Bayesian Hierarchical CACE Model Accounting for Incomplete Noncompliance with Application to a Meta-Analysis of Epidural Analgesia on Cesarean Section." *Journal of the American Statistical Association* 116 (536): 1700–1712.

- Zhou, Jincheng, James S. Hodges, M. Fareed K. Suri, and Haitao Chu. 2019. "A Bayesian Hierarchical Model Estimating CACE in Meta-Analysis of Randomized Clinical Trials with Noncompliance." *Biometrics* 75 (3): 978–87.
- Zhou, Ting, Jincheng Zhou, James S Hodges, Lifeng Lin, Yong Chen, Stephen R Cole, and Haitao Chu. 2021. "Estimating the Complier Average Causal Effect in a Meta-Analysis of Randomized Clinical Trials with Binary Outcomes Accounting for Noncompliance: A Generalized Linear Latent and Mixed Model Approach." American Journal of Epidemiology 191 (1): 220–29.

Jincheng Zhou Gilead Inc. Clinical Data Science Foster City, CA 94404, USA jeni.zhou9@gilead.com

Jinhui Yang University of Minnesota Twin Cities Department of Computer Science and Engineering Minneapolis, MN 55455, USA yang7004@umn.edu

James S. Hodges University of Minnesota Twin Cities Division of Biostatistics School of Public Health Minneapolis, MN 55455, USA hodge003@umn.edu

Lifeng Lin University of Arizona Department of Epidemiology and Biostatistics Mel and Enid Zuckerman College of Public Health Tucson, AZ 85724, USA lifenglin@arizona.edu

Haitao Chu (Affliation 1) Pfizer Inc. (Affliation 2) University of Minnesota Twin Cities (1) Statistical Research and Data Science Center New York, NY 10017, USA (2) Division of Biostatistics School of Public Health Minneapolis, MN 55455, USA (1) Haitao.Chu@Pfizer.com (2) chux0051@umn.edu